

Alkali pyroxenes and amphiboles: a window on rare earth elements and other high field strength elements behavior through the magmatic-hydrothermal transition of peralkaline granitic systems

Cyrielle Bernard, Guillaume Estrade, Stefano Salvi, Didier Béziat, Martin J

Smith

▶ To cite this version:

Cyrielle Bernard, Guillaume Estrade, Stefano Salvi, Didier Béziat, Martin J Smith. Alkali pyroxenes and amphiboles: a window on rare earth elements and other high field strength elements behavior through the magmatic-hydrothermal transition of peralkaline granitic systems. Contributions to Mineralogy and Petrology, 2020, 10.1007/s00410-020-01723-y. hal-02989854

HAL Id: hal-02989854 https://hal.science/hal-02989854

Submitted on 5 Nov 2020

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.

1	Alkali pyroxenes and amphiboles: a window on Rare Earth Elements and other High Field Strength
2	Elements behavior through the magmatic-hydrothermal transition of peralkaline granitic systems
3	
4	Cyrielle Bernard ^a , Guillaume Estrade ^{a□} , Stefano Salvi ^a , Didier Béziat ^a , Martin Smith ^b
5	^a GET, CNRS, UPS, Université de Toulouse III, Toulouse, France
6	^b School of Environment and Technology, University of Brighton, Brighton, BN2 4GJ, UK
7	
8	Corresponding author: guillaume.estrade@get.omp.eu
9	ORCID: 0000-0001-6907-7469
10	
11	
12	Acknowledgments
13	This work was supported by an INSU/TelluS grant from CNRS (French National Center for Scientific
14	Research). We thank Anthony Williams-Jones, Olga Vasyukova and Sam Broom-Fendley for providing some of
15	the rock samples from the Strange Lake and Amis complexes and David Chew who helped with LA-ICP-MS
16	mapping. Kathryn Goodenough provided helpful discussion. We also wish to thank CPM reviewers Charles
17	Beard and Adrian Finch for their suggestions and constructive criticisms, which greatly improved the final
18	version of this manuscript.
19	
20	
21	
22	
23	
24	

1

25

26

27 <u>Abstract</u>

28 Peralkaline granites and pegmatites are a prime repository of REE and HFSE, critical raw materials. 29 Although it is accepted that magmatic processes are fundamental in concentrating these metals, the role of 30 hydrothermal fluids in concentrating and fractionating these elements remains unclear. This paper investigates 31 the global reproducibility of the magmatic-hydrothermal evolution of alkaline silica-saturated systems using 32 alkali pyroxene and amphiboles from six alkaline complexes. These minerals contain significant amounts of 33 REE and other HFSE, and pyroxene is stable throughout the magmatic and hydrothermal stages. Amphibole 34 consists of mostly unzoned arfvedsonite, leakeite, and katophorite, while pyroxene is always aegirine. Two types 35 of aegirine were defined. In all complexes, type-I aegirine is zoned; its core is enriched in Ca, REE, Zr, Hf, Sc and Sn, and the rims in Na, Fe³⁺ and contain secondary rare-metal bearing minerals and fluid inclusions. Type-II 36 37 aegirine replaces amphibole and is oscillatory zoned. We interpret the amphiboles and REE-rich cores of type-I 38 aegirine to have grown during the magmatic stage, whereas the rims of REE-poorer type-I and II aegirine formed 39 during the hydrothermal stage. During magmatic crystallization, REE intake into amphiboles and pyroxene as 40 well as LREE-HREE fractionation were favored by their crystallographic properties and by competition among 41 them and other minerals. During subsequent hydrothermal stages, REE and other HFSE were remobilized, 42 locally reconcentrated and fractionated in mineral pseudomorphs and secondary pyroxene. These observations 43 point out the importance of studying rock-forming minerals such as pyroxenes and amphiboles to unravel 44 geological events controlled by common processes globally.

45

46 Keywords

47 Peralkaline granite, pegmatite, pyroxene, amphibole, rare earth elements, high field strength elements

48

49 Introduction

High Field Strength Elements (HFSE, namely Nb, Ta, Zr, Hf, U, Th, Sn, and Ti), and in particular Rare
Earth Elements (REE), are among raw materials considered the most critical today (European Commission
2018). They are used in many modern technologies linked to the transition to renewable energy infrastructure

53 (Lucas et al. 2014; Goodenough et al. 2018), but their production is quite restricted, worldwide. REE deposits 54 can be primary (igneous, carbonatites and alkaline systems) or secondary (placers and ion-adsorption) 55 (Chakhmouradian and Wall 2012) and economic concentrations of heavy REE (HREE, Gd to Lu) are rather rare 56 compared to those of light REE (LREE, La to Eu). The latter occur in high amounts mostly in carbonatites and 57 placer deposits and are commonly hosted in fluorcarbonates such as bastnäsite-(Ce), parisite-(Ce) and 58 synchysite-(Ce), and phosphate such as monazite-(Ce). Ion-adsorption deposits provide most of the global HREE 59 production (Chakhmouradian and Wall 2012; U. S. Geological Survey 2019). In alkaline igneous rocks, REE are 60 hosted mainly in monazite-(Ce), xenotime-(Y), fergusonite-(Y), loparite-(Ce) and eudialyte-group minerals 61 (EGM) (Larsen and Sørensen 1987; Chakhmouradian and Wall 2012). Most alkaline igneous rocks are silica-62 undersaturated, with the silica-saturated variety (i.e., granitic) being quite rare (e.g. Larsen and Sørensen 1987; 63 Ranløv and Dymek 1991; Foland et al. 1993; Kramm and Kogarko 1994; Sheard et al. 2012). However, it is the 64 latter that have high Yb/La ratios compared to other varieties (Fig 1). Pegmatites, which form from the last and 65 most evolved part of silica saturated melts, concentrate the highest amounts of REE and other HFSE.

66 There is a general consensus in the literature that magmatic processes play a role for a pre-enrichment 67 of alkaline rocks in REE and other HFSE (Marks and Markl 2017). These include sourcing from undepleted, 68 more or less metasomatized mantle, crustal contamination, fractional crystallization, and melt-melt immiscibility 69 during magma cooling (Bonin 2007; Nardi and Bitencourt 2009; Chakhmouradian and Zaitsev 2012; Veksler et 70 al. 2012). There is also evidence that hydrothermal processes may play a key role in concentrating these 71 elements to ore grades (e.g. Salvi and Williams-Jones 1990; William-Jones et al. 2012). However, to date, the 72 relative importance of magmatic versus hydrothermal processes is still debated, partly because the processes 73 occurring at the magmatic-hydrothermal transition are hard to document and most studies were carried out in the 74 laboratory (e.g. Migdisov et al. 2016).

75 To fill this gap, it is important to understand the timing of concentration of the REE in natural occurrences. 76 To do so, in this paper we propose for the first time a detailed study of common minerals that carry REE and 77 other HFSE from the magmatic through the hydrothermal stages, from six silica-saturated alkaline occurrences 78 worldwide. The goal is to make out the meaning of textures and link them to the major events that impacted the 79 selected complexes, i.e. the magmatic-hydrothermal transition. Once it is clear which zones of the crystals grew 80 during the magmatic phase and which grew during hydrothermal circulations, we can analyze the REE and other 81 HFSE compositions in the different zones, and infer the timing of concentration and fractionation of the REE. 82 Ideal candidates to do so are the pyroxene and amphibole mineral groups, because of their ubiquitous presence in

83 these rocks and because they have been documented to form during a large span of the evolution of alkaline 84 rocks, from early magmatic to late hydrothermal stages (e.g. Nielsen 1979; Jones and Peckett 1981; Ranløv and Dymek 1991; Marks et al. 2004). In addition, they incorporate a relatively high amount of REE and other HFSE 85 86 compared to other rock-forming minerals (e.g. alkali feldspar and quartz), even though they are not considered as 87 ore as their partition coefficients for HFSE (except Eu) are not high enough (Fedele et al. 2015). Primary 88 zirconosilicates can also incorporate high levels of HFSE and are common in alkaline rocks, however, they are 89 frequently altered to secondary minerals during the hydrothermal stage. Hence, pyroxenes and amphiboles are 90 the best minerals to evidence the behavior of HFSE throughout the entire process of alkaline granite and 91 pegmatite crystallization.

92

93 1. <u>Geological background</u>

94 The six alkaline igneous complexes on which this work focuses are: Ambohimirahavavy (Estrade et al. 95 2014a) and Manongarivo (Donnot 1963) in Madagascar, Amis in Namibia (Schmitt et al. 2002), Evisa in Corsica 96 (Bonin et al. 1978), Khan Bogd in Mongolia (Kovalenko et al. 2006), and Strange Lake in Canada (Currie 1985). 97 These complexes were chosen because all present evidence of hydrothermal circulation and contain high 98 amounts of REE (from 800 ppm in a granite from Khan Bogd to 12 % in a pegmatite from Strange Lake, 99 unpublished data). All complexes contain peralkaline rocks, i.e. their ratio of $(Na_2O+K_2O)/Al_2O_3$ is >1. Some of 100 them have been the subject of in-depth studies, such as the Strange Lake complex which was first studied in the 101 late 1980's (e.g. Miller, 1986; Salvi and Williams-Jones 1990) and still is today (Vasyukova and Williams-Jones 102 2020); others are poorly known, as is the case for the Manongarivo complex, for which the only available 103 information is a mere mention in a French compilation (Donnot, 1963).

104 Although they were emplaced in different geodynamic complexes and at different periods, it appears that the 105 melt source was a metasomatized upper mantle for all six complexes. Crustal contamination and feldspars 106 fractionation, if occurring, are two processes that can enrich alkaline granites in REE. However, the rate of 107 partial melting at the origin of these complexes is still debated, which is an essential parameter given that a lower 108 rate implies a greater accumulation of incompatible elements, REE included, in the newly generated melt. All 109 complexes underwent at least one alteration episode from an orthomagmatic fluid, and some of them an 110 additional late hydrothermal event (identified at Amis, Evisa, and Strange Lake). The composition of the fluids 111 that circulated was measured in fluid inclusions at Ambohimirahavavy and Strange Lake, and inferred from the

112 study of secondary mineral assemblages for the other complexes. The presence of significant amounts of Na⁺ and Fe³⁺ in an orthomagmatic fluid is reported for Evisa, the two Malagasy complexes and Strange Lake, and 113 Ca²⁺ was identified at Khan Bogd, Madagascar and Strange Lake. Locally, CO₂ and/or CH₄ were reported at 114 Khan Bogd and Strange Lake, and CO_3^{2-} at Amis. The resulting conclusions are similar for all the complexes: 115 116 independently of its orthomagmatic or late origin, a F-, Ca-rich fluid is systematically observed and inferred to 117 have mobilized and fractionated the REE. The importance of this process compared to magmatic ones in 118 concentrating and fractionating the REE is still a matter of debate. Comparing six complexes that were emplaced 119 in different geodynamic settings and witnessed different fluid conditions (composition, temperature), is a 120 strategy to highlight common features in order to better constrain and generalize the timing of REE and other 121 HFSE enrichment in alkaline complexes. Below, follows a rapid summary of the main geological features of 122 each complex; more detailed information is provided in Online Resource 1.

123

124 1.1. <u>Amis complex, Namibia</u>

125 The Amis complex and the larger Brandberg complex onto which it is juxtaposed are of the same age (132.5 126 to 130.5 My) and are hosted in metasediments of the Damara orogen (~ 550 My; Miller 1983) and volcanic 127 rocks from the Paraná-Etendeka igneous province. Three granite and 4 pegmatite-aplite samples were used for 128 this study. The Amis complex is mainly made of an arfvedsonite granite and aegirine-rich pegmatite-aplites. The 129 Amis complex is highly enriched in REE, other HFSE and volatiles (H₂O, F). Pegmatites and aplites in the 130 north-western part of the Amis complex are particularly enriched in REE mineralizations and uranium. Schmitt 131 et al. (2002) infer that hydrothermal fluids had a very limited impact on the rocks of Amis complex. They affirm that minerals are all mainly magmatic, with the exception of replacement of arfvedsonite by quartz-hematite 132 133 overgrowths. Based on the study of melt inclusions, they propose that the main process of REE and other HFSE 134 enrichment is magmatic. In contrast, Diehl (1990) proposed that hydrothermal fluids played a significant role, 135 and inferred the replacement of arfvedsonite by aegirine.

136

1.2. <u>Evisa complex, Corsica</u>

137

The Evisa complex (290 My; Cocherie et al. 2005) was emplaced among metaluminous granitoids. We collected 7 granite and 5 pegmatite samples from the central part of the complex. The complex mainly comprises hypersolvus (perthitic feldspar) and subsolvus (albitic and alkali feldspar) peralkaline granites (Bonin 1990) and

141 associated pegmatites rich in REE-bearing minerals such as monazite-(Ce), apatite, and allanite-(Ce). The 142 hypersolvus granite probably experienced two hydrothermal events: a minor, early episode at the end of 143 crystallization, and a more important second episode, characterized by F-rich fluids, around 200 My (Poitrasson 144 et al. 1998; Bonin et al. 2008). The subsolvus granite is intrusive into the perthitic unit and only experienced the 145 second hydrothermal event, which triggered the replacement of the primary zirconosilicate elpidite by a 146 secondary assemblage (Bonin 1990).

147

148 *1.3. <u>Khan Bogd complex, Mongolia</u>*

149 The Khan Bogd complex is located in the southern Gobi Desert, at the transition between island-arc calc-alkaline 150 differentiated volcanics (329±5 My) and rift-related bimodal basalt-comendite-alkali granite association (318-151 290 My), and is dated at approximately 290 Ma (Kovalenko et al. 2006). It is among the largest alkaline granite 152 plutons in the world, with a surface area of 1500 km². We collected 3 granite and 7 pegmatite samples, mostly on 153 the western side of the complex. The pluton consists of two ring bodies: a western peralkaline arfvedsonite-154 bearing granite and a later emplaced, eastern aegirine granite. They are both associated with pegmatites. The 155 pluton has been investigated for mining purposes and estimates indicate grades between 0.3 and 4.5 % REE, the 156 highest grade being located at the top of the granite plutons (estimation of the tonnage is not available, 157 Koyalenko and Yarmolyuk 1995). The origin of this pluton is linked to a mantle plume and the continental 158 subduction of the South Mongolian Hercynides, but the exact magmatic process is still debated (Kovalenko et al. 159 2006; Kynicky et al. 2011). Kynicky et al. (2011) propose the release of a silica-saturated orthomagmatic fluid 160 that, similarly to Evisa, triggered the replacement of primary elpidite.

161

162 *1.4. <u>Manongarivo and Ambohimirahavavy complexes, Madagascar</u>*

163

Ambohimirahavavy and Manongarivo are the biggest complexes of the Ampasindava province in northwestern Madagascar, yet are not well documented. They were emplaced at circa 24 Ma into marine-shelf carbonates and marine-fluvial siliciclastic sediments of the Isalo Group (Thomas et al. 2009; Cucciniello et al. 2016). We collected 3 granite and 4 pegmatite samples from the southern part of the Ambohimirahavavy complex. Due to poor outcropping conditions and low occurrence of pegmatites, we could only obtain 3 169 pegmatite samples from Manongarivo. The samples from Ambohimirahavavy were selected according to 170 previous studies from Estrade et al. (2014a, b). Both complexes were likely emplaced in a rifting context and are 171 made of two ring-shaped intrusions side by side, composed of nepheline syenite, alkali feldspar syenite, biotite 172 granite, peralkaline pegmatites, and various volcanic rocks. Due to the recent discovery of rare metals in 173 economic concentrations in ion-adsorption clays, the province drew the attention of a mining exploration 174 company (Estrade et al. 2014b). Granite and pegmatite dykes are the most enriched lithologies in REE and other 175 HFSE. An orthomagmatic fluid metasomatized the complexes in the very last stages of magmatic evolution 176 (Estrade et al. 2014b) and caused the replacement of primary zirconosilicates (eudialyte group minerals (EGM) 177 at Ambohimirahavavy, unknown at Manongarivo) by an assemblage of secondary HFSE-bearing minerals 178 (Lacroix 1923; Rakotovao et al. 2009; Estrade et al. 2018).

179

180 1.5. <u>Strange Lake complex, Canada</u>

181 The Strange Lake complex was emplaced 1240 My ago into Paleoproterozoic gneisses and quartz 182 monzonite (Miller 1996) and is considered to represent an extension of the Gardar rift in Greenland (Pillet et al. 183 1989; Boily and Williams-Jones 1994; Siegel et al. 2017a). We chose 3 granite and 4 pegmatite samples which 184 were selected according to previous studies from Salvi and Williams-Jones (1990, 2006; 1995). The Strange 185 Lake complex consists of a hypersolvus granite, while a younger, largely metasomatized transsolvus granite 186 (rare perthite plus two distinct feldspars, previously classified as subsolvus, e.g. Boily and Williams Jones, 1994) 187 surrounds it, forming the majority of the complex (Gysi et al. 2016; Siegel et al. 2017b). These plutons are 188 partially bounded by an outwardly-dipping fracture associated with fluorite and hematite breccia. Two zones of 189 pegmatites have been reported: the Main zone in the center of the complex, and the B zone on the northwestern 190 edge. These magmatic events correlate with a progressive enrichment in REE and other HFSE, from HFSE-poor 191 hypersolvus granite to HFSE-rich transsolvus granite and pegmatites (Miller 1996). Most of the REE are 192 concentrated in the pegmatites. Reserves are estimated at 278 Mt grading 0.93% REE₂O₃ of which 39% are 193 HREE, and 214 Mt at 0.85 % REE₂O₃ in total (Gowans et al. 2017). The complex also contains significant 194 amounts of ZrO₂, Nb₂O₅, and BeO. Two main extensive hydrothermal events affected the complex. The first 195 event is attributed to circulation of a hot (≥ 300 °C) orthomagmatic brine consisting of K- and Na-enriched 196 aqueous and carbonic phases (Salvi and Williams-Jones 2006; Vasyukova and Williams-Jones 2019). The second hydrothermal event consists of a cooler fluid (100 - 200 °C) resulting from mixing of a meteoric fluid
and a fluid originating from the granites (Gysi et al. 2016).

199

200

201

2. <u>Features common to pegmatites and granites from all six complexes</u>

Pegmatites in all complexes are heterogeneous in texture, i.e. they show mineralogical layering visible from the scale of the outcrop to that of a thin section (Fig 3), whereas granites are generally homogeneous. In pegmatites, the layers have grain sizes ranging from a few μ m to about 40 cm and locally more (Fig 3). Therefore, to obtain meaningful data, samples for this study were selected from zones of relatively fine grain size (less than a few cm).

All pegmatites and granites are dominated by quartz, alkali feldspar (perthites, albite, orthoclase, and/or microcline) that are commonly strongly albitized, plus alkali amphiboles and, mostly in pegmatites, aegirine (Fig 4). Common minerals in both rock types include zircon, pyrochlore group minerals (PGM) and Fe- and Tioxides. Other minerals present mostly in pegmatites include Ca and Na zirconosilicates, fluorite, astrophyllite and REE-bearing minerals (typically bastnäsite-(Ce), monazite-(Ce), xenotime-(Y), chevkinite-(Ce), allanite-(Ce), aeschynite-(Y), fergusonite-(Y), britholite-(Y), synchysite-(Ce)). Ti-bearing minerals such as chevkinite-(Ce) and astrophyllite show signs of alteration such as dissolved edges (Fig 4a).

214 Ca and Na zirconosilicates, which are characteristic minerals of agpaitic rocks, are mostly eudialyte-group 215 minerals (EGM) at Ambohimirahavavy (Lacroix 1923; Estrade et al. 2018), and elpidite at Evisa, Khan Bogd 216 and Strange Lake (Bonin 1988; Salvi and Williams-Jones 1990; Grigor'eva et al. 2011). These zirconosilicates 217 were partially to totally replaced by secondary miaskitic mineral assemblages, which commonly form 218 pseudomorphs (Poitrasson et al. 1998; Estrade et al. 2014b; Gysi et al. 2016). The mineralogy of the 219 pseudomorphs varies in the different complexes and consists either of different Zr- and/or REE-bearing minerals 220 (Fig. 4b), or exclusively zircon plus quartz, i.e. typical miaskitic mineral assemblage (Fig 4c). Both 221 pseudomorph types are found in all complexes, except in Amis where our samples only contain the miaskitic 222 zircon-quartz type. Elpidite can also be extensively replaced by other zirconosilicates, namely armstrongite and 223 gittinsite (e.g. Salvi and William-Jones 1995), whereas EGM is only partially replaced by these phases. At Khan 224 Bogd, cathodoluminescence reveals the elpidite is altered from its core, replaced by zircon and armstrongite (Fig 225 4d). Pseudomorphs from Evisa consist of Ca-rich mineral assemblages such as fluorite and parisite-(Ce), the latter growing mostly at the expense of bastnäsite-(Ce). At Manongarivo and Amis, the precursor mineral has notbeen identified because partial replacement has not been observed in these complexes, so far.

228

229 3. <u>Analytical methods</u>

Polished thin sections were prepared from all samples from the six complexes and were studied using
 transmitted light microscopy, cathodoluminescence, Scanning Electron Microscopy (SEM), microprobe, and
 Laser-Ablation Inductively Coupled Plasma Mass Spectrometry (LA-ICPMS) in order to determine mineral
 textural relationships and amphiboles and pyroxene zoning.

SEM images in back-scattered electron (BSE) mode were collected with a Jeol JSM6360LV SEM coupled
with a Bruker silicon drift detector and interpreted with the Esprit software at the Géosciences Environnement
Toulouse laboratory, using an accelerating voltage of 20 kV, and a detection time of 40 s.

237 Mineral compositions were determined at the Centre Raymond Castaing in Toulouse, using a CAMECA 238 SX Five electron microprobe (EPMA), with an accelerating voltage of 15 kV, a beam current of 20 nA for 239 pyroxene, 10 nA for amphibole, and a beam diameter of about 2 μ m. Quantitative elemental maps were 240 performed on the same instrument, using an accelerating voltage of 15 kV, a beam current of 20 nA and 100 nA 241 for major and minor elements respectively. Counting time was 1 s on each pixel with a step of 0.5 to 4 μ m 242 depending on crystal size. Uncertainties were calculated using the method of Ancey et al. (1978). Standards used 243 for calibration as well as detection limits based on repeated measurements are provided in Online Ressource 2.

To resolve the oxidation state of iron measured by microprobe, calculations of the valence were made 244 following Droop (1987). In order to verify these measurements, a ⁵⁷Fe Mössbauer spectrum was collected at the 245 246 Laboratoire de Chimie de Coordination - University of Toulouse on a constant-acceleration conventional 247 spectrometer, at 80 K and with a 1.85 GBq source of 57Co, over a ±12 mm/s range with 512 channels. The 248 spectra were acquired via a compact detector c-system and recorded with a Canberra multichannel analyzer, 249 coupled to a computer with Recoil Mossbauer Analysis Software 1.05 from Lagarec and Rancourt (1997). 250 Uncertainties were calculated using the covariance matrix. Pyroxene end-members were then calculated 251 following Mann et al. (2006). Amphiboles end-members were calculated using the spreadsheet available from 252 Locock (2014).

253 Trace element concentrations were determined in situ by LA-ICPMS at the Géosciences Environnement 254 Toulouse laboratory, University of Toulouse, on 30-µm polished thin-sections, using a New Wave Research ESI 255 213 laser coupled to a Thermo-Fisher Element-XR high-resolution ICP-MS. Laser beam diameter varied from 30 256 to 50 µm depending on mineral size, and data were acquired by ablating lines instead of spots in order to prevent 257 intersecting solid inclusion or internal zoning. NIST synthetic glass certified reference materials SRM 610 and 258 SRM 612 were used as external and quality control standards, respectively. Each analysis was normalized using 259 SiO₂ values previously determined by EMPA. The relative precision ranged from 5 to 10% and detection limits, 260 based on repeated measurements, are provided in Online Ressource 3. Trace elements maps were performed at 261 the Trinity College in Dublin according to the method provided in Ubide et al. (2015) using photon machines 262 G2 193 nm UV laser with a Helex two-volume cell coupled to a Thermo iCAPQ ICP-MS. Laser beam spot for 263 maps was 12 µm square, scan speed 8 µm/s and repetition rate 41 Hz. NIST SRM 610 glass was used as 264 standard. A beam size of 12 μ m square typically allows a spatial resolution of 7-10 μ m in x and y. This relatively 265 new ablation technique removes only 3-4 μ m of material, which allows several maps to be performed on the 266 same area. A first map with all REE and other HFSE was first established, before acquiring a second map with 267 only 5 elements in order to obtain a better signal. LA-ICP-MS maps provide detailed spatial information on 268 zonation in crystals that would not be available otherwise. Analysis were semi-quantitative and compared to 269 microprobe data to obtain the real concentration.

270 To check for the absence of REE mineral inclusions in the pyroxene, we analyzed one sample from the 271 Amis complex by Scanning Transmission Electron Microscopy (STEM) at the Centre Raymond Castaing in 272 Toulouse. Unlike TEM, with STEM the electron beam is focused on a specific point of the sample and scans its 273 surface instead of being focused on the entire surface. This allows a better resolution as well as local chemical 274 identification (e.g. Pennycook and Nellist 2011). The Focused Ion Beam (FIB) was used to prepare the sample 275 for STEM. The FIB is a Hélios 600i operated at 5 kV, and the TEM a JEOL cold-FEG JEM-ARM200F operated 276 at 200 kV equipped with a Cs probe corrector reaching a spatial resolution of 0.078 nm. Images were acquired in 277 high-angle dark field (HAADF) as well as bright field (BF) modes. EDX spectra were recorded on a 278 JEOL CENTURIO SDD detector.

279

280

4. Amphibole and pyroxene textural and chemical description

281 *4.1. <u>Textural features</u>*

4.1.1. Amphiboles

283 Amphiboles occur in granites and pegmatites as well-developed euhedral crystals (Fig 4e). No significant 284 textural difference was found between amphiboles of the six complexes. They all appear pleochroic, dark green 285 or dark blue to black under plain polarized light (PPL). Their size is highly variable from one complex to another 286 as well as inside pegmatites (typically ranging from 20 µm to 10 cm), but is always comparable with the size of 287 quartz and alkali feldspar crystals in the same rock. Except at Evisa (Fig 4j), amphiboles are not zoned, but they 288 display dissolution/corrosion textures on their rims. They commonly have small quartz and feldspar inclusions, 289 and are in equilibrium with these minerals. Replacement of amphiboles by aegirine and Ti- and/or Fe-oxides is a 290 common feature of peralkaline granitoids (Marks and Markl 2017) and occurs in all pegmatites and some 291 granites (Fig 4e).

292

293 *4.1.2. Pyroxene*

294 Pyroxene is ubiquitous in pegmatites and granites of all six complexes but present three different 295 textural features. (1) In pegmatites which contain only minor amphibole, it occurs as light to dark green euhedral 296 crystals under PPL ranging from 50 µm to 10 cm. These pyroxene crystals present a core-to-rim zonation 297 pattern, with additional sector zoning in the rims, which are visible under PPL (Fig 4f). Similarly to amphiboles, 298 these crystals are in equilibrium with quartz and alkali feldspar grains. In the rims of this pyroxene, we also 299 found fluid inclusions (Fig 4g) and mineral inclusions that match the composition of those met in pseudomorphs 300 (Fig 4b and h). These pyroxene crystals were found in all studied complexes but Strange Lake. At Khan Bogd, 301 these crystals present an additional zoning area between the core and the sector zoned rims (Fig 4k). (2) In some 302 pegmatites, pyroxene occurs as anhedral crystals of size similar to the one we just described. These crystals are 303 oscillatory zoned, from yellowish to green under PPL (Fig 4i). They commonly include Fe- and/or Ti- oxides. 304 (3) In granites as well as in some pegmatites, anhedral pyroxene is found replacing amphibole (Fig 4e). This 305 pyroxene is also weakly oscillatory zoned. As it replaces amphibole only partially, it's size is smaller than the 306 previously described pyroxene crystals and depends on the original amphibole size. This pyroxene is associated 307 with Fe- and/or Ti- oxides and can sometimes keep the 120° cleavages from the original amphibole.

308 Based on occurrence, size, color, zoning and mineralogical associations, we distinguish two main types of 309 pyroxene in the six complexes we focus on in this study. Type-I matches euhedral zoned pyroxene, and our 310 study focuses mainly on this type. Based on the observation that they present anhedral shape, oscillatory zoning, and are associated with Fe- and/or Ti- oxides, we regroup anhedral single pyroxene crystals and pyroxene
 replacing amphibole under the denomination type-II.

313

314 <u>4.2. Major element composition</u>

315 *4.2.1. Amphiboles*

316 Since they present the same textural features, all amphiboles were identified based solely on their 317 chemistry, following the classification of Hawthorne et al. (2012) and in accordance with Leake et al. (1997). The general formula of amphibole is $AB_2C_5T_8O_{22}W_2$ where, in this study, A = Na; B = Na, Ca; $C = Fe^{2+}$, Fe^{3+} , 318 319 Li; T = Ca, Al; and W = OH, F. The rootname is given to these amphiboles based on the composition of the sites 320 A, B and C, and the prefix is added according to the composition of the sites C and W. By strictly following 321 these rules, it appears that some complexes contain only one amphibole type, common to granites and 322 pegmatites, and some contain different amphiboles (Table 1). Ambohimirahavavy (ferro-ferri-fluoro-leakeite; Na₃Fe₄LiSi₈O₂₂F₂), Manongarivo and Strange Lake (fluoro-arfvedsonite; Na₃Fe₅Si₈O₂₂F₂), and Khan Bogd 323 (arfvedsonite; Na₃Fe₅Si₈O₂₂(F,OH)₂) contain only one amphibole. At Amis, amphibole is ferro-ferri-fluoro-324 325 leakeite in pegmatites and fluoro-arfvedsonite in the granite. At Evisa, granites and pegmatites contain a zoned 326 amphibole with a core of fluoro-arfvedsonite and a rim of ferro-ferri-fluoro-leakeite (Fig 4L). In addition, the 327 hypersolvus granite from Evisa contains zoned crystals with ferro-ferri-katophorite (Na₂Ca(Fe²⁺,Mg)₄Fe³⁺(Si₇Al)O₂₂(OH)₂) in their core and fluoro-arfvedsonite in their rims (Fig 4j). 328

329 However, it is important to keep in mind that amphibole is a solid solution, thereby two amphiboles can 330 have different names but be close in composition, and inversely can have the same name but present a 331 compositional range. To avoid these artificial hiatuses and subsets, we focused on the concentration of elements 332 involved, when attributing amphibole names in this study. Hence, the distinction between katophorite and other 333 amphiboles is based on the Ca/(Ca+Na) ratio; being >0.25 (close to 0.4), this makes it a Na-Ca amphibole, 334 compared to values <0.1 for other amphiboles, which are then considered Na-amphiboles. Thereby, ferro-ferri-335 katophorite at Evisa exhibits a significant compositional gap with the other amphiboles from this study. The distinction between Na-amphiboles is more subtle, and is based on the amount of Li and Fe²⁺/Fe³⁺ ratio in the C 336 337 site. Ideally, leakeite has 1 Li apfu; this value is on average 0.6 apfu at Ambohimirahavavy and Evisa, and 0.7 338 apfu at Amis. Arfvedsonites from the other complexes have average Li contents of 0.1-0.3 apfu, validating the 339 distinction proposed here, although these values are intermediate between the two categories. The prefixes ferroand ferri- refer to the presence of Fe^{2+} and Fe^{3+} , respectively, in the C site. The prefix fluoro- indicates that the F concentration is systematically higher than the concentration of other elements in the W site. In this study, the amounts of F and OH, respectively, are close to 1.1 and 0.9 apfu in arfvedsonite from all complexes but Khan Bogd, where it is closer to 0.9 apfu F and 1.1 apfu OH. Strictly complying with the definition would result in two different names (F and OH varieties), which would vary with each complex. However, this would lead to add unnecessary confusion to the nomenclature, therefore, because the compositional differences are only minor, we chose to ignore this parameter.

347 Within a given category, amphiboles are globally similar in composition in all complexes, including Fe²⁺/Fe³⁺ (around 3 for fluoro-arfvedsonite, 2 for leakeite, and 3.5 for katophorite), with only local variations 348 349 (Table 1). Chlorine was not detected by microprobe analyses in any of the amphiboles. The main distinction 350 regarding major element composition is found in (fluoro)-arfvedsonite from Manongarivo and Khan Bogd, 351 which both have high contents of Mn (respectively 1.92 and 1.66 % vs 0.50 % in the other complexes, Fig 5d) 352 and Fe²⁺/Fe³⁺ (4.00 and 5.90 vs 2.40). Amphibole from Ambohimirahavavy is also rich in Mn (1.99 %) but does not show a high Fe^{2+}/Fe^{3+} value. Fluoro-arfvedsonite from Amis is as rich in F as ferro-ferri-fluoro-leakeite from 353 354 all other complexes (around 1.6 apfu), which is unusual (Deer et al. 1997b). Amphiboles from pegmatites are generally slightly richer in HFSE, Li and Fe³⁺ than amphiboles from granites. 355

356

357 *4.2.2. Pyroxene*

Based on composition, pyroxene in granites and pegmatites consists of 85 to 97 % aegirine endmember. Therefore, according to the classification of Morimoto (1988), we will refer to it as aegirine in the following sections of this paper. Pyroxene from Ambohimirahavavy and Khan Bogd is more Ca-rich than the other complexes, but it is still classified as aegirine.

Type-I aegirine crystals show a characteristic zoning pattern in pegmatites from all locations: a core-torim zoning visible in BSE images (Fig 6n) and a sector zoning visible even by optical microscopy (Fig 6o). This pattern defines 3 main zones, where zone A is the core and zones C and D form sector-zoned rims. Zone C matches the crystallographic sector (110), and zone D sector (100) (Ubide et al. 2019). In a few cases, an additional zone matching the crystallographic sector (010) is also present (e.g. ESM 6). All zones are optically distinguished by different tones of green, attributed to different Ti contents (Fig 6g) (e.g. Strong 1969; Ferguson 1973; Nielsen 1979). The contacts between different zoning are sharp in all occurrences (Fig 6). Zone A, the 369 core, is systematically enriched in Ca, Sn, Hf, Zr (Fig 7a, b, c, g) and depleted in Na, Al and Fe³⁺ compared to 370 the other zones (Table 3). In pegmatites, we measured up to 2 wt% ZrO_2 in aegirine cores; although uncommon, 371 Zr-rich aegirine was also described in other localities such as in nepheline syenites from the Motzfeldt Centre, 372 South Greenland (Jones and Peckett 1981; up to 7 wt% ZrO₂) and in metaluminous trachytes from the 373 Warrumbungle Volcano, Australia (Duggan 1988; up to 14.5 wt% ZrO₂). Sector zone C is particularly enriched 374 in Ti and Ca (Fig 6e, g), while D is in Fe (Fig 6c). Two major growth layers are visible in the rims with, in sector 375 C, Ti and Ca decreasing towards the rims of each layer (Fig 6e, g). Al content is low in aegirine from all six 376 complexes (about 0.3 % Al₂O₃ compared to standard average values of 1.2 % reported in aegirine; e.g. Deer et 377 al. 1997a). At Khan Bogd, an additional zone, referred to as zone B, is found between the core and the sector-378 zoned rims of the crystals (Fig 4k); it shows random oscillations due to variations in many elements, including Ti 379 and Mn. The same zoning pattern, including zone B, was also observed by Ranløv and Dymek (1991) in 380 nepheline syenite from the Narssaq Peninsula. However, they only documented variations in Zr, Ti, Al, Na, Ca 381 and Fe. Larsen (1976) and Piilonen et al. (1998) respectively described a core-to-rim zoning in aegirine from 382 nepheline syenite in the Ilímaussaq and Mont Saint-Hilaire complexes. They report cores enriched in Ca, Fe²⁺, Mg, Mn and Zr compared to rims, which in turn contain higher Al, Ti, Na and Fe³⁺. Shearer and Larsen (1994) 383 384 supplemented the description of aegirine from Ilímaussaq, by reporting Sr and REE enrichment in cores 385 compared to rims. In our analyses, we did not observe a systematic difference in Mn and Sr contents between 386 core and rims.

387 Type-II aegirine is not sector-zoned but shows oscillatory zoning and we observed a wide variety of 388 compositions depending on which layer was analyzed (Table 2). This zoning is irregular, with no core, made of 389 compositionally varying layers with variable thickness. No common pattern was found among any type-II 390 aegirine crystals, beside their heterogeneity. Many elements are involved, including Fe, Mn, Ca, and Ti (see 391 Online Resource 8). No significant chemical differences were found between type-II aegirine from granites and 392 pegmatites. Type-II aegirine is commonly depleted in Ti and Sn compared to type-I aegirine, except at Khan 393 Bogd. Mg is commonly found in higher concentration in type-II aegirine (Table 2). No significant compositional 394 difference was found between type-II aegirine replacing amphibole in granite and those in pegmatites. At 395 Strange Lake, all observed aegirine crystals are of type-II (Table 2; Salvi and Williams-Jones 1990; Roelofsen 396 1997).

397

399 *4.3.1. Amphiboles*

400 Amphiboles from all six complexes have very similar trace element patterns (Table 1). Fluoro-401 arfvedsonite and ferri-fluoro-leakeite have practically the same composition, with Li contents alone being 402 responsible for the transition from one kind to the other. This cation enters the amphibole structure according to the substitution: ${}^{C}Li^{+} + {}^{C}Fe^{3+} = 2{}^{C}Fe^{2+}$, described by Hawthorne et al. (1996). Ferro-ferri-katophorite, which is 403 more calcic than the two other amphiboles, relates to fluoro-arfvedsonite through the substitution: ${}^{B}Ca^{2+} + {}^{T}Al^{3+}$ 404 = ^BNa⁺ + ^TSi⁴⁺ (Deer et al. 1997b). This last substitution increases the size of the C site, leaving more space to 405 incorporate LREE. Similarly, the coupled substitution (Mg, Fe²⁺) \rightarrow Fe³⁺ occurring in the C site of amphiboles 406 407 enhances HREE incorporation (Bottazzi et al. 1999; Siegel et al. 2017b). Amphibole chondrite-normalized REE 408 patterns are systematically enriched in HREE relative to LREE, with a depression in medium REE (MREE, from 409 Eu to Dy). This pattern is unusual compared to more Ca-rich amphibole compositions in which REE patterns are 410 usually enriched in LREE or in MREE (e.g. Marks et al. 2004; Coint et al. 2013). All spectra display a negative 411 Eu anomaly suggesting early plagioclase fractionation (Fig 8). The absolute REE concentrations, though 412 somewhat varying from one complex to another, are quite similar (Fig 8).

413 Despite their similar patterns, the absolute concentrations of REE and other HFSE can vary in 414 amphiboles from one complex to the other. Fluoro-arfvedsonite from Manongarivo and Khan Bogd both have 415 low quantities of Sn (4 and 10 ppm) and HREE (35 and 50 ppm on average). Fluoro-arfvedsonite from 416 Manongarivo also has less LREE than fluoro-arfvedsonite from other complexes (11 vs 60 ppm on average). 417 Ferro-ferri-katophorite from Evisa displays high enrichments in LREE and HREE (respectively 224 and 290 418 ppm) (Fig 8). Ferro-ferri-fluoro-leakeite from Amis is notably different from other complexes in terms of trace 419 elements: it is richer than ferro-ferri-fluoro-leakeite from other complexes in Ti (0.18 vs 0.07 apfu), Sn (360 vs 420 32 ppm), Zn (0.24 vs 0.05 apfu), Pb (97 vs 7 ppm), LREE (168 vs 37 ppm on average) and HREE (140 vs 83 421 ppm on average) (Table 1, Fig 5d). Mn and Zn vary together on a 1:1 relation at Strange Lake, as mentioned in 422 the paper by Hawthorne et al. (2001). Mn and Zn covary in amphiboles from Ambohimirahavavy and Amis as 423 well, but in a different way (respectively 4:1 and 1:3), while at Manongarivo and Khan Bogd they do not appear 424 to be related.

425

427 Type-I aegirine, similarly to amphiboles, presents comparable trace elements patterns but different 428 absolute concentrations, from one complex to the other (Table 2). Cores of type-I aegirine from the two 429 Malagasy complexes have high Zr concentrations compared to other complexes (> 8000 vs 2700 ppm on average 430 for the other complexes). Khan Bogd aegirine is poorer in Li (17 vs 115 ppm), LREE (12 vs 80 ppm on average) 431 and HREE (21 vs 125 ppm on average). Type-I aegirine from Amis, similarly to ferro-ferri-fluoro-leakeite in this 432 complex, shows quite a different trace chemistry than aegirine from other complexes. Average Li is low (50 vs 433 115 ppm), while other metals are high, e.g. Zn (1780 vs 435 ppm), Sn (3740 vs 700 ppm), Pb (39 vs 7 ppm) and 434 U (29 vs 0.5 ppm) (Table 2 and Fig 7g). REE and Sc contents in cores of type-I aegirine (zone A) are higher than 435 in the rims (zones B, C and D) for all complexes (Fig 6, 9 and Table 3; similar maps of aegirine crystals from 436 each complex are provided in Online Resource 4 to 8). Inside the rims, sector zone C is systematically enriched 437 in REE compared to sector D (Fig 6k, 1, m). Nb and Ta do not have a constant behavior: their contents can be 438 higher in sector zone C, be present only in fractures, or be enriched in an independent zonation, with all three 439 cases being possible within the same sample. Similarly to amphiboles, the chondrite-normalized REE patterns of 440 aegirine show high values for HREE relative to LREE, with lower medium REE. This particular pattern is 441 known to be specific to Na- and Fe-enriched aegirine (Mahood and Stimac 1990; Shearer and Larsen 1994). The 442 absolute REE concentrations, are quite similar from one complex to another, although small variations do exist 443 (Fig 9).

Trace element concentrations in type-II aegirine are highly dependent on the zoning pattern. This is notably true for Sn, Hf and Zr. Compared to type-I aegirine, Sn covaries with Ti instead of Zr. The shape of type-II aegirine REE patterns is similar to that of type-I, i.e. high HREE relative to LREE, with a depression in medium REE and a negative Eu anomaly. There is no common pattern in the REE contents of type-II aegirine: REE contents are higher in type-II than in type-I aegirine at Khan Bogd, generally lower at Evisa, and have similar values in Madagascar and at Amis (Fig 7g).

To verify that the concentrations of REE measured in aegirine truly result from their incorporation in the crystal structure and not from small mineral inclusions, we used a STEM approach to investigate the core of a type-I aegirine, where REE are most concentrated. We did not detect any REE mineral inclusion, validating the premise that the REE are indeed incorporated in the pyroxene's structure. The main substitution involved in this process is common to type-I aegirine from all pegmatites. The classic aegirine substitution is

455
$${}^{M2}Ca^{2+} + {}^{M1}(Mg^{2+}, Fe^{2+}) \leftrightarrow {}^{M2}Na^{+} + {}^{M1}Fe^{3+}.$$
 (Eq. 1)

456 However, in aegirine from this study there is not enough Fe^{3+} to equilibrate this reaction, hence Al, Zr and Ti, 457 present in aegirine, also contribute to compensate this deficiency, resulting in a new substitution mechanism:

458
$${}^{M2}Ca^{2+} + {}^{M1}(Mg^{2+}, Fe^{2+}) \leftrightarrow {}^{M2}Na^{+} + {}^{M1}(Fe^{3+}, Al^{3+}, Zr^{4+}, Ti^{4+}).$$
 (Eq. 2)

However, REE can enter both M1 and M2 sites along with Fe³⁺ and Na⁺ (Beard et al. 2019). In clinopyroxenes, 459 incorporation of REE, as well as other HFSE, is controlled by five crystal-chemistry driven mechanisms, with 460 461 site parameters being determined by the size of the major elements. The first mechanism is decreasing 462 temperature and increasing melt alkalinity, which usually lowers the REE compatibility. The second mechanism is similar to that for amphiboles and is defined by the substitution $Ca \rightarrow Na$, which increases the size of the M2 463 464 site leaving more space to incorporate LREE. It also decreases the size of the M1 site, making it more adequate 465 to incorporate HREE. The third mechanism takes place in the M1 crystallographic site, where the combined substitution (Mg, Fe²⁺) \rightarrow Fe³⁺ leads to a charge effects that allows more HREE to enter this site along with Fe³⁺ 466 467 (Marks et al. 2004; Beard et al. 2019). The fourth mechanism, described by Mollo et al. (2017), is also due to a 468 charge effect. These authors propose the coupled substitution

469
$$^{M1}(Al, Fe^{3+}) + ^{T}Al \leftrightarrow ^{M2}(Mg, Fe^{2+}) + ^{T}Si,$$
 (Eq. 3)

whereby the increase in Al at the expense of Si creates a charge deficiency compensated by the incorporation of
REE in the M2 site. In addition, the growth from an environment rich in REE can result in incorporation of high
levels of REE in aegirine. Indeed, the adsorption of REE onto aegirine surface during supersaturation may be too
fast to equilibrate as crystal growth rate exceeds the internal diffusion rate (Smith et al. 2004).

All of the above mechanisms have the potential of favoring the incorporation of REE into pyroxene, however, aegirine in our study having low ^TAl and practically no Mg or Fe²⁺ in the M2 site, the mechanism of Mollo et al. is probably not significant in this case. The high REE concentration in aegirine of this study is therefore probably the result of the four other mechanisms that rely on size, charge, and kinetics effect at M sites of aegirine.

479

481 Some amphibole and aegirine chondrite-normalized spectra, mostly from one pegmatite sample from
482 Amis, display a peak in Ce concentration (Fig 8 and 9). The occurrence of this peak being restricted to a few
483 samples, mass interference is excluded. In addition, ICP-MS signals showing peaks during ablation were all

excluded so the presence of this Ce peak is not due to mineral inclusions. This peak may be due to the presence of Ce⁴⁺ instead of the usual Ce³⁺. Indeed, Ce⁴⁺ has a smaller ionic radius (0.97 Å) than Ce³⁺ (1.143 Å) for an eight-fold coordination, thereby Ce⁴⁺ substitutes more easily with Fe²⁺ (0.92 Å) and Mg²⁺ (0.89 Å; Shannon 1976). The change in oxidation state is commonly interpreted to indicate an increase in oxygen fugacity in the environment that can be either low temperature magmatic (<600 °C, Trail et al. 2012) or hydrothermal (Xu et al. 2017).

490

491 <u>5. Discussion</u>

492 *5.1. Origin of amphiboles*

493 In the previous sections we have seen that three different amphiboles can be distinguished based on 494 major and trace chemistry: fluoro-arfvedsonite (Na₃Fe₅Si₈O₂₂(F,OH)₂), ferri-fluoro-leakeite their 495 (Na₃Fe₄LiSi₈O₂₂F₂), and ferro-ferri-katophorite (Na₂Ca(Fe²⁺,Mg)₄Fe³⁺(Si₇Al)O₂₂(OH)₂). All of them occur in 496 granites and pegmatites as euhedral crystals and contain inclusions of quartz and feldspar grains (Fig 4e), 497 suggesting co-crystallization. Only amphiboles from Evisa have a variable composition, from fluoro-498 arfvedsonite or ferro-ferri-katophorite in cores, respectively to ferri-fluoro-leakeite or fluoro-arfvedsonite in the 499 rims. In the other complexes, only one amphibole type is present in granites and in pegmatites. Based on textural 500 observations, we believe these changes are the sign of magmatic evolution. Indeed, the concentration of Li and 501 Na rises in the melt as differentiation occurs, until Li- and Na-rich amphiboles crystallize instead of their Li-502 poor, Mg-, Ca-rich equivalents. Based on various pieces of evidence, such as having a similar age to that of 503 granite emplacement (Kovalenko et al. 2006), presence of melt inclusions (Schmitt et al. 2002), occurrence as 504 well-developed euhedral crystals, and dissolution/corrosion textures (Estrade et al. 2014a; Gysi et al. 2016), it is 505 accepted in the literature that amphiboles are magmatic phases in peralkaline granites. Hence, in accordance with 506 our observations and those in the literature about the six studied complexes, we conclude that amphiboles of all 507 types in our samples are magmatic. Their variation in trace element composition is directly linked to the 508 environment they grew in, a highly evolved melt, enriched in incompatible elements. Among incompatible 509 elements are the REE, which are known to be preferentially incorporated in alkaline amphiboles along with Na for LREE and Fe³⁺ for HREE (Bottazzi et al. 1999; Siegel et al. 2017b). Due to this crystallo-chemical control, 510 511 amphiboles can incorporate a lot of REE at the magmatic stage, with a preference for HREE. Slight 512 compositional differences in REE, Zn, Sn, Nb, Zr and Pb between amphiboles of the 6 complexes (Fig 5d) likely reflect parental melt compositions. For example, the presence of ferro-ferri-katophorite in Evisa probablyindicates an effect of melt contamination in Ca by the surrounding aluminous granites.

515 As previously described, amphiboles are commonly altered to Fe- and Ti-oxides and to type-II aegirine. 516 Based on textural evidence that at Strange Lake this replacement occurs preferentially along cleavages in altered 517 granites and pegmatites, as well as on mass-balance calculations, Salvi and Williams Jones (1990) identified this 518 pyroxene occurrence as hydrothermal. Because we observed this replacement in all of our samples, it follows 519 that a late hydrothermal event is a common feature in the complexes studied and, by extrapolation, most likely in 520 alkaline rocks in general; indeed, a hydrothermal event was also documented in silica-undersaturated alkaline 521 rocks such as at Ilímaussaq (e.g. Borst et al. 2016) or at Khibina and Lovozero (e.g. Arzamastsev et al. 2011). 522 Gysi et al. (2016) and Vasyukova and Williams-Jones (2019) show that at Strange Lake this event is due to a 523 relatively high temperature, Ca-rich fluid. Based on the similarity of amphiboles alteration in all six complexes, 524 we infer that the circulation of an orthomagmatic fluid is a process common to all complexes.

525

526

527

5.2. <u>Origin of pyroxene</u>

5.2.1. Core-to-rim zonation

Even though both types of aegirine identified in all complexes show some kind of zonation, only type-I shows a core-to-rim zonation, with a core enriched in Ca, Zr, Sn, Hf, Sc, REE and depleted in Na, Al and Fe compared to the rims (Fig 6). The rims also have a higher Fe^{3+}/Fe^{2+} ratio than the core. This pattern is systematic for type-I aegirine crystals from all complexes. Type-II aegirine is oscillatory zoned and can replace amphibole.

533 The sharp compositional change between the core and the rims in type-I aegirine (Fig 6) suggests a 534 sudden change in the crystallizing environment. Based on the scarce presence of type-I aegirine where 535 amphiboles are present, we infer that they grew in competition. Only the rims C and D contain fine oscillatory 536 zoning (e.g. TiO_2 on Fig 6g), fluid inclusions (Fig 4g), and, at Ambohimirahavavy, mineral inclusions that match 537 the composition of those measured in hydrothermal pseudomorphs (e.g. zircon, bastnäsite-(Ce), Fig 4b and h). 538 Therefore, we infer that type-I aegirine has a magmatic core (zone A on Fig 6p) while the rims are hydrothermal 539 (zones C and D in Fig 6p). Zone B being only present at Khan Bogd, it will be discussed lower in this section. 540 Chemical zoning in aegirine of our rocks matches that observed by Piilonen et al. (1998) in aegirine from Mont 541 Saint-Hilaire, which they describe as a magmatic pyroxene overgrown by hydrothermal fibrous aegirine. More
542 details about each element in the different zones is provided in the following paragraphs.

543 The variations in Na, Ca, Al, and Fe can be explained by a common process. Type-I aegirine magmatic cores are enriched in Ca, and its hydrothermal rims are richer in Na, Al and Fe³⁺. Evidence such as widespread 544 545 feldspar albitization, fluid inclusions composition, and the hydrothermal growth of aegirine without amphibole 546 indicate that circulating hydrothermal fluids are oxidizing (above the magnetite-hematite buffer), of relatively 547 high pH and rich in NaCl among other phases (Salvi and Williams-Jones 1996; Smith 2007; Estrade 2014; Gysi 548 et al. 2016). Depletion in Ca and enrichment in Na and Fe while moving on from aegirine magmatic core to 549 hydrothermal rims can then be explained by intake of Na from the hydrothermal fluids and simultaneous 550 crystallization of Ca-rich secondary minerals in the pseudomorphs (fluorite, bastnäsite-(Ce)). This process 551 increases the amount of aegirine endmember compared to hedenbergite endmember which, along with the hydrothermal environment being more oxidizing than the silicate melt, increases the Fe^{3+}/Fe^{2+} ratio in the rims. 552 553 The absolute amount of Al is very low in aegirine of this study, so the observed increase in this element in aegirine rims can be linked only to charge effects, with Al³⁺ being incorporated along with Fe³⁺. 554

555 Because of their similar behavior, Zr and Hf both occur in the same growth zones. Based on the many 556 arguments in favor of a magmatic core and hydrothermal rims, we propose two hypotheses as to the 557 crystallization timing of type-I aegirine. Jones and Peckett (1981) asserted that formation of Zr-enriched aegirine 558 is enhanced by low oxygen fugacity, and can only occur if no other Zr-bearing phase is growing. Pseudomorphs 559 after primary zirconosilicates were found around aegirine crystals (Fig 4b), hence it is impossible that aegirine 560 grew after the magmatic zirconosilicates. Our first hypothesis is that, in accordance with the observations made 561 by Jones and Peckett (1981), type-I aegirine cores grew before magmatic zirconosilicates, in the magma 562 chamber. In the absence of any mineral Zr has a strong affinity for, Zr partitioned mostly into aegirine. These 563 primary phenocrysts were brought into pegmatites, where primary zirconosilicates began to crystallize, 564 incorporating most of the available Zr.. Moving on to the hydrothermal stage, primary zirconosilicates were 565 destabilized and replaced by minerals richer in Zr, including zircon, which form the pseudomorphs. Aegirine 566 rims were then in competition with zircon, which has a high Zr uptake and therefore accounts for the decrease in 567 Zr concentration in aegirine rims. In this hypothesis, aegirine necessarily stopped growing during their migration 568 from the magma chamber to the pegmatites, or it would have resulted in another zonation in the magmatic core. 569 Our second hypothesis is that aegirine core and primary zirconosilicates grew at the same time. In this case the 570 conclusions of Jones and Peckett (1981) do not apply. This may be because their study was based on rocks from 571 the magmatic chamber of Ilímaussaq which had plenty of time to equilibrate during crystallization, whereas our 572 study is based on pegmatites, which crystallize quickly. Anyhow, in this hypothesis the competition between the 573 two minerals resulted in the preference of Zr to go into primary zirconosilicates, but because of the high content 574 of Zr in the melt and its low mobility (Duggan 1988), affinity for aegirine was still high enough to enrich it up to 575 the 2 wt% that we have measured. The subsequent hydrothermal stage is similar to the one in our previous 576 hypothesis, with higher Zr uptake by zircon rather than by zirconosilicates accounting for the decrease in Zr 577 concentration in aegirine rims. A similar behavior was observed for Sn and is most likely due to a similar phenomenon. No Sn-based mineral was found in pseudomorphs, but Sn⁴⁺ can easily enter in the composition of 578 secondary Ti-oxides because its radius is similar to the Ti⁴⁺ ion in eight-fold coordination (Shannon 1976). In 579 580 addition, Sn is known to have a high solubility in Cl-rich fluids (e.g. Keppler and Wyllie 1991), making it easily 581 mobilized. This results in a much higher Sn content in the magmatic zone A than in the rims of aegirine. At 582 Amis, amphibole in pegmatite as well as aegirine are both enriched in Sn, Zn and Pb in comparison with the 583 other complexes of this study (Fig 7g). Ambohimirahavavy and Manongarivo have similar trace elements 584 concentration (Fig 7g) and formed in the same geological context, a few km away from each other. Both 585 amphibole and pyroxene from Khan Bogd have low trace elements concentrations. All these observations tend to 586 indicate that the variations of absolute concentration in REE, Zn, Sn, Zr, U and Pb between aegirine crystals of 587 the 6 complexes (Fig 7g), similarly to amphiboles, likely reflect parent melt compositions.

588

589

5.2.2. Sector zoning

590 Sector zoning occurs in the most outer rims in type-I aegirine, involves variations in Ti, Ca, Fe and 591 REE, and was observed in all complexes. Many explanations have been invoked as to the origin of sector zoning, 592 and the latest agree on the fact that it is linked to a crystal growth faster than element diffusion rates (Ubide et al. 593 2019). If this condition is respected, it will result a disequilibrium between the different crystal faces that will not 594 all incorporate the same elements as it normally would (Strong 1969; Ferguson 1973; Larsen 1981). Elements 595 are however incorporated on the different crystal faces in a matter so that charge balance is respected (Ubide et 596 al. 2019). The crystallization rate could also impact the size of protosites (partially formed site on the surface of 597 a growing crystal), which are in equilibrium with the crystallizing environment, and preferentially incorporate 598 elements depending on their charge/radius ratio. If the growth rate is higher than ionic diffusion in the crystal, 599 the protosites will not have enough time to re-equilibrate in size and incorporate different elements on the

different faces of the crystal (Nakamura 1973). This higher growth rate is likely related to a lower temperature
due to the change from magmatic to hydrothermal environment (Barkov and Martin 2015). However, these
considerations must be taken carefully, as most of the research is based on augite and not aegirine.

603 Incorporation of Ti in aegirine is uncommon (Rønsbo et al. 1977; Dyulgerov and Platevoet 2006). Many 604 explanations have been evoked: high temperature, low fO₂, high fO₂, low pressure, high activities of TiO₂ and 605 SiO₂, high activity of alkalis, or a peculiar chemical composition of the environment such as low Zr and Na and 606 high Ca (Ferguson 1973; Flower 1974; Larsen 1976; Rønsbo et al. 1977; Nielsen 1979; Brousse and Rançon 607 1984; Dyulgerov and Platevoet 2006). In our samples, a high temperature is ruled out as Ti incorporation occurs 608 during the hydrothermal stage. As described in the previous section, Zr and Ca concentration are low in the 609 crystallizing environment, and Na is high. Hence, and in accordance with Dyulgerov and Platevoet (2006), 610 Larsen (1976), and Flower (1974), we suggest that in our rocks the incorporation of Ti into hydrothermal 611 aegirine is favored by a high activity of Na and the low availability of Zr. Titanium may have been provided by 612 the alteration of magmatic Ti-bearing minerals, such as astrophyllite (found at Manongarivo, Strange Lake, Khan Bogd and Amis, Fig 4a) chevkinite-(Ce) (found at Ambohimirahavavy, Manongarivo and Evisa), or others such 613 614 as narsarsukite, låvenite, and aenigmatite.

615

616

5.2.3. Oscillatory zoning

617 Oscillatory zoning is present in both types of aegirine. In type-I, it is found in the sector zoned rims C 618 (crystallographic sector (110)) and D (crystallographic sector (100)) as well as in zone B, intermediate between 619 the core and the rims. Considering that zone B in type-I aegirine is only present in our samples in Khan Bogd, 620 and was locally described in aegirine from the South Gardar Province, South Greenland by Ranløv and Dymek 621 (1991), a general process to explain its formation is difficult to establish. Like zones C and D, zone B is poorer 622 in Ca, Zr, Hf, Sn and REE than zone A. It is not specifically enriched in any element, but shows oscillatory 623 zoning in Ti and Mn. We believe that zone B development is related to a specificity in the history of the Khan 624 Bogd complex (Kynicky et al. 2011). In this complex, magmatic zone A aegirine would crystallize before or in 625 competition with primary elpidite. At the hydrothermal stage, primary elpidite altered mostly into a secondary 626 hydrated Ca- and REE-rich elpidite (Fig 4i), and zone B grew simultaneously. The uptake of Ca and Zr by 627 secondary elpidite from the fluid led to the lower concentration of these elements observed in zone B. The final

stage, similarly to other complexes, was a more or less extensive pseudomorphism of elpidite and the formationof zones C and D on aegirine.

630 Oscillating concentrations in zones C and D are most likely produced by autocatalytic surface 631 attachment, or boundary layer effect (Ortoleva et al. 1987; London 2008), which occurs when the crystal growth 632 rate exceeds the diffusion rate of components in the fluid. Consider an initial solution saturated in component A, 633 whose adsorption on the crystal surface is fast. As the adsorption progresses, the interface between the crystal 634 and the solution becomes depleted in component A, and switches to saturation in component B. Component B is 635 then adsorbed at the surface of the crystal, until the interface crystal-solution becomes saturated in component A 636 again. This process results in a rhythmic but irregular zoning of aegirine crystals in zones C and D, consistent 637 with a growth in a rapidly changing environment, which is a hydrothermal fluid.

Type-II aegirine either replaces amphibole or forms isolated crystals. Both kinds are found associated with Fe- and/or Ti- oxides, and are randomly zoned in many elements (Ti, Ca, Mn, Fe, Hf, Sn, Zr at least). The formation of aegirine from amphibole is known to be hydrothermal (e.g. Salvi and Williams-Jones 1990; Gysi et al. 2016). For arfvedsonite, it follows the reaction provided by Salvi and Williams-Jones (1997):

$$642 \qquad 3Na_3Fe_5Si_8O_{22}(F,OH)_2 + 2H_2O = 9NaFeSi_2O_6 + 2Fe_3O_4 + 6SiO_2 + 5H_2. \tag{Eq. 4}$$

The released Fe generally forms hematite or magnetite. Considering that zoning patterns are the same for all type-II aegirine crystals and that they are systematically found associated with Fe- and/or Ti- oxides, we consider that they all grew replacing amphibole, totally or partially. Hence, we consider type-II aegirine as hydrothermal.

647

648

5.3. REE behavior in amphiboles and aegirine

As mentioned in the introduction, amphiboles and pyroxenes, in agreement with their crystal-chemical parameters, can incorporate up to several hundreds of ppm of REE. In addition, their alkaline nature enhances the fractionation of light versus heavy REE through the preferential incorporation of HREE in their structure, resulting in the globally HREE-enriched patterns observed (Fig. 9; Beard et al. 2019). From the amphibole compositions, we know that the melt was rich in F (Table 1). Beard et al. (2020) showed that the more F in the melt, the lower the pyroxene-melt partition coefficient, hence the less REE are incorporated in pyroxene; this affects the LREE more than the HREE and hence contributes to the observed globally HREE-enriched patterns. 656 The REE compositional patterns observed in type-I aggirine are similar to the compositional trends for 657 Zr, Hf and Sn, i.e. they are globally enriched in the core compared to the rims, except for Khan Bogd where the opposite is true (Fig 10, Table 3). Given that the REE concentrations in aegirine at Khan Bogd are very low (2-7 658 659 ppm for Ce, 4-15 ppm for Y), we believe that this reverse trend is actually not significant (Fig 10). The 660 difference in REE concentration between aegirine zone A and zones B, C and D is due to the competition 661 between aegirine and zirconosilicates for these elements, during the magmatic stage for zone A, and during the 662 hydrothermal stage for the zones B, C and D. Scandium is sometimes grouped with REE, hence, it is not 663 surprising to observe that, similarly to REE, it is enriched in the core of type-I aegirine.

664 During the hydrothermal stage, aegirine is in competition with an altered, REE-enriched elpidite for 665 zone B, and with REE-bearing minerals in pseudomorphs (e.g. bastnäsite-(Ce)) for zones C - (110) and D -666 (100). We therefore suggest that it is a competition for REE between aggirine and coprecipitating minerals 667 having each a different REE affinity which explains the variations in REE concentrations among the aegirine 668 core-to-rim zones. REE are also affected by sector zoning and are more concentrated in zone C than D, along 669 with Ti and Ca. Beard et al. (2019) studied experimentally the fractionation of REE in clinopyroxene in 670 peralkaline melts and showed that the more Na and less Ca in M2 site of pyroxene, the smaller the M1 site, and 671 the more HREE are incorporated. In addition, the authors mention that physico-chemical parameters not 672 recorded in the composition of clinopyroxene, such as pressure, also play a significant role in the size of the M1 673 site. However, this mechanism is well defined only for pyroxenes whose aegirine component is not higher than 674 50 %. In our samples, the substitution Ca \rightarrow Na simply cannot be responsible for the zonation observed, as more 675 REE are found in the C sector, where Ca concentration is also higher. In addition, in Ca-rich pyroxenes such as diopside and augite, Ca was not reported to fractionate between sectors (e.g. Ubide et al. 2019), which is another 676 677 argument for the occurrence of a different exchange mechanism in our samples.

678 In addition to the above variations in bulk abundance, cores and rims also preferentially incorporate 679 REE with lower or higher atomic numbers, creating a fractionation between the heavy and light REE. To 680 evaluate this process, we calculated the percentages of REE enrichment or depletion from core to rim in type-I 681 pyroxene. To do so, data obtained by LA-ICPMS were assigned either to the core or to the rim, based on their Zr 682 content and zoning SEM observations. By comparing these two parameters, we were able to determine a 683 threshold value, unique to each complex, below which a given aegirine analysis should be considered as rim, and 684 above which core. These data were averaged out to obtain a single value each for core and rim (Fig 10). Finally, 685 we calculated the extent of loss or enrichment between core and rim for each REE, according to:

686
$$d = \frac{C-R}{C} \times 100$$
 (Eq. 5)

where *d* represents the percentage difference between core and rim, *C* and *R* the mean values obtained for core and rim, respectively. Error bars shown in Fig 11 were obtained using common error calculation formulae, namely (Eq. 6) for the mean and (Eq. 7) for the percentage.

690
$$\overline{err} = \sqrt{\frac{\sum_{x=1}^{n} (err_x)^2}{n}}$$
(Eq. 6)

691
$$d_{err} = \frac{C-R}{C} \sqrt{\frac{err_{C}^{2} - err_{R}^{2}}{(C-R)^{2}} + \frac{err_{C}^{2}}{C^{2}}}$$
 (Eq. 7)

with \overline{err} being the mean error, *n* the number of measurements, d_{err} the error on the percentage variation from core to rim, err_c the mean error on core data, and err_R the mean error on rim data. Resulting values indicate an enrichment in that element in the rim compared to the core when positive, while negative ones indicate a depletion (Fig 11).

From the calculation of the percentage of depletion of REE from core to rim (Fig 11) we see that Amis,
Manongarivo and Evisa have a trend with a more important depletion of LREE than HREE: from 40-80 % to 2040 % depletion, respectively. Ambohimirahavavy shows the opposite trend: around 30 % of LREE depletion for
45% of HREE depletion. The latter trend is not as obvious as for other complexes, but it is still significant.

700

701

5.4. <u>General model</u>

We propose a global model of formation for amphiboles and aegirine in pegmatites (Fig 12). During the magmatic stage, two hypotheses remain. In the first one, amphibole and zone A of type-I aegirine grew at early magmatic stage, concentrating Zr and REE. In the second hypothesis, type-I aegirine core, amphibole and zirconosilicates grew in competition. Most Zr was incorporated in the zirconosilicates, as well as REE if the primary zirconosilicates was EGM. However, highly differentiated alkaline melts being particularly enriched in Zr and REE, there is an extra supply in these metals which, combined with a fast crystallization, could be incorporated by the structures of amphiboles and aegirine (Fig 12a).

During the orthomagmatic hydrothermal stage, amphiboles did not grow anymore, and type-I aegirine was in competition with secondary Zr- and REE-bearing minerals (generally zircon and bastnäsite-(Ce)) that formed pseudomorphs after primary zirconosilicates. Since HFSE have much higher affinity for these secondary minerals, their concentrations globally decreased in the co-crystallizing aegirine rims. It is also during this stage
that type-II aegirine crystals nucleated. Nevertheless, competition does not explain the observed LREE-HREE
fractionation. Residual melt, depleted in incompatible elements, likely crystallized late quartz and feldspar.

715 Given the overwhelming evidence for circulation of an orthomagmatic fluid in all complexes, we propose 716 that fractionation results from variations in fluid composition. From the presence of secondary minerals such as 717 bastnäsite-(Ce) and fluorite, and the study of fluid inclusions at Ambohimirahavavy and Strange Lake (Estrade 718 2014; Vasyukova et al. 2016), we know that the fluids contained at least the anions Cl⁻, F⁻, and $CO_3^{2^-}$. These species have the ability to form stable complexes with the REE, to different extents. F⁻ and CO₃²⁻ together are 719 720 mostly depositional ligands (Migdisov et al. 2016), thus precipitate with REE typically forming the LREE-721 bearing bastnäsite-(Ce). On the other hand, Cl⁻ is a weak but abundant ligand that can transport LREE more 722 easily than the heavy ones (Migdisov et al. 2016). If present as a transporting ligand, F⁻ also transports more 723 LREE than HREE (Beard et al. 2020). We can thus infer that, depending on the relative concentrations of each 724 ligand, fractionation of light vs heavy REE would be specific to each complex (Fig 12b). As long as fluid flow 725 continued, aegirine rims kept growing, although REE incorporation and LREE-HREE fractionation kept 726 diminishing. By the time the hydrothermal stage waned, the two aegirine types were in equilibrium with the 727 pseudomorphs and the partially replaced amphiboles (Fig 12c).

728

729 6. Conclusions

730

By comparing amphibole and pyroxene in six alkaline complexes worldwide, our study highlights that despite different geodynamic contexts, mineralogy, and REE enrichment, there is a general crystallization process common to all studied alkaline pegmatites. We believe this process can be generalized to all alkaline SiO₂-saturated pegmatites worldwide, with local variations in mineralogy and REE-enrichment and fractionation rates. This crystallization process includes at least two main stages, respectively magmatic and hydrothermal.

From the study of the composition of amphibole and type-I aegirine core, we document the magmatic stage, during which a systematic primary enrichment in REE, HFSE and a LREE-HREE fractionation occurs. This enrichment is common to all complexes and linked to the properties of the extremely differentiated alkaline magma. The enrichment rate, however, differs from one complex to another. The REE fractionation is most represent an inherent properties of aegirine and alkali amphiboles that preferentially incorporate HREEand do not represent an inherent property of the melt.

742 From the study of the texture and composition of type-I aegirine rims and type-II aegirine, we document 743 similarities between complexes during the hydrothermal stage, at which amphiboles stop growing, type-I 744 aegirine grows sector-zoned rims, and type-II aegirine begins replacing amphibole along with Fe- and Ti-oxides. 745 The drop in REE concentration in hydrothermal aegirine of all complexes shows that REE hydrothermal 746 enrichment is globally less effective than magmatic REE enrichment, but still effective. The observed REE 747 fractionation in all complexes show that hydrothermal fluids also have the ability to fractionate LREE-HREE 748 through their transportation associated with ligands such as F⁻, Cl⁻. The rate of fractionation as well as the 749 amount of mobilized REE depends on the local properties of the circulating fluid(s). These observations show 750 that hydrothermal flow is mandatory in order to enhance the concentration of the REE, and specially HREE, in 751 alkaline pegmatites to ore levels.

752

753 <u>References</u>

- Ancey M, Bastenaire F, Tixier R (1978) Application des méthodes statistiques en microanalyse. Microanal
 Microsc Électronique À Balayage 323:11–16
- Arzamastsev AA, Arzamastseva LV, Zaraiskii GP (2011) Contact interaction of agpaitic magmas with basement
 gneisses: an example of the Khibina and Lovozero massifs. Petrology 19:109–133
- Barkov AY, Martin RF (2015) Anomalous Cr-rich zones in sector-zoned clinopyroxene macrocrysts in gabbro,
 Mont Royal, Montreal, Quebec, Canada. Can Mineral 53:895–910
- Beard C, van Hinsberg V, Stix J, Wilke M (2020) The effect of fluorine on clinopyroxene / melt trace-element partitioning. Contrib Mineral Petrol. https://doi.org/doi: 10.1007/s00410-020-1672-5
- Beard CD, van Hinsberg VJ, Stix J, Wilke M (2019) Clinopyroxene/melt trace element partitioning in sodic
 alkaline magmas. J Petrol. https://doi.org/10.1093/petrology/egz052
- Boily M, Williams-Jones AE (1994) The role of magmatic and hydrothermal processes in the chemical evolution
 of the Strange Lake plutonic complex, Quebec-Labrador. Contrib Mineral Petrol 118:33–47
- Bonin B (1988) Peralkaline granites in Corsica: some petrological and geochemical constraints. Rendiconti Della
 Soc Ital Mineral E Petrol 73:1191–1194
- Bonin B (2007) A-type granites and related rocks: evolution of a concept, problems and prospects. Lithos 97:1–
 29
- 770 Bonin B (1990) Les granites des complexes annulaires, BRGM. Office des publications universitaires
- Bonin B, Grelou-Orsini C, Vialette Y (1978) Age, origin and evolution of the anorogenic complex of Evisa (Corsica): A K-Li-Rb-Sr study. Contrib Mineral Petrol 65:425–432

- Bonin B, Platevoet B, Poitrasson F, Renna MR (2008) Eurogranites-IGCP510 2008 Joint Field-meeting—
 Alkaline The Permian–Triassic A-type Volcanic–Plutonic Igneous Suite of Corsica. In: 33th
 International Geological Congress in Oslo, Norway Convention Centre, Lillestrom, Norway
- Borst AM, Friis H, Andersen T, et al (2016) Zirconosilicates in the kakortokites of the Ilímaussaq complex,
 South Greenland: Implications for fluid evolution and high-field-strength and rare-earth element
 mineralization in agpaitic systems. Mineral Mag 80:5–30
- Bottazzi P, Tiepolo M, Vannucci R, et al (1999) Distinct site preferences for heavy and light REE in amphibole
 and the prediction of Amph/L D REE. Contrib Mineral Petrol 137:36–45
- 781 Brousse R, Rançon JP (1984) Crystallization trends of pyroxenes from agpaitic phonolites (Cantal, France).
 782 Mineral Mag 48:39–45
- 783 Chakhmouradian AR, Wall F (2012) Rare earth elements: minerals, mines, magnets (and more). Elements
 784 8:333-340
- 785 Chakhmouradian AR, Zaitsev AN (2012) Rare earth mineralization in igneous rocks: sources and processes.
 786 Elements 8:347–353
- 787 Chengyu W, Dianhao H, Zhongxun G (1990) REE Geochemistry in the Weathered Crust of Granites, Longnan
 788 Area, Jiangxi Province. Acta Geol Sin Engl Ed 3:193–209. https://doi.org/10.1111/j.1755 789 6724.1990.mp3002006.x
- Cocherie A, Rossi P, Fanning CM, Guerrot C (2005) Comparative use of TIMS and SHRIMP for U–Pb zircon dating of A-type granites and mafic tholeiitic layered complexes and dykes from the Corsican Batholith (France). Lithos 82:185–219
- Coint N, Barnes CG, Yoshinobu AS, et al (2013) Use of trace element abundances in augite and hornblende to
 determine the size, connectivity, timing, and evolution of magma batches in a tilted batholith.
 Geosphere 9:1747–1765
- Cucciniello C, Tucker RD, Jourdan F, et al (2016) The age and petrogenesis of alkaline magmatism in the
 Ampasindava Peninsula and Nosy Be archipelago, northern Madagascar | SpringerLink. Mineral Petrol
 110:309–331. https://doi.org/10.1007/s00710-015-0387-1
- 799 Currie KL (1985) An unusual peralkaline granite near lac Brisson, Quebec-Labrador. Curr Res 73–80
- Boer WA, Howie RA, Zussman J (1997a) Rock-Forming Minerals: Single-chain Silicates, Volume 2A.
 Geological Society of London
- B02 Deer WA, Howie RA, Zussman J (1997b) Rock-forming Minerals: Double-Chain Silicates, Volume 2B.
 B03 Geological Society of London
- Biehl M (1990) Geology, mineralogy, geochemistry and hydrothermal alteration of the Brandberg alkaline
 complex, Namibia. Geological Survey of Namibia
- 806 Donnot M (1963) Côte Nord-Ouest du complexe intrusif alcalin; Ampasindava-Manongarivo
- BO7 Droop GTR (1987) A general equation for estimating Fe 3+ concentrations in ferromagnesian silicates and oxides from microprobe analyses, using stoichiometric criteria. Mineral Mag 51:431–435
- Buggan MB (1988) Zirconium-rich sodic pyroxenes in felsic volcanics from the Warrumbungle Volcano,
 Central New South Wales, Australia. Mineral Mag 52:491–496
- Byulgerov MM, Platevoet B (2006) Unusual Ti and Zr aegirine-augite and potassic magnesio-arfvedsonite in the
 peralkaline potassic oversaturated Buhovo-Seslavtzi complex, Bulgaria. Eur J Mineral 18:127–138
- 813 Estrade G (2014) Le complexe cénozoïque alcalin d'Ambohimirahavavy à Madagascar : origine, évolution et
 814 minéralisations en métaux rares. Toulouse 3

- 815 Estrade G, Béziat D, Salvi S, et al (2014a) Unusual evolution of silica-under-and-oversaturated alkaline rocks in
 816 the Cenozoic Ambohimirahavavy Complex (Madagascar): Mineralogical and geochemical evidence.
 817 Lithos 206:361–383
- Estrade G, Salvi S, Béziat D, et al (2014b) REE and HFSE mineralization in peralkaline granites of the
 Ambohimirahavavy alkaline complex, Ampasindava peninsula, Madagascar. J Afr Earth Sci 94:141–
 155
- 821 Estrade G, Salvi S, Béziat D (2018) Crystallization and destabilization of eudialyte-group minerals in peralkaline
 822 granite and pegmatite: a case study from the Ambohimirahavavy complex, Madagascar. Mineral Mag
 823 82:375–399. https://doi.org/10.1180/minmag.2017.081.053
- European Commission (2018) Report on Critical Raw Materials and the Circular Economy.
 http://ec.europa.eu/docsroom/documents/27348. Accessed 25 Apr 2018
- Fedele L, Lustrino M, Melluso L, et al (2015) Trace-element partitioning between plagioclase, alkali feldspar,
 Ti-magnetite, biotite, apatite, and evolved potassic liquids from Campi Flegrei (Southern Italy). Am
 Mineral 100:233–249
- Ferguson AK (1973) On hour-glass sector zoning in clinopyroxene. Mineral Mag 39:321–325.
 https://doi.org/10.1180/minmag.1973.039.303.08
- Flower MF (1974) Phase relations of titan-acmite in the system Na2O-Fe2O3-Al2O3-TiO2-SiO2 at 1000 bars
 total water pressure. Am Mineral J Earth Planet Mater 59:536–548
- Foland KA, Landoll JD, Henderson CMB, Chen J (1993) Formation of cogenetic quartz and nepheline syenites.
 Geochim Cosmochim Acta 57:697–704. https://doi.org/10.1016/0016-7037(93)90380-F
- Goodenough KM, Wall F, Merriman D (2018) The Rare Earth Elements: Demand, Global Resources, and
 Challenges for Resourcing Future Generations. Nat Resour Res 27:201–216.
 https://doi.org/10.1007/s11053-017-9336-5
- 838 Gowans RM, Lewis WJ, Zalnieriunas RV (2017) Quest Rare Minerals Ltd.: Strange Lake Resource Estimation
- Grigor'eva AA, Zubkova NV, Pekov IV, et al (2011) Crystal chemistry of elpidite from Khan Bogdo (Mongolia)
 and its K-and Rb-exchanged forms. Crystallogr Rep 56:832
- Gysi AP, Williams-Jones AE, Collins P (2016) Lithogeochemical vectors for hydrothermal processes in the
 Strange Lake peralkaline granitic REE-Zr-Nb deposit. Econ Geol 111:1241–1276
- Hatch GP (2015) TMR Advanced Rare-Earth Projects Index Technology Metals Research.
 http://www.techmetalsresearch.com/metrics-indices/tmr-advanced-rare-earth-projects-index/. Accessed
 27 Jun 2018
- Hawthorne FC, Oberti R, Cannillo E, et al (2001) Li-bearing arfvedsonitic amphiboles from the Strange Lake
 peralkaline granite, Quebec. Can Mineral 39:1161–1170
- Hawthorne FC, Oberti R, Harlow GE, et al (2012) Nomenclature of the amphibole supergroup. Am Mineral
 97:2031–2048
- Hawthorne FC, Oberti R, Ungaretti L, et al (1996) Fluor-ferro-leakeite, NaNa2 (Fe2+ 2Fe3+ 2Li) Si8O22F2, a
 new alkali amphibole from the Canada Pinabete pluton, Questa, New Mexico, USA. Am Mineral
 81:226–228
- Ishihara S, Hua R, Hoshino M, Murakami H (2008) REE abundance and REE minerals in granitic rocks in the
 Nanling range, Jiangxi Province, southern China, and generation of the REE-rich weathered crust
 deposits. Resour Geol 58:355–372
- Jones AP, Peckett A (1981) Zirconium-bearing aegirines from Motzfeldt, south Greenland. Contrib Mineral
 Petrol 75:251–255

- Keppler H, Wyllie PJ (1991) Partitioning of Cu, Sn, Mo, W, U, and Th between melt and aqueous fluid in the systems haplogranite-H 2 O- HCl and haplogranite-H 2 O- HF. Contrib Mineral Petrol 109:139–150
- Kogarko LN, Williams CT, Woolley AR (2002) Chemical evolution and petrogenetic implications of loparite in the layered, agpaitic Lovozero complex, Kola Peninsula, Russia. Mineral Petrol 74:1–24
- Kovalenko VI, Yarmoluyk VV, Sal'nikova EB, et al (2006) Geology, geochronology, and geodynamics of the
 Khan Bogd alkali granite pluton in southern Mongolia. Geotectonics 40:450–466
- Kovalenko VI, Yarmolyuk VV (1995) Endogenous rare metal ore formations and rare metal metallogeny of
 Mongolia. Econ Geol 90:520–529
- Kramm U, Kogarko LN (1994) Nd and Sr isotope signatures of the Khibina and Lovozero agpaitic centres, Kola
 Alkaline province, Russia. Lithos 32:225–242. https://doi.org/10.1016/0024-4937(94)90041-8
- Kynicky J, Chakhmouradian AR, Xu C, et al (2011) Distribution and evolution of zirconium mineralization in peralkaline granites and associated pegmatites of the Khan Bogd complex, southern Mongolia. Can Mineral 49:947–965
- 871 Lacroix A (1923) Minéralogie de Madagascar. A. Challamel, éditeur, Librairie maritime et coloniale
- Lagarec K, Rancourt DG (1997) Extended Voigt-based analytic lineshape method for determining N dimensional correlated hyperfine parameter distributions in Mössbauer spectroscopy. Nucl Instrum
 Methods Phys Res Sect B Beam Interact Mater At 129:266–280
- Larsen LM (1976) Clinopyroxenes and coexisting mafic minerals from the alkaline Ilimaussaq intrusion, South
 Greenland. J Petrol 17:258–290
- Larsen LM (1981) Sector zoned aegirine from the Ilímaussaq alkaline intrusion, South Greenland. Contrib
 Mineral Petrol 76:285–291
- Larsen LM, Sørensen H (1987) The Ilímaussaq intrusion—progressive crystallization and formation of layering
 in an agpaitic magma. Geol Soc Lond Spec Publ 30:473–488
- Leake BE, Woolley AR, Arps CE, et al (1997) Nomenclature of amphiboles; report of the Subcommittee on Amphiboles of the International Mineralogical Association Commission on new minerals and mineral names. Mineral Mag 61:295–310
- Li X-H, Li W-X, Li Q-L, et al (2010) Petrogenesis and tectonic significance of the 850 Ma Gangbian alkaline
 complex in South China: evidence from in situ zircon U–Pb dating, Hf–O isotopes and whole-rock
 geochemistry. Lithos 114:1–15
- Liu Y, Chen Z, Yang Z, et al (2015) Mineralogical and geochemical studies of brecciated ores in the Dalucao
 REE deposit, Sichuan Province, southwestern China. Ore Geol Rev 70:613–636.
 https://doi.org/10.1016/j.oregeorev.2015.03.006
- Locock AJ (2014) An Excel spreadsheet to classify chemical analyses of amphiboles following the IMA 2012
 recommendations. Comput Geosci 62:1–11
- London D (2008) Pegmatites. Sp. Pub. 10. Can Miner 347:
- 893 Lucas J, Lucas P, Le Mercier T, et al (2014) Rare earths: science, technology, production and use. Elsevier
- Mahood GA, Stimac JA (1990) Trace-element partitioning in pantellerites and trachytes. Geochim Cosmochim
 Acta 54:2257–2276. https://doi.org/10.1016/0016-7037(90)90050-U
- Mann U, Marks M, Markl G (2006) Influence of oxygen fugacity on mineral compositions in peralkaline melts:
 The Katzenbuckel volcano, Southwest Germany. Lithos 91:262–285.
 https://doi.org/10.1016/j.lithos.2005.09.004

- Marks M, Halama R, Wenzel T, Markl G (2004) Trace element variations in clinopyroxene and amphibole from
 alkaline to peralkaline syenites and granites: implications for mineral-melt trace-element partitioning.
 Chem Geol 211:185–215
- 902 Marks MA, Markl G (2017) A global review on agpaitic rocks. Earth-Sci Rev 173:229–258
- 903 Migdisov A, Williams-Jones AE, Brugger J, Caporuscio FA (2016) Hydrothermal transport, deposition, and
 904 fractionation of the REE: Experimental data and thermodynamic calculations. Chem Geol 439:13–42
- 905 Miller RM (1983) The Pan-African Damara Orogen of South West Africa/Namibia. Evol Damara Orogen South
 906 West Afr
- 907 Miller RR (1996) Structural and textural evolution of the Strange Lake peralkaline rare-element (NYF) granitic
 908 pegmatite, Quebec-Labrador. Can Mineral 34:349–371
- Mollo S, Blundy JD, Giacomoni P, et al (2017) Clinopyroxene-melt element partitioning during interaction
 between trachybasaltic magma and siliceous crust: Clues from quartzite enclaves at Mt. Etna volcano.
 Lithos 284–285:447–461. https://doi.org/10.1016/j.lithos.2017.05.003
- Moore M, Chakhmouradian AR, Mariano AN, Sidhu R (2015) Evolution of rare-earth mineralization in the Bear
 Lodge carbonatite, Wyoming: Mineralogical and isotopic evidence. Ore Geol Rev 64:499–521.
 https://doi.org/10.1016/j.oregeorev.2014.03.015
- 915 Morimoto N (1988) Nomenclature of Pyroxenes. Mineral Petrol 39:55–76
- 916 Nakamura Y (1973) Origin of sector-zoning of igneous clinopyroxenes. Am Mineral 58:986–990
- 917 Nardi LVS, Bitencourt M de F (2009) A-type granitic rocks in post-collisional settings in southernmost Brazil:
 918 their classification and relationship with tectonics and magmatic series. Can Mineral 47:1493–1503.
 919 https://doi.org/10.3749/canmin.47.6.1493
- 920 Nielsen TFD (1979) The occurrence and formation of Ti-aegirines in peralkaline syenites. Contrib Mineral
 921 Petrol 69:235-244
- 922 Ortoleva P, Merino E, Moore C, Chadam J (1987) Geochemical self-organization I; reaction-transport feedbacks
 923 and modeling approach. Am J Sci 287:979–1007
- Pennycook SJ, Nellist PD (2011) Scanning Transmission Electron Microscopy: Imaging and Analysis. Springer
 Science & Business Media
- Piilonen PC, McDonald AM, Lalonde AE (1998) The crystal chemistry of aegirine from Mont Saint-Hilaire,
 Quebec. Can Mineral 36:779–791
- Pillet D, Bonhomme MG, Duthou JL, Chenevoy M (1989) Chronologie Rb/Sr et K/Ar du granite peralcalin du lac Brisson, Labrador central, Nouveau-Québec. Can J Earth Sci 26:328–332
- Poitrasson F, Duthou J-L, Pin C (1995) The relationship between petrology and Nd isotopes as evidence for
 contrasting anorogenic granite genesis: example of the Corsican Province (SE France). J Petrol
 36:1251–1274
- Poitrasson F, Paquette J-L, Montel J-M, et al (1998) Importance of late-magmatic and hydrothermal fluids on the
 Sm–Nd isotope mineral systematics of hypersolvus granites. Chem Geol 146:187–203
- 935 Rakotovao S, Rakotondrazafy R, Beziat D, et al (2009) Pétrologie du complexe alcalin cénozoïque
 936 d'Ambohimirahavavy, presqu'île d'Ampasindava, nord-ouest de Madagascar. Mada-Géo 13:2–19

937 Ranløv J, Dymek RF (1991) Compositional zoning in hydrothermal aegirine from fenites in the Proterozoic 938 Gardar Province, South Greenland. Eur J Mineral 837–854. https://doi.org/10.1127/ejm/3/5/0837

- Roelofsen JN (1997) The primary and secondary mafic silicates of two alkaline anorogenic complexes: Strange
 Lake (Quebec-Labrador) and Amba Dongar (Gujarat, India). PhD Thesis, McGill University Libraries
- 941 Rønsbo JG, Pedersen AK, Engell J (1977) Titan-aegirine from early Tertiary ash layers in northern Denmark.
 942 Lithos 10:193–204
- Salvi S, William-Jones A (1995) Zirconosilicate phase relations in the Strange Lake (Lac Brisson) pluton,
 Quebec-Labrador, Canada: American Mineralogist. Am Mineral 80:1031–1040.
 https://doi.org/10.2138/am-1995-9-1019
- Salvi S, Williams-Jones AE (1990) The role of hydrothermal processes in the granite-hosted Zr, Y, REE deposit
 at Strange Lake, Quebec/Labrador: evidence from fluid inclusions. Geochim Cosmochim Acta
 54:2403-2418
- Salvi S, Williams-Jones AE (2006) Alteration, HFSE mineralisation and hydrocarbon formation in peralkaline
 igneous systems: Insights from the Strange Lake Pluton, Canada. Lithos 91:19–34
- Salvi S, Williams-Jones AE (1996) The role of hydrothermal processes in concentrating high-field strength
 elements in the Strange Lake peralkaline complex, northeastern Canada. Geochim Cosmochim Acta
 60:1917–1932
- Salvi S, Williams-Jones AE (1997) Fischer-Tropsch synthesis of hydrocarbons during sub-solidus alteration of
 the Strange Lake peralkaline granite, Quebec/Labrador, Canada. Geochim Cosmochim Acta 61:83–99
- Sanematsu K, Kon Y, Imai A, et al (2013) Geochemical and mineralogical characteristics of ion-adsorption type
 REE mineralization in Phuket, Thailand. Miner Deposita 48:437–451. https://doi.org/10.1007/s00126 011-0380-5
- Schmitt AK, Trumbull RB, Dulski P, Emmermann R (2002) Zr-Nb-REE mineralization in peralkaline granites
 from the Amis Complex, Brandberg (Namibia): evidence for magmatic pre-enrichment from melt
 inclusions. Econ Geol 97:399–413
- 962 Shannon RD (1976) Revised effective ionic radii and systematic studies of interatomic distances in halides and
 963 chalcogenides. Acta Crystallogr 32:751–767
- Sheard ER, Williams-Jones AE, Heiligmann M, et al (2012) Controls on the Concentration of Zirconium,
 Niobium, and the Rare Earth Elements in the Thor Lake Rare Metal Deposit, Northwest Territories,
 Canada. Econ Geol 107:81–104. https://doi.org/10.2113/econgeo.107.1.81
- 967 Shearer CK, Larsen LM (1994) Sector-zoned aegirine from the Ilimaussaq alkaline intrusion, South Greenland:
 968 Implications for trace-element behavior in pyroxene. Am Mineral 79:340–352
- Siegel K, Williams-Jones AE, Stevenson R (2017a) A Nd-and O-isotope study of the REE-rich peralkaline
 Strange Lake granite: implications for Mesoproterozoic A-type magmatism in the Core Zone (NE Canada). Contrib Mineral Petrol 172:54
- Siegel K, Williams-Jones AE, van Hinsberg VJ (2017b) The amphiboles of the REE-rich A-type peralkaline
 Strange Lake pluton–fingerprints of magma evolution. Lithos 288:156–174
- 974 Smith MP (2007) Metasomatic silicate chemistry at the Bayan Obo Fe–REE–Nb deposit, Inner Mongolia, China:
 975 contrasting chemistry and evolution of fenitising and mineralising fluids. Lithos 93:126–148
- Smith MP, Henderson P, Jeffries TER, et al (2004) The rare earth elements and uranium in garnets from the
 Beinn an Dubhaich Aureole, Skye, Scotland, UK: constraints on processes in a dynamic hydrothermal
 system. J Petrol 45:457–484
- 979 Strong DF (1969) Formation of the hour-glass structure in augite. Mineral Mag 37:472–479
- Sun S-S, McDonough W (1989) Chemical and isotopic systematics of oceanic basalts: implications for mantle
 composition and processes. Geol Soc Lond Spec Publ 42:313–345

- Thomas RJ, De Waele B, Schofield DI, et al (2009) Geological evolution of the Neoproterozoic Bemarivo Belt,
 northern Madagascar. Precambrian Res 172:279–300. https://doi.org/10.1016/j.precamres.2009.04.008
- Trail D, Bruce Watson E, Tailby ND (2012) Ce and Eu anomalies in zircon as proxies for the oxidation state of magmas. Geochim Cosmochim Acta 97:70–87. https://doi.org/10.1016/j.gca.2012.08.032
- 986 U. S. Geological Survey (2019) Rare Earths Statistics and Information. In: Miner. Commod. Summ. https://www.usgs.gov/centers/nmic/rare-earths-statistics-and-information. Accessed 28 Aug 2019
- Ubide T, McKenna CA, Chew DM, Kamber BS (2015) High-resolution LA-ICP-MS trace element mapping of
 igneous minerals: In search of magma histories. Chem Geol 409:157–168.
 https://doi.org/10.1016/j.chemgeo.2015.05.020
- Ubide T, Mollo S, Zhao J, et al (2019) Sector-zoned clinopyroxene as a recorder of magma history, eruption
 triggers, and ascent rates. Geochim Cosmochim Acta 251:265–283.
 https://doi.org/10.1016/j.gca.2019.02.021
- 994 Vasyukova O, Williams-Jones A (2020) Partial melting, fractional crystallisation, liquid immiscibility and
 995 hydrothermal mobilisation–A `Recipe'for the formation of economic A-Type granite-hosted HFSE
 996 deposits. Lithos 356:105300
- 997 Vasyukova O, Williams-Jones AE (2014) Fluoride-silicate melt immiscibility and its role in REE ore formation:
 998 Evidence from the Strange Lake rare metal deposit, Québec-Labrador, Canada. Geochim Cosmochim
 999 Acta 139:110–130
- Vasyukova OV, Williams-Jones AE (2019) Closed system fluid-mineral-mediated trace element behaviour in
 peralkaline rare metal pegmatites: Evidence from Strange Lake. Chem Geol 505:86–99
- 1002 Vasyukova OV, Williams-Jones AE, Blamey NJF (2016) Fluid evolution in the Strange Lake granitic pluton,
 1003 Canada: Implications for HFSE mobilisation. Chem Geol 444:83–100
- 1004 Veksler IV, Dorfman AM, Dulski P, et al (2012) Partitioning of elements between silicate melt and immiscible
 1005 fluoride, chloride, carbonate, phosphate and sulfate melts, with implications to the origin of
 1006 natrocarbonatite. Geochim Cosmochim Acta 79:20–40. https://doi.org/10.1016/j.gca.2011.11.035
- Wang Q, Deng J, Liu X, et al (2010) Discovery of the REE minerals and its geological significance in the Quyang bauxite deposit, West Guangxi, China. J Asian Earth Sci 39:701–712. https://doi.org/10.1016/j.jseaes.2010.05.005
- William-Jones AE, Migdisov AA, Samson IM (2012) Hydrothermal mobilisation of the Rare Earth Elements–a
 Tale of" Ceria" and" Yttria. Elements 8:355–360
- Xu C, Kynickỳ J, Smith MP, et al (2017) Origin of heavy rare earth mineralization in South China. Nat Commun
 8:14598
- Yang X-M, Yang X-Y, Zheng Y-F, Le Bas MJ (2003) A rare earth element-rich carbonatite dyke at Bayan Obo,
 Inner Mongolia, North China. Mineral Petrol 78:93–110
- Zaitsev AN, Terry Williams C, Jeffries TE, et al (2014) Rare earth elements in phoscorites and carbonatites of the Devonian Kola Alkaline Province, Russia: Examples from Kovdor, Khibina, Vuoriyarvi and Turiy Mys complexes. Ore Geol Rev 61:204–225. https://doi.org/10.1016/j.oregeorev.2014.02.002
- 1019
- 1020
- 1021 <u>Tables caption</u>
- 1022

Table 1 Mean values for the major and trace element composition, obtained by microprobe and LA-ICPMS
 respectively, of amphiboles from pegmatites and granites from the six complexes. Abbreviations: no.: number of
 replicates; F-arf: fluoro-arfvedsonite; F-f-f-lea: ferro-ferri-fluoro-leakeite; F-f-kato: ferro-ferri-katophorite; b.d.l:

1026 below detection limit; Ambo: Ambohimirahavavy; KB: Khan bogd; M: Manongarivo; SL: Strange Lake

1027

Table 2 Mean values (number of analysis given in bold) for the major and trace element composition, obtained
by microprobe and LA-ICPMS respectively, of aegirine from pegmatites and granites from six complexes
worldwide in both pegmatites and granites. Granites only comprise type-II aegirine. T I, T II: aegirine type-I, II.
Abbreviations: no.: number of replicates; b.d.l: below detection limit; Ambo: Ambohimirahavavy; KB: Khan
bogd; M: Manongarivo; SL: Strange Lake

1033

Table 3 Mean composition of cores and rims of type-I aegirine from 5 complexes. Abbreviations: no.: number of
 replicates; b.d.l: below detection limit; Ambo: Ambohimirahavavy; KB: Khan bogd; M: Manongarivo

1036

1037 Figures caption

1038

1039 Fig. 1 A logarithmic diagram plotting chondrite-normalized (Sun and McDonough 1989) values of whole-rock 1040 Yb/La vs Yb contents for different alkaline complexes worldwide. Five groups are distinguished: alkaline 1041 granites, alkaline pegmatites, carbonatites, nepheline syenites, and ion-adsorption deposits. Data are from this 1042 study plus from Estrade et al. (2014b), Boily and Williams-Jones (1994), Estrade (2014), Kynicki et al. (2011), 1043 Poitrasson et al. (1995), Vasyukova and Williams-Jones (2014), Schmitt et al. (2002), Moore et al. (2015), Yang 1044 et al. (2003), Xu et al. (2017), Liu et al. (2015), Zaitsev et al. (2014), Kogarko et al. (2002), Li et al. (2010), 1045 Sørensen et al. (1987), Hatch (2015), Sanematsu et al. (2013), Ishihara et al. (2008), Wang et al. (2010), and 1046 Chengyu et al. (1990)

1047

1048 Fig. 2 Map localizing the six complexes studied in this paper. The geodynamic context and ages are given1049 below their names in the rectangular boxes

1050

Fig. 3 Photographs of outcrop or hand samples of pegmatites from the six complexes. Most display a well-marked layering defined by mineralogy and textural variations

1053

1054 Fig. 4 a Altered astrophyllite from Amis, b REE-bearing pseudomorph (Manongarivo), c Zircon-quartz 1055 pseudomorph (Evisa), d Elpidite (Khan Bogd), altered to armstrongite and zircon in its center, e Arfvedsonite 1056 replaced by type-II aegirine (Evisa), f A type-I aegirine showing core to rim zoning (Ambohimirahavavy), g 1057 Two tiny fluid inclusions in the rim of a type-I crystal of aegirine (Khan Bogd), h Mineral inclusions contained 1058 within the rim of a type-I aegirine (Ambohimirahavavy) that are of the same nature than minerals in REE-1059 bearing pseudomorph, i A single crystal of type-II aegirine (Strange Lake), j Zoned amphibole, from katophorite 1060 in the core to arfvedsonite in the rims (Evisa), \mathbf{k} A type-I aggiring showing core to rim zoning with zone B (Khan 1061 Bogd), I Zoned amphibole, from arfvedsonite in the core to leakeite in the rims (Evisa). Photographs a, c, e, f, g, 1062 i and k were taken with an optical microscope; photographs b, h, j and l with SEM EDS; and d with optical 1063 cathodoluminescence. pseudomorph; zrn: zircon; aeg: aegirine; amph: amphiboles; qtz: quartz; afs: 1064 alkali feldspar; bsn: bastnäsite-(Ce); arf: arfvedsonite; ox: Ti- and Fe-oxides; calh: calciohilairite; CKZ: 1065 unidentified Ca- and K-bearing zirconosilicates; CMZ: unidentified Ca- and Mn-bearing zirconosilicates; 1066 NbYsil: unidentified Nb- and Y-bearing silicate; arm: armstrongite; elp: elpidite; astr: astrophyllite

1067

1068 Fig. 5 Chemical composition of amphiboles. a, b, c Ternary diagrams representing amphibole poles for the a 1069 Evisa and Manongarivo complexes; b Ambohimirahavavy and Amis complexes; c Khan Bogd and Strange Lake 1070 complexes. Poles are plotted as 10 Mg and 10 Ca for a better visualization of the dataset. d Boxplots showing the trace-element composition of amphiboles from the different complexes. Thick black line represents the median, 1071 1072 upper and lower boxplot limits the first and third quartile respectively, upper and lower dashed lines are the 1073 maximum and minimum values respectively, and the thin black line is the detection limit. Areas are shaded to 1074 distinguish data for each complex M: Manongarivo; Amb: Ambohimirahavavy; SL: Strange Lake; KB: Khan 1075 Bogd

1076

1077 Fig. 6 Microprobe and LA-ICPMS maps on a type-I aegirine crystal at Ambohimirahavavy. Visible zonations
1078 were also found in the other complexes (see Supplementary material). 3 zones are distinguished: the core A, and
1079 the sector-zoned rims Cand D. Microprobe maps are in wt%, and LA-ICPMS maps are in ppm. Limits of

aegirine on LA-ICPMS maps are blurry because the signal from the crystal and the matrix mix and do not reflectan actual concentration

1082

1083 Fig. 7 Chemical composition of type-I and type-II aegirine crystals. Type-I aegirine is split between core and 1084 rim. Core and rim analyses are identified for Type-I aegirine. a, b, c are for aegirine, (Ti, Zr)-aegirine, and 1085 clinoferrosilite; when figures; \mathbf{d} , \mathbf{e} , \mathbf{f} are for aegirine + (Ti, Zr)-aegirine, hedenbergite, and clinoferrosilite. \mathbf{a} , \mathbf{d} 1086 complexes Evisa and Manongarivo; b, e complexes Ambohimirahavavy and Amis; c, f Khan Bogd and Strange 1087 Lake; g Box diagrams comparing the trace element contents of aegirine from the different complexes. Thick 1088 black line represents the median, upper and lower boxplot limits the first and third quartile respectively, upper 1089 and lower dashed lines are the maximum and minimum values respectively, and the thin black line is the 1090 detection limit. Colours are keyed to the legend given in a-f. Areas are shaded to distinguish data for each 1091 complex

1092

1093 Fig. 8 Logarithmic diagrams plotting chondrite-normalized REE composition obtained by LA-ICP-MS for1094 amphiboles (normalization from Sun and McDonough, 1989) for all six complexes

1095

Fig. 9 Logarithmic diagrams plotting chondrite-normalized REE composition obtained by LA-ICPMS for
 aegirine (normalization from Sun and McDonough, 1989) for all six complexes. Large spectra presents all LA ICP-MS analyses for type-I and type-II aegirine, whereas small spectra on the side show an example of analyses
 performed on a single type-I crystal

1100

Fig. 10 Diagram plotting REE concentration obtained by LA-ICPMS in the core and rim of type-I pyroxene in 5 complexes. The typical analytical error is provided by the cross on the top right corner and is similar for all provided analyses

1104

Fig. 11 Histograms plotting the difference of REE concentration between the core and the rims of type-Iaegirine, and thereby REE fractionation from core to rim. Negative values stand for a higher REE concentration

in the core, and positive ones for a higher concentration in the rims. The rim enrichment in Khan Bogd is anartifact. Calculated error is shown by black lines at the end of the histogram bars

1109

1110 Fig. 12 A schematic illustration of a model for aegirine and amphiboles formation. The arrows signify transfer of 1111 elements (in the boxes) to the different minerals from the melt or fluid (light blue), the line thickness is 1112 proportional to the amount of transfer, and a dashed line indicates a possible transfer. Ligands in the fluids are 1113 also shown. During the early magmatic stage, the competition for REE, Zr, Hf, Zr and Sn is low to moderate 1114 (depending on the co-presence or not of amphibole, pyroxene and complex zirconosilicate) and allows a relative 1115 enrichment in these elements in type-I aegirine and, mostly for REE, in amphibole. During the hydrothermal 1116 stage, the competition for these elements is stronger as it involves zircon and REE-bearing minerals; 1117 consequently, type-I aegirine rims are globally depleted. Fluids composition leads to a different mobilization of 1118 light and heavy REE, originating REE fractionation. Meanwhile, hydrothermal fluids replace amphiboles by 1119 type-II secondary aegirine and Fe-Ti-oxides 1120 1121 **Electronic Supplementary Material** 1122 Online Resource 1 Text and maps detailing the geological background of all six studied complexes 1123 1124 Online Resource 2 Table of standards used for calibration of EPMA and associated detection limits. 1125 Abbreviation: n.a.: not analyzed 1126 1127 Online Resource 3 Table of detection limits for in situ LA-ICPMS measurements 1128 1129 Online Resource 4 Microprobe maps on a type-I aegirine crystal at Amis. 3 zones are distinguished: A, the core 1130 rich in Ca, Zr, Sn, Hf and poor in Na, Fe; C, sector zoning rich in Ti, Ca; and D, sector zoning rich in Fe. The 1131 thick line represents limits of the map 1132

1133	Online Resource 5 Microprobe maps on a type-I aegirine crystal at Evisa. 3 zones are distinguished: A, the
1134	core rich in Ca, Zr, Sn, Hf and poor in Na, Fe; C, sector zoning rich in Ti, Ca; and D, sector zoning rich in Fe.
1135	The thick line represents limits of the map
1136	
1137	Online Resource 6 Microprobe maps on a type-I aegirine crystal at Khan Bogd. 4 zones are distinguished: A,
1138	the core rich in Ca, Zr, Sn, Hf and poor in Na, Fe; B; C, sector zoning rich in Ti, Ca; and D, sector zoning rich in
1139	Fe
1140	
1141	Online Resource 7 Microprobe maps on a type-I aegirine crystal at Manongarivo. 3 zones are distinguished: A,
1142	the core rich in Ca, Zr, Sn, Hf and poor in Na, Fe; C, sector zoning rich in Ti, Ca; and D, sector zoning rich in Fe
1143	
1144	Online Resource 8 Microprobe maps on a type-II aegirine crystal at Strange Lake. No core-to-rim zoning is
1145	observed, but an oscillatory zoning occurs for most elements

1146