

**Distinguishing magmatic and metamorphic processes in peralkaline rocks of the Norra Kärr complex (Southern Sweden) using textural and compositional variations of clinopyroxene and eudialyte-group minerals.**

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Originally published:

May 2017

**Journal of Petrology 58(2017), 361-384**

DOI: <https://doi.org/10.1093/petrology/egx019>

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2 complex (Southern Sweden) using textural and compositional variations of clinopyroxene and  
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4

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17

18 Running Title: Magmatic and Metamorphic Processes in the Norra Kärr Alkaline Complex

19

20 **Abstract**

21 The 1.49 Ga old Norra Kärr complex in Southern Sweden contains rocks characterized by a  
22 very high ratio of  $(\text{Na}+\text{K})/\text{Al} \geq 1.2$  and a complex and highly unusual mineralogy, including  
23 rock-forming catapleiite, eudialyte-group minerals as well as minor rinkite- and britholite-  
24 group minerals. In contrast to other well-studied examples of agpaitic rocks, the Norra Kärr  
25 rocks have been deformed and partially metamorphosed during the  
26 Sveconorwegian/Grenvillian orogeny, and are now preserved in a westward dipping synform.  
27 Magmatic and metamorphic processes at the Norra Kärr complex are distinguished by  
28 combining rock fabrics of clinopyroxene and eudialyte-group minerals. Both mineral groups  
29 are stable over a large P-T range, which makes them excellent monitors of the geochemical  
30 evolution of such systems and enables the reconstruction of magmatic and subsequent  
31 metamorphic conditions.

32 The magmatic mineral assemblage crystallized from a subsolvus syenite at  
33 continuously decreasing temperatures (700 - 450°C) and silica activity (0.6 - 0.3). Due to  
34 initially relatively low peralkalinity and reducing conditions, Zr was first incorporated in Zr-  
35 aegirine. Subsequent destabilization of the latter indicates increasing peralkalinity, oxygen  
36 fugacity and water activity, which resulted in the crystallization of early magmatic catapleiite.  
37 Crystallization of presumably later magmatic Mn- and REE-poor eudialyte-group minerals,  
38 happened as soon as sufficient Cl, REE and HFSE were enriched in the residual melt.

39 Metamorphic conditions during the Sveconorwegian/Grenvillian orogeny are  
40 constrained to T between 400 - 550°C and  $a_{\text{SiO}_2}$  range of 0.25 - 0.4. Due to deformation and  
41 interaction with fluids, post-magmatic Al-rich aegirine as well as post-magmatic eudialyte-  
42 group minerals enriched in REE, Y and Mn formed. Subsequently, the eudialyte-group  
43 minerals were destabilized and decomposed to post-magmatic catapleiite and secondary REE-

44 bearing minerals. During the whole history of the complex,  $a_{\text{SiO}_2}$  remains very similar,  
45 indicating very little interaction with the surrounding granitic rocks.

46           Regardless of the intense deformation due to folding of the Norra Kärr body during  
47 the Sveconorwegian/Grenvillian orogeny, indications for primary magmatic layering of the  
48 intrusion are retained on the deposit scale. In addition, the compositional changes of  
49 magmatic eudialyte-group minerals from the outer to the inner subunit indicate a primary  
50 geochemical evolution feature due to fractional crystallization.

51

52 Keywords: deformed nepheline syenite; rare earth elements; clinopyroxene, eudialyte-group minerals

53

## 54 INTRODUCTION

55

56 During the last decade the economic interest in high field strength elements (HFSE) and rare  
57 earth elements (REE) increased due to their broad application in high-tech products. In  
58 addition to carbonatites, alkaline and especially peralkaline rocks (molar  $\text{Na}+\text{K} > \text{Al}$ ) are  
59 particularly enriched in HFSE and comprise one of the most promising sources for future  
60 REE supply (e.g., Mitchell, 2015; Smith et al., 2015; Goodenough et al., 2016). The  
61 extraordinary enrichment of HFSE and REE in such rocks was explained by the high  
62 alkalinity and the reducing conditions during crystallization of the magma, which minimizes  
63 the loss of volatile components and maximizes the ore precipitation potential during  
64 magmatic differentiation (e.g., Kogarko, 1990; Marks et al., 2011).

65 This study investigates peralkaline rocks of the Norra Kärr complex in southern  
66 Sweden, which are characterized by an unusual mineralogy including rock-forming  
67 catapleiite, eudialyte-group minerals (EGM) and minor REE-bearing minerals of the rinkite-  
68 and britholite-groups (Adamson, 1944; Eckermann, 1968; Sjöqvist et al., 2013). Such  
69 lithologies are classified as agpaitic rocks, a term originally introduced by Ussing (1912) and  
70 later complemented by Sørensen (1960, 1997) and Marks et al. (2011). Minerals of the  
71 eudialyte group are the most common index minerals in agpaitic rocks (e.g., Sørensen, 1997;  
72 Schilling et al., 2011b). Such minerals can incorporate significant concentrations of more than  
73 30 elements, including Zr, Nb and naturally occurring lanthanides and Y, with LREE being  
74 commonly more abundant than the HREE.

75 Unlike other well-studied examples of agpaitic complexes such as the Ilímaussaq  
76 complex in Greenland (see recent reviews by Sørensen et al., 2006; Upton, 2013; Marks &  
77 Markl, 2015; Dostal, 2015) or the Lovozero and Khibina complexes in Russia (e.g. Kogarko

78 et al., 1982; Kogarko, 1987; Pekov, 1998; Arzamastsev et al., 2001, 2005), the igneous Norra  
79 Kärn body has been deformed and partially recrystallized during subsequent metamorphic  
80 events. This specific feature is used to amplify the restricted knowledge on the behavior of  
81 agpaitic rocks during metamorphism, which is to date mainly limited to studies of: the Red  
82 Wine and Kipawa complexes in Canada (Curtis & Gittins, 1979; Curtis & Currie, 1981;  
83 Currie & van Breemen, 1996; van Breemen & Currie, 2004); nepheline syenites in Malawi  
84 (e.g. Woolley et al., 1996); and peralkaline gneisses in India (Nanda et al., 2008; Goswami &  
85 Basu, 2013; Chakrabarty et al., 2016). We present detailed observations on rock and mineral  
86 textures combined with compositional data for the major rock-forming minerals to provide  
87 some direct insight into the petrological and geochemical processes accompanied by the  
88 deformation and metamorphism of agpaitic rocks.

89

## 90 **GEOLOGICAL SETTING**

91

92 The Norra Kärn complex is located approx. 2 km E of Lake Vättern, approx. 15 km NNE of  
93 the township of Gränna in Southern Sweden (Fig. 1). It is situated in the Trans-scandinavian  
94 Igneous Belt, which intruded into the Svecofennian domain of Fennoscandia in the course of  
95 juvenile crust reworking between 1.85–1.75 Ga (Andersson et al., 2007 and references  
96 therein). The Trans-scandinavian Igneous Belt rocks form a voluminous array of  
97 monzodioritic to granitic batholiths of alkali-calcic to calc-alkaline composition (Högdahl et  
98 al., 2004; Fig. 1). They extend over ca. 1400 km from NW Norway to SE Sweden. Significant  
99 igneous activity between 1.65–1.50 Ga was responsible for the formation of an active margin  
100 along the south-western border of the Fennoscandian Shield (McLelland, 1989). From ca.  
101 1470 Ma onwards A- and I-type granitoids and syenitoids were emplaced in a typical

102 volcanic-arc setting, marking the pre-collisional stage of the Danopolonian or Hallandian  
103 orogeny in the area east and southeast of Lake Vättern (Brander, 2011, Brander & Söderlund,  
104 2009). In the time of 1.5-1.4 Ga two possible settings for intra-cratonic magmatism are  
105 discussed in the literature, an anorogenic and an orogenic extension. Aberg (1988) interpreted  
106 the emplacement of alkali-calcic and calc-alkaline intrusions into the southernmost parts of  
107 the Trans-scandinavian Igneous Belt as a product of continental rifting between 1.48 and 1.35  
108 Ga (Aberg et al., 1984; 1985a; 1985b; Smellie & Stuckless, 1985).

109         During the period of around 1140 – 900 Ma wide areas of the southern and central  
110 Trans-scandinavian Igneous Belt, as well as broad areas of south-western Fennoscandia were  
111 subject to the collision of Fennoscandia with Amazonia during the  
112 Grenvillian/Sveconorwegian orogeny (e.g. Bingen et al., 2008). These areas form today the  
113 Sveconorwegian orogenic belt. The area east of Lake Vättern belongs to the frontal zone of  
114 the orogen and was mainly affected by metamorphism during the late stages of the orogeny,  
115 the so-called Falkenberg and Dalane phases (e.g. by Bingen et al., 2008 and references  
116 therein). During the Falkenberg phase (980-970 Ma) crustal thickening and associated  
117 eclogite facies conditions affected the Fennoscandian crust to at least 50 km depth. This was  
118 followed by decompression, relaxation and gravitational collapse during the Dalane phase at  
119 970-900 Ma (Bingen et al., 2006; 2008; Möller et al., 2007). During this time the N-S  
120 trending Protogine Zone was formed as a steep, sheer zone at the very front of the  
121 Sveconorwegian orogen (Andréasson & Rodhe, 1994; Wahlgren et al., 1994; Söderlund et al.,  
122 2004).

123

## 124 **The Norra Kärr complex**

125

126 The relatively small Norra Kärr complex covers an area of approx. 350 x 1100 m and is  
127 located in the Väjö Granite Suite (TIB I), between the Vimmerby Batholith in the east and  
128 the major Protogine shear zone (PZ) in the west (Fig. 1b). The body is preserved within a  
129 westwards dipping (approx. 40°) synform (Fig. 2), deformed by moderate E-W and late, weak  
130 N-S directed compression under ductile conditions (Rankin, 2011).

131 The granitic host rocks at the western contact of the Norra Kärr complex and some  
132 deeper parts in the east (only known from drilling cores) exhibit clear signs of fenitization  
133 (Adamson, 1944; Eckermann, 1968; Sjöqvist et al., 2015). These fenites are up to 100 m wide  
134 massive horizons, but also occur as decimeter-thick veins in the surrounding granite. They are  
135 characterized by the absence of quartz, which is replaced by albite, magnetite, hematite and  
136 fluorite, and the replacement of biotite by aegirine (Adamson, 1944; Sjöqvist et al., 2015).  
137 Fenitization of the alkaline rocks itself is less prominent and rarely occurs in the outer  
138 subunit, only at the direct contact to the fenitized host rocks. Recently, the fenitization of the  
139 granitic gneisses was determined to have a U-Pb zircon age of  $1.49 \pm 0.01$  Ga (Sjöqvist et al.,  
140 2014), which is interpreted coeval with the age of magmatic emplacement of the Norra Kärr  
141 complex in a presumably pre-collisional rifting setting. Younger ages determined by Sjöqvist  
142 et al. (2014) constrain the deformation during the Sveconorwegian orogeny from  $1148 \pm 5$  Ma  
143 onwards. At Norra Kärr neither the surrounding country rocks (granitic- to- syenitic gneisses)  
144 nor the metamorphosed syenites of the complex itself provide any clear evidence indicating  
145 the grade of metamorphism of the complex. Regional geological studies (Wahlgren &  
146 Stephens, 2004) report lower greenschist facies conditions for the area to the East of Lake  
147 Vättern and the Protogine shear Zone (Fig. 1). Considering this and the actual position of the  
148 complex within this major shear zone (Fig. 1), moderate- to- high greenschist facies  
149 conditions can be assumed.



150 For this study, the peralkaline rocks of the Norra Kärr complex were collectively  
151 classified as meta-nepheline syenites (after Gillespie & Styles, 1999). The most common rock  
152 type (85 vol.% of the exposed area), the so-called grennaite (Adamson, 1944) is a catapleiite-  
153 and EGM-bearing aegirine meta-nepheline syenite (Fig. 2). The texture of this unit varies  
154 systematically across the deposit (Figs 3, 4 and Electronic Appendix 1). On the basis of the  
155 frequency of medium- to- coarse-grained lenses and bands or schlieren (Törnebohm, 1906;  
156 Adamson, 1944; Sjöqvist et al., 2015), the frequency of catapleiite and the degree of  
157 deformation, the following three subunits with gradual transitions between them are  
158 distinguished: The central part (inner subunit) was strongly foliated (Figs 3a-b and 4a-c) and  
159 is best described as an aegirine leuco-meta-nepheline syenite (a.k.a. migmatitic grennaite;  
160 lithologic nomenclature by Tasman Metals, Sjöqvist et al., 2013). This area is surrounded by  
161 a zone rich in pegmatoidal schlieren (mid subunit, Figs 3e-h and 4e and i-k) named pegmatite-  
162 bearing aegirine meta-nepheline syenite (a.k.a. pegmatitic grennaite; Adamson, 1944). The  
163 border area (outer subunit, Figs 3c and 4f-h) of the complex is defined by a foliated  
164 catapleiite-bearing aegirine meta-nepheline syenite (a.k.a. catapleiite grennaite; Adamson,  
165 1944).

166 Minor rock types of the Norra Kärr complex include the strongly folded fluoro-  
167 leakeite-aegirine meta-nepheline syenite (previously called kaxtorpita, Fig. 4d), the aegirine-  
168 amphibole-biotite meta-nepheline syenite (a.k.a. pulaskite) and the arfvedsonite-aegirine  
169 meta-nepheline syenite (former lakarpita), well-known from the discovery outcrop of Norra  
170 Kärr (Sjöqvist et al., 2013).

171 The most common deformation features of all rock types include the alignment and  
172 stretching of certain minerals (e.g. clinopyroxene, EGM, catapleiite, Fig. 4a-e) and the  
173 presence of porphyroclasts (mainly microcline, Fig. 4f-h), which have tails of recrystallized

174 material (Fig. 4h). They are commonly preserved in a finer-grained, probably recrystallized  
175 matrix and form typical shear related patterns (sigma- and delta-clast, Fig. 4f).

176

## 177 **PETROGRAPHY**

178

179 The phase assemblages and micro-textural characteristics of the investigated samples are  
180 described in the following. A list of investigated samples is given in Table 1, sample locations  
181 are indicated in Fig.2. The modal mineralogy of the main subunits is summarized in Table 2.

182

### 183 **Catapleiite-and EGM-bearing aegirine meta-nepheline syenite (grennaite)**

184 Inner subunit (aegirine leuco-meta-nepheline syenite)

185

186 This unit displays strong foliation and in places intense folding, with rather common  
187 crenulated foliation and gneissic textures (Figs 3a and b and 4a-c). The main felsic minerals  
188 are subhedral- to- anhedral and fine-grained albite (up to 30 wt.%), microcline (up to 20  
189 wt.%) and nepheline (approx. 10 wt.%), the latter being partly- to- significantly-replaced by  
190 natrolite (10-35 wt.%). Fine- to- medium-grained aegirine (15-20 wt.%) occurs as subhedral-  
191 to- anhedral, needle or columnar-shaped crystals, which commonly exhibit irregular rims  
192 and/or replacement textures of darker BSE contrast (Fig. 5c). Fine- to- medium-grained EGM  
193 (approx. 15 wt.%), can be concentrated in hinge zones of folds and be partly enriched in mm-  
194 wide bands, mainly composed of feldspar and clinopyroxene (Fig. 4c). In these bands  
195 elongated minerals are aligned while coarser-grained minerals form  $\sigma$ - and/or  $\delta$ -clasts.

196 Eudialyte-group minerals are commonly elongated and ductile deformed in the direction of  
197 foliation (Figs 3a, 4a and 6c). The crystals show no zoning or overgrowth textures but are  
198 commonly intergrown with fine-grained, anhedral catapleiite and minor britholite-group  
199 minerals. The most common accessories include galena and sphalerite.

200

201 Mid subunit (pegmatite-bearing aegirine meta-nepheline syenite)

202

203 This subunit contains variable amounts of leucocratic, mid- to coarse-grained and partly-  
204 pegmatitic nepheline syenite schlieren (up to ten cm in thickness) set in a fine-grained  
205 aegirine meta-nepheline syenite matrix. The dark green-grey matrix (grain size < 1mm) is  
206 characterized by a homogeneous mineral distribution with no specific accumulation of  
207 minerals (Fig. 3d). Deformation is displayed by the alignment of most of the mineral grains  
208 (Fig. 4e) and stretching of rarely occurring larger crystals (e.g. catapleiite needles). The main  
209 rock-forming minerals are anhedral albite (55 wt.%), nepheline (10 wt.%) and clinopyroxene  
210 (20 wt.%) with minor microcline (5 wt.%), catapleiite (3 wt.%), EGM (2 wt.%), natrolite (3  
211 wt.%) and accessory britholite- and rinkite-group minerals ( $\leq 0.25$  wt.%). The latter are  
212 always spatially associated with EGM.

213 The schlieren (Figs 3e-h and 4i-k) generally follow the foliation and the folding of the  
214 rock. However, they also may be intercalated with the fine-grained matrix. The following  
215 types of schlieren can be distinguished:

216 (A) Fine- to- medium-grained schlieren with no alignment of crystals, mainly  
217 consisting of feldspar, clinopyroxene, euhedral to subhedral EGM and nepheline (e.g. Figs 3f  
218 and h, 4i, 6a and b and 7d and e). In general, the relative content of clinopyroxene increases

219 towards the rims of these schlieren. Eudialyte-group minerals are medium- to- coarse-grained  
220 reaching several mm in size (Figs 3f and h, 4k and 6a and b). Larger crystals are commonly  
221 sector- or oscillatory-zoned. Porous and irregular zones in the central parts of euhedral  
222 crystals can be complexly overgrown by areas with lower BSE contrast and along the rims of  
223 such crystals flame textures with higher BSE contrast may be present (Figs 4k and 6a and b).  
224 Rims and cracks are commonly associated with catapleiite and britholite-group minerals,  
225 occurring as filling of interstitial spaces or replacing EGM (Fig. 6a and b). Catapleiite occurs  
226 as two textural varieties, (1) catapleiite spatially associated or intergrown with EGM,  
227 preferably occurring at the border area between clinopyroxene and EGM (Fig. 7e) and (2)  
228 catapleiite inclusions in fan-shaped sectors of radiating clinopyroxene aggregates (Figs 4i and  
229 7d). These radial aggregates of subhedral clinopyroxenes (Fig. 7d) are unique to this schlieren  
230 type. They are irregularly rimmed by areas with darker BSE contrast.

231 (B) Medium- to- coarse-grained schlieren with no general alignment of crystals. They  
232 are mainly composed of subhedral- to- euhedral feldspar, nepheline, clinopyroxene and EGM  
233 (Figs 3e and g, 4j, 5a and b and 7c). Euhedral- to subhedral eudialyte-group minerals mainly  
234 exhibit sector zoning. Similar to schlieren (A) these are overgrown and/or replaced by  
235 irregular or flamy areas with brighter BSE contrast (Fig. 4k). In cases EGM are extensively  
236 altered and replaced by catapleiite and britholite-group minerals (Fig. 3g). In contrast to  
237 schlieren (A), catapleiite is not necessarily spatially associated with EGM or clinopyroxene. It  
238 may form needles, laths (rarely with a bluish color in hand specimen) or subhedral crystals  
239 (Figs 3e and 7b). Clinopyroxene forms euhedral, coarse-grained and oscillatory- and/or  
240 sector-zoned crystals (Fig. 5a and b). An exceptionally large (> 1 cm) and euhedral  
241 clinopyroxene crystal was observed in sample 5408 (Figs 5a and 7c). This crystal exhibits a  
242 partly-resorbed Zr-rich core, surrounded by an area rich in anhedral catapleiite inclusions,  
243 followed by well-developed Ti-rich sectors and late irregular rims of Al-rich aegirine.

244 In both schlieren types, large, patchy aggregates composed of numerous euhedral fine-  
245 grained clinopyroxene crystals (Fig. 3e and h) occur. Nepheline and feldspar are extensively  
246 altered to natrolite (Figs 3e-h and 4e and j).

247

248 Outer subunit (catapleiite-bearing aegirine meta-nepheline syenite)

249

250 Both, magmatic and metamorphic terminology is used to describe the textural appearance of  
251 this subunit, although the classification of some of the features is not clear. This will be  
252 further commented in the discussion section.

253 This unit is up to several hundred meters in width, is schistose and fine-grained- to-  
254 aphanitic with porphyroclastic textures (Figs 3c and 4f-h). The contact towards the  
255 surrounding host rocks is characterized by an intense bleaching zone. Mineralogically, this  
256 rock type is very similar to the matrix of the mid subunit described above (Table 2), but is  
257 characterized by the absence of schlieren. Characteristic for this unit are medium-grained  
258 feldspars that form  $\sigma$ - and/or  $\delta$ -clasts (Fig. 4f) as well as distinct EGM and catapleiite  
259 textures: In addition to fine-grained EGM in the matrix, larger poikilitic EGM porphyroclasts  
260 (Figs 3c, 4h and 6d) with abundant catapleiite (Figs 4h and 6d, Zr map), feldspar, nepheline  
261 and clinopyroxene inclusions occur, commonly surrounded by columnar rinkite-group  
262 minerals (Figs 4h and 6d). Furthermore, EGM in the outermost bleaching zones are partly- to-  
263 entirely replaced by zircon. Catapleiite forms up to 2 cm large, porphyroclasts (Fig. 4g) as  
264 well as elongated aggregates composed of anhedral, recrystallized catapleiite crystals (Fig.  
265 7f). Clinopyroxene occurs as fine-grained euhedral- to- anhedral crystals rimed by a later  
266 clinopyroxene generation.

267

268 **Other rock units**

269 Arfvedsonite-aegirine meta-nepheline syenite (a.k.a. lakarpite)

270

271 Although being only a minor rock type restricted to the eastern contact of the mid subunit  
272 with the outer subunit (Fig. 2), this rock type is best known from the so-called “*discovery*  
273 *outcrop*” of the complex (Sjöqvist et al., 2013). This fine- to medium-grained unit mainly  
274 consists of feldspar (35-60 wt.%), arfvedsonite (up to 20 wt.%), clinopyroxene (up to 15  
275 wt.%) and EGM (up to 12 wt.%) with minor nepheline and natrolite. Feldspar-rich schlieren  
276 contain cm-sized, pinkish and mostly poikilitic EGM commonly surrounded by an amphibole-  
277 and clinopyroxene-rich matrix. Clinopyroxene forms elongated to euhedral crystals, most  
278 other minerals are anhedral- to- subhedral in their habit.

279

280 Fluoro-leakeite-aegirine meta-nepheline syenite (a.k.a. kaxtorpite)

281

282 This fine- to- medium-grained rock with dark green- to- black colour was first described by  
283 Adamsson (1944) and is located in the very centre of the intrusion (Fig. 2). The rock is  
284 variably deformed showing alignment of elongated minerals culminating in intense isoclinal  
285 folding and crenulated foliation (Fig. 4d). Most minerals are anhedral, elongated and aligned  
286 forming small alternating bands. The fine-grained matrix consists of albite, fluoro-leakeite  
287 (Oberti et al., 2015), pectolite, clinopyroxene, nepheline and natrolite ( $\pm$  catapleiite, titanite,

288 fluorite and rosenbuschite) with large microcline augen preserved. The rock is commonly  
289 interfolded with the surrounding inner subunit.

290

291 Aegirine-amphibole-biotite meta-nepheline syenite (a.k.a. pulaskite)

292

293 This rock is only present in the western and northern part of the complex, mostly forming  
294 elongated bodies in the outer subunit (catapleiite-bearing aegirine meta-nepheline syenite;  
295 Fig. 2). This unit is dark bluish and fine- to- coarse-grained with white- to- colourless feldspar  
296 schlieren and contains boudin-shaped pods (up to decimetre size) of mafic rocks not yet  
297 investigated in detail. The rock consists of large and anhedral microcline crystals embedded  
298 or engulfed in fine- to medium-grained bands consisting of albite, clinopyroxene, biotite,  
299 amphibole  $\pm$  fluorite. Most of these minerals are aligned, elongated and anhedral, only  
300 clinopyroxene and amphibole form euhedral crystals.

301

## 302 **MINERAL COMPOSITIONS**

### 303 **Analytical techniques**

304

305 Drill core samples were collected along two drill sections (J and P, nomenclature by Tasman  
306 Metals) at different depths of the intrusion (Fig. 2b and c). The investigated samples were  
307 selected to represent different styles of mineralization and degrees of alteration. Polished  
308 thick sections (120  $\mu\text{m}$ ) have been used for all analyses and a list of analyzed samples is given  
309 in Table 1.

310 Carbon-coated thick sections were studied by SEM-based Mineral Liberation  
311 Analyzer (MLA) and an Electron Probe Microanalyzer (EPMA) at the Helmholtz-Institute  
312 Freiberg for Resource Technology. The MLA studies used an FEI MLA Quanta 650F  
313 scanning electron microscope equipped with a field emission gun and two energy dispersive  
314 X-ray spectrometers (Bruker AXS Xflash® 5010) combined with image analysis software  
315 (Fandrich et al., 2007). Grain-based X-ray mapping (GXMAP) with resolution 500 x 500  
316 pixels by a pixel size of 2µm with 5 pixel step size was applied at an acceleration voltage of  
317 25 kV and a spot size of 5. The data was classified using the software package MLA Suite  
318 3.0.

319 Electron Probe Microanalyzer measurements were performed with a JEOL JXA  
320 8530F equipped with a field emission electron gun and five wavelength-dispersive  
321 spectrometers (WDS). Two of these spectrometers are provided with H-type crystals for high  
322 sensitivity and one with L-type crystals for good energy resolution. The X-ray element  
323 distribution maps were acquired with an acceleration voltage of 20 kV, a beam current of 50  
324 nA, beam diameter of 14 µm and a dwell time of 140 ms. Aluminum, Y, Cl, and Ce were  
325 measured using the WDS, while Si, K, Ca, Mn, Zr, and Fe were measured with an energy  
326 dispersive spectrometer (EDS). Quantitative analyses of EGM were performed at an  
327 accelerating voltage of 20 kV and a beam current of 20 nA applying scanning mode as further  
328 details are described in Atanasova et al. (2015). Clinopyroxene, catapleiite, feldspar,  
329 nepheline and natrolite single point analyses were conducted at an accelerating voltage of 20  
330 kV and a beam current of 20 nA. All further details are given in Electronic Appendix 2. Only  
331 aggregated analytical results are reported in Tables 3-7 and in Electronic Appendix 3. All  
332 detailed data is available on request by the senior author.

333



334 **Clinopyroxene**

335

336 Structural formula calculations for clinopyroxene are based on 4 cations and 6 oxygens,  
337 assuming stoichiometry. Average clinopyroxene analyses are given in Table 3. For  
338 classification we used the International Mineralogical Association (IMA/CNMMN)  
339 nomenclature (Morimoto et al., 1988; Rock, 1990) applying the classification diagram Jadeite  
340 (Jd) – Aegirine (Ae) – Quad (Q) with Quad (Q) being the sum of Ca-Mg-Fe pyroxene  
341 components (enstatite, ferrosilite, diopside and hedenbergite)

342 In the investigated samples, clinopyroxene is invariably sodic with a compositional  
343 range of  $Jd_{2-40}Ae_{45-98}Q_{0-18}$  (Table 3, Fig. 8). Based on their textural appearance (see above),  
344 we distinguish the following five varieties, showing distinct compositions in an aegirine –  
345 (Ti+Zr)aegirine – jadeite ternary (Fig. 5).

346 (1) Resorbed cores (Fig. 5a) of zirconian aegirine (Zr-Ae) with compositions of  $Ae_{68-}$   
347  $_{75}Ti/Zr-Ae_{14-16}Jd_{3-4}$  featuring high Zr (up to 0.06 apfu) but low Ti (<0.02 apfu) contents.

348 (2) Clinopyroxene areas containing anhedral catapleiite inclusions (Fig. 5a), having  
349 compositions of  $Ae_{77-88}Ti/Zr-Ae_{2-7}Jd_{4-10}$  with low Zr and Ti (<0.01 apfu) contents.

350 (3) Subhedral to anhedral, needle- or columnar-shaped clinopyroxene (Fig. 5c)  
351 exhibiting compositions of  $Ae_{76-88}Ti/Zr-Ae_{1-6}Jd_{5-14}$ . Most of these data fall into the field for  
352 aegirine (sensu stricto) with  $Ae \geq 80$ , some of the data indicate calcian aegirine ( $Q \geq 10$ ) and a  
353 notable number of analyzes represent aluminian aegirine ( $Jd > 10$ ; Fig. 8).

354 (4) Titanian aegirine (Ti-Ae) with compositions of  $Ae_{67-87}Ti/Zr-Ae_{3-15}Jd_{6-12}$  exhibiting  
355 low Zr (up to 0.03 apfu) but high Ti (up to 0.06 apfu) occurring as well-defined sectors and  
356 irregular areas. (Fig. 5a-c).

357 (5) Aluminian aegirine (Al-Ae) with compositions of  $\text{Ae}_{56-75}\text{Ti/Zr-Ae}_{1-5}\text{Jd}_{17-35}$  forming  
358 anhedral rims (Fig. 5a-c) and/or irregular patchy areas crosscutting clinopyroxene types (3)  
359 and (4).

360 The frequency of the different clinopyroxene types varies strongly throughout the  
361 complex: In the mid subunit (1)-(5) occur, but in samples from the inner and the outer  
362 subunits (1) and (2) are absent, while (4) is rarely present. While (3) and (5) occur in similar  
363 amounts in the inner unit, in the outer unit (5) predominates over (3) (Fig. 5d).

364 The various clinopyroxene types are not only observed in the different subunits, but in  
365 single crystals too. As illustrated for samples 7315 and 6105 (Fig. 5b and c) from the inner  
366 unit, clinopyroxene (3) is anhedrally overgrown by (5). All five above-defined clinopyroxene  
367 types are present in a single crystal from the coarse-grained, undeformed schlieren (type B) of  
368 the mid subunit (sample 5408; Fig. 5a). Here, a Zr-aegirine core (1) is irregularly overgrown  
369 by aegirine (2) containing numerous anhedral catapleiite inclusions. Titanian-aegirine sectors  
370 (4) and aegirine (3) overgrow this most central part of the crystal. Aluminian-aegirine (5)  
371 anhedrally rims the crystal.

372

### 373 **Eudialyte-group minerals**

374

375 Formula calculations for EGM were carried out by normalizing the sum of (Si + Zr + Ti + Nb  
376 + Al + Hf) to 29 apfu (see details in Pfaff et al., 2010). In total 91 EGM grains from 13  
377 samples were analyzed. The compiled data for each sample including an average for the  
378 particular units is given in Electronic Appendix 3.

379 The textural and compositional diversity of EGM analyzed is illustrated in Table 4 and  
380 Figs 3, 6 and 9. In common with clinopyroxene, four textural varieties of EGM, which show  
381 clear compositional differences are distinguished (Figs 6 and 9):

382 (1) Sector-zoned EGM enriched in Zr and depleted in REE and Nb.

383 (2) Oscillatory-zoned EGM that overgrow EGM (1); slightly enriched in Ca, Fe and  
384 Cl, but depleted in REE.

385 (3) Irregular, up to several tens of micrometers wide areas with bright BSE contrast,  
386 which replace and/or infiltrate EGM (1) and (2) (Fig. 4k). These are enriched in LREE and  
387 show highest #Mn and  $\sum$ REE.

388 (4) Few micrometers wide rims or areas around vugs and/or cracks being enriched in  
389 Y, HREE and #Mn, depleted in LREE with the highest  $\sum$ REEY and Nb.

390 In a given sample, sector and oscillatory-zoned EGM (1) and (2) are comparably REE-  
391 poor and exhibit low #Mn (defined as  $Mn/(Fe+Mn)$ ), whereas types (3) and (4) are REE-rich  
392 and exhibit higher #Mn (Figs 6 and 9).

393 The most significant textural and compositional complexity is observed for EGM from  
394 schlieren of the mid subunit, as was the case for clinopyroxene (see above). Sector-zoned  
395 EGM (1) is observed in samples 7308 and 5618 (Figs. 6a and b), where sectors with high BSE  
396 contrast are richer in Zr, REE and Y compared to sectors with low BSE contrast. These most  
397 central parts of the crystals are overgrown either by oscillatory-zoned EGM (2) with  
398 comparably higher Si, Ca, MREE and HREE but low LREE (Fig. 6b) or by irregular shaped  
399 areas of type (3) with very high BSE contrast being enriched in REE and Y. Higher Y  
400 contents are documented at the rim of the crystals as well as in poikilitic areas of type (4).

401 EGM type (3) mainly occurs in the inner subunit. These are fine-grained, deformed  
402 and texturally less complex crystals with no significant compositional variation (Fig. 6c).  
403 They coexist with clinopyroxene types (3) and (5) (Fig. 5c) and have the highest LREE and  
404 Mn enrichments (Fig. 9). EGM in the outer subunit are fine-grained of types (1) and (2) or  
405 poikilitic type (4) (Fig. 6d). Poikilitic EGM have the lowest LREE content compared to all  
406 EGM studied and show a particular depletion of REE at the rims (Fig. 6d, BSE and Ce map).

407 Furthermore, regional compositional differences in EGM compositions exist for Ca,  
408 Na, Y and the  $\Sigma$ HREE, where a general decrease from the outer to the inner subunit is present  
409 (Fig. 9). The opposite trend is noted for #Mn,  $\Sigma$ REE and  $\Sigma$ LREE, which increase from the  
410 border to the centre of the intrusion (Fig. 9). Niobium (apfu) and Zr/Hf ratios however, do not  
411 show such systematic changes, and Cl contents are mostly relatively low ( $Cl_{apfu} < 0.5$ ), with  
412 one sample containing exceptionally Cl-rich EGM. These observations are in accordance with  
413 observations made by Sjöqvist et al. (2013).

414

### 415 **Catapleiite**

416

417 Structural formula calculations for catapleiite are based on 10 cations and 11 oxygens,  
418 assuming stoichiometry. Representative compositions are given in Table 5 and Fig. 7a.  
419 Catapleiites from two samples of the mid subunit were analyzed. In sample 5408 both textural  
420 varieties of Ca-catapleiite, namely coarse-grained subhedral crystals (Fig. 7b) and fine-  
421 grained anhedral inclusions in clinopyroxene (Fig. 7c) have very similar compositions with up  
422 to 7 wt.% CaO (Table 5). Similarly, in sample 5618 catapleiite occurrences at the border area  
423 between clinopyroxene and EGM and catapleiite inclusions in clinopyroxene (Fig. 7d and e)

424 exhibit quite similar compositions with CaO = 0.3-1.3 wt.% and Na<sub>2</sub>O = 13.0-14.0 wt.%  
425 (Table 5).

426

### 427 **Feldspars, feldspathoids and zeolithes**

428

429 Feldspars occur in all investigated samples as distinct albite and microcline grains and no  
430 perthitic exsolution textures were observed. There are no significant differences between the  
431 various samples and units with albite being almost pure (< 1 mol.% microcline) and  
432 microcline containing up to 5 mol.% albite (Table 6).

433 Nepheline compositions (Ne<sub>74-77</sub>Ks<sub>21-25</sub>An<sub>0</sub>Qz<sub>0-1</sub>) in all samples are very similar (Table  
434 7). According to the diagram after Hamilton (1961), the majority of analyses plot between the  
435 ideal compositions of Morosewicz (M) and Buerger (B) (Fig. 10) and generally fall below the  
436 isotherm at T = 700°C, with two exceptional analyses indicating T > 775°C.

437 Albite and nepheline are variably replaced by zeolithes. Natrolite is largely dominant  
438 and shows a very restricted compositional variation. Analcime was found only in one sample  
439 (Table 7).

440

## 441 **DISCUSSION**

442 The aim of this study is to distinguish the magmatic signature of an agpaitic complex from  
443 features related to subsequent deformation and metamorphism using mineral textures and  
444 compositions. In the course of the study, minerals of the feldspar-group and nepheline were  
445 proven less beneficial probably due to their relatively simple structure and composition. In

446 contrast, structurally and compositionally more complex minerals of the eudialyte-group,  
447 catapleiite as well as clinopyroxene appeared more appropriate to reflect physico-chemical  
448 changes.

449

#### 450 **Discrimination between magmatic and metamorphic features**

451 The agpaitic Norra Kärr complex was subsequently overprinted and deformed during two  
452 successive orogenic events and is now preserved in a well-defined syncline with near parallel  
453 occurrence of major rock units (Fig. 2b und c). In the following we assign the various textures  
454 and compositions of the rock forming minerals to either magmatic relics or metamorphic  
455 features in order to reconstruct the magmatic and the subsequent metamorphic history of the  
456 Norra Kärr rocks.

457

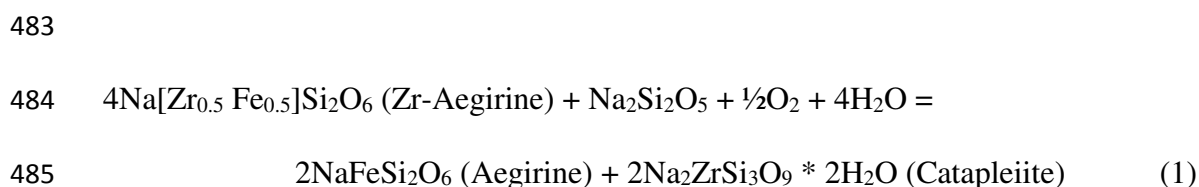
#### 458 Preserved magmatic features

459

460 Feldspars from the Norra Kärr rocks do not exhibit perthite or antiperthite textures preserved  
461 (in agreement with Adamson 1944). We take this as evidence for the crystallization from a  
462 subsolvus (low T) syenite (Sood & Edgar, 1970; Larsen & Sørensen, 1987; Markl, 2001;  
463 Markl & Baumgartner, 2002; Chakrabarty et al., 2016). Alternatively, this could also reflect  
464 the metamorphic recrystallization, but considering the observation that even feldspar  
465 inclusions in EGM and clinopyroxene represent near end member compositions, a primary  
466 subsolvus origin is considered most probable. This hypothesis is further supported by  
467 temperatures derived from nepheline compositions, which generally do not exceed 700°C  
468 (Fig. 10). The two exceptional nepheline compositions that indicate  $T > 775\text{ °C}$  might  
469 represent early magmatic (phenocrystic) relict nepheline.

470 Relics of an early magmatic crystallization stage of the Norra Kärr rocks might be  
471 represented by the Zr-aegirine cores of clinopyroxene (1) (Figs 5a and 7c). At constant P and  
472 T, Zr-aegirine is stable at moderate  $a_{\text{NdS}}$  (activity of  $\text{Na}_2\text{Si}_2\text{O}_5$ ) and under relatively reducing  
473 conditions (Andersen et al., 2012). At this stage clinopyroxene could act as a sink for Zr,  
474 preventing the crystallisation of early magmatic Zr-silicates. Similar relic clinopyroxene  
475 crystals were reported from the deformed Red Wine intrusion (Currie & Curtis, 1976),  
476 resulting from the large stability field of clinopyroxene and its ability to adapt its  
477 composition to the prevailing P-T-X conditions.

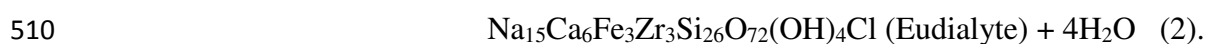
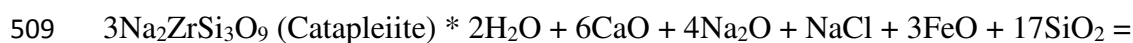
478 Subsequent destabilization of Zr-aegirine (Fig. 7c and d) results in the exsolution of  
479 catapleiite and Zr-poor clinopyroxene (2). The catapleiite inclusions occur in well-defined  
480 areas in clinopyroxene and are characterized by irregular shapes (Figs 5a and 7c). This  
481 process happened probably due to an increase of  $a_{\text{NdS}}$ ,  $a_{\text{H}_2\text{O}}$  or  $f_{\text{O}_2}$  during a later crystallization  
482 stage and can be described by the schematic reaction:



487 This hypothesis was tested by reintegrating the Zr-aegirine composition, using the  
488 molar volumes of catapleiite inclusions contained in well-defined areas of aegirine (2) (Fig.  
489 7d) as well as measured mineral compositions (Tables 3 and 5). Reintegrated Zr-aegirine  
490 compositions correspond very well with measured compositions (Table 8) supporting the  
491 argument for catapleiite formation by Zr-aegirine destabilization and subsequent exsolution.  
492 Considering the textural relationships illustrated in Fig. 7e, we assume that further catapleiite  
493 crystallized prior to EGM, forming euhedral crystals, which were later encased in EGM. In  
494 contrast, the textural relationships illustrated in Fig. 7d provide evidence that both minerals

495 may have replaced each other. In addition, it is not clear if the above-described catapleiite  
 496 exsolutions in clinopyroxene and coarse, euhedral catapleiite intergrown with EGM represent  
 497 different stages during the magmatic evolution of these rocks. Indeed, they may be two  
 498 different generations. Compositionally, however, there are no differences between these two  
 499 types of catapleiite (Fig. 7a). In general, both catapleiite and EGM occur in Zr-rich  
 500 peralkaline magmatic systems. However, the formation of magmatic EGM seems to require  
 501 relatively low water activities, but high Cl activities, conditions that are typically met in  
 502 reduced nepheline syenitic systems, whereas magmatic catapleiite seems to be restricted to  
 503 peralkaline (granitic) systems being relatively H<sub>2</sub>O-rich (e.g., Andersen et al. 2010; Marks et  
 504 al., 2011; Andersen & Friis, 2015). Note that catapleiite can also form (along with many other  
 505 phases) during the hydrothermal alteration of EGM (e.g., Salvi et al., 2000; Karup-Møller et  
 506 al., 2010; Karup-Møller & Rose-Hansen, 2013; Borst et al., 2016). The catapleiite-eudialyte  
 507 equilibrium can be expressed by the following schematic reaction:

508



511

512 The Cl contents in EGM from the Norra Kärr rocks mostly vary between 0.2-0.4 apfu  
 513 (Fig. 9a) and are relatively low compared to EGM in other alkaline complexes with Cl<sub>apfu</sub> up  
 514 to 1.9 (see Schilling et al., 2011b). EGM from Norra Kärr may contain up to about 3 wt.%  
 515 H<sub>2</sub>O (Atanasova et al., 2015). This implies that EGM may be stable even at relatively-low  
 516 a<sub>HCl</sub> but relatively-high a<sub>H2O</sub>. We propose that in the case of Norra Kärr a<sub>H2O</sub> was not high  
 517 enough to stabilize catapleiite over a long crystallization interval. In addition, initial



518 catapleiite formation lead to subsequent REE enrichment of the melt, which probably was the  
519 crucial promoting factor stabilizing EGM in this relatively Cl-poor environment.

520 On the deposit scale, we observe a continuous decrease in Ca, Na, Y and the HREE  
521 with increasing #Mn,  $\Sigma$ REE and LREE from the outer to the inner subunit as demonstrated in  
522 Fig. 9. Such compositional changes of EGM are attributed to primary magmatic evolution  
523 (Sjöqvist et al., 2013) also known from EGM-bearing layered intrusions (e.g. Pfaff et al.,  
524 2008; Schilling et al., 2011a; Sheard et al., 2012; Lindhuber et al., 2015; Ratschbacher et al.,  
525 2015; Möller & Williams-Jones, 2016). These studies report compositional evolution in terms  
526 of the Mn/(Fe+Mn) ratio and the content of the different REE, which is considered to be the  
527 result of fractional crystallization during primary, magmatic layering of agpaitic intrusions  
528 (Fig. 11). Early magmatic EGM are enriched in Fe and the HREE, which leads to subsequent  
529 Mn- and- LREE-enrichment of the residual melt and to crystallization of late magmatic Mn-  
530 and LREE-dominated EGM. Similarly, primary layering of the Norra Kärr complex (Fig. 11),  
531 is indicated by early Fe- and- HREE-dominated EGM from the outer subunit as well as by  
532 continuous Mn and LREE enrichment in the mid and inner subunits (Fig. 11) and are still  
533 preserved within the fold structure.

534

535

536 Textures assigned to metamorphic overprint

537

538 The magmatic textures described above such as sector and oscillatory zoning in  
539 clinopyroxene and EGM are partly-overgrown and/or replaced by late anhedral Al-aegirine  
540 and REE-enriched EGM, respectively. The formation of Al-aegirine is a general feature of

541 metamorphosed syenites (Wooley et al., 1996) and is probably formed by the following  
542 schematic reaction (Curtis & Gittins, 1979):

543



545

546 Same reaction is observed in samples studies from Norra Kärr as well. According to  
547 (Curtis & Gittins, 1979) equation (3) could take place in a huge P- and- T-range of conditions,  
548 between 450-700°C and 2-10 kbar, covering both, higher greenschist and lower- to- moderate  
549 amphibolite facies conditions. In contrast to Red Wine, where amphibolite facies conditions  
550 are reported, at Norra Kärr moderate- to- high greenschist facies conditions are only roughly  
551 estimates. These estimates are supported by the observation that relic magmatic textures are  
552 preserved due to assumed relatively low P and T during deformation of the rocks (Wooley et  
553 al., 1996).

554 In EGM, flamy BSE textures (Figs 4k, 6a and b) as well as very late Y enrichments  
555 and LREE depletions at the rim of crystals, along cracks, vugs (Figs 6a, b and d) and in highly  
556 poikilitic areas (Figs 6b and d) are observed. Flamy textures as shown in Fig. 4k indicate  
557 both, new formation of compositionally different EGM of type 3, (straight boundaries  
558 between EGM (1) and (3)) as well as recrystallization or replacement of EGM types (1)  
559 and/or (2) by EGM of type (3) (flamy EGM (3) permeating EGM (1) and/or (2)). Both can be  
560 explained by the interaction with fluids, presumably generated during the metamorphic  
561 overprint of the rocks. Yttrium and HREE enrichments accompanied by LREE depletion at  
562 the rims of crystals (Fig. 6a, b and d) might document an even later event causing further  
563 compositional adaptation of EGM, in this case accompanied by the re-distribution of REE and  
564 Y, spatially-associated with the formation of late catapleiite and secondary LREE-rich

565 minerals (Fig. 6a, Ce map). We suggest that these textures could potentially be caused by  
566 recrystallization of EGM in response to metamorphic deformation.

567         The widespread alteration of albite and nepheline to natrolite (Fig. 3) is also a late  
568 process, probably related to fluid-assisted overprint during deformation. Importantly,  
569 analcime is only very rarely observed (actually in only one sample, see above). This  
570 corresponds to observations from the Mont Saint-Hilaire complex and Shushina Hill  
571 (Schilling et al., 2011a; Chakrabarty et al., 2016, respectively) and it was suggested that  
572 under such conditions analcime is only stable at low  $T < 150^{\circ}\text{C}$ . Accordingly, this argues for  
573 higher temperatures for natrolite formation. In the case of Norra Kärr a further indication for  
574 the “metamorphic” origin is given by the spatial distribution of natrolite alteration, which  
575 preferably occurs in the mid and inner units, while the very fine-grained rocks from the outer  
576 unit are less affected, although these rocks are in direct contact with the granitic country rock.

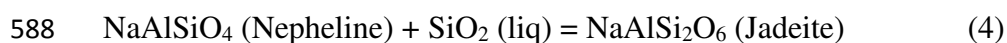
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### 578 **Thermodynamic modelling to constrain the p-T-conditions**

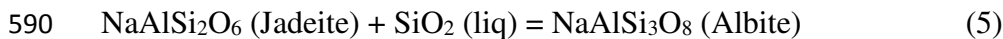
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580 The albite – nepheline – aegirine assemblage of the Norra Kärr rocks permit estimations of  
581 temperature and silica activity at the assumed pressure of the magmatic stage (e.g. Mitchell &  
582 Platt, 1978; Markl et al., 2001, Schilling et al., 2011a) and may shed light into the so far-  
583 poorly constrained conditions during metamorphic overprint during the Sveconorwegian  
584 (Grenvillian) orogeny. We performed thermodynamic calculations applying the GEOCALC  
585 software of Berman et al. (1987) and Liebermann & Petrakakis (1990) with the database of  
586 Berman (1988) using the following three schematic equilibria.

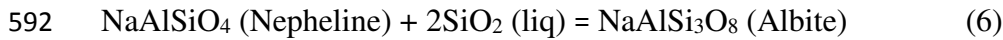
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591



593

594 End member activities were calculated using a one-site mixing model for nepheline  
595 and applying the solution activity formulations of Green et al. (2007) and Holland & Powell  
596 (2003) for clinopyroxene and feldspar, respectively.

597 Assuming typical emplacement pressures of 1-2 kbar for syenitic peralkaline  
598 intrusions (e.g., Konnerup-Madsen & Rose-Hansen, 1982; Salvi et al., 2000) and using the  
599 mean compositions of Zr-aegirine (1), which we interpret as early magmatic clinopyroxene  
600 (see above), equilibrium conditions of 600 - 700°C at  $a_{\text{SiO}_2}$  values of 0.5-0.6 are indicated  
601 (Fig. 12a). Such conditions are consistent with estimations for other peralkaline complexes  
602 and are in agreement with the absence of perthitic alkali feldspar (see above) and the  
603 temperature estimates based on nepheline thermometry (Fig. 10). Compositions of  
604 presumably late-magmatic aegirine (2) and (3) indicate equilibration to lower temperatures (<  
605 450°C) and  $a_{\text{SiO}_2}$  values of < 0.3 (Fig. 12a).

606 Taken together, we interpret these data as representing the magmatic evolution of the  
607 Norra Kärr rocks, indicating a relatively low liquidus temperature of <700°C and a relatively  
608 extensive crystallization interval down to temperatures < 400°C, which has been  
609 demonstrated before, based on both field studies (Markl et al., 2001; Marks et al., 2003;  
610 Marks & Markl, 2003) and experimental work (Piotrowski & Edgar, 1970; Sood & Edgar,  
611 1970; Giehl et al., 2013; 2014).

612 To constrain the metamorphic conditions of the Norra Kärr rocks during the  
613 Sveconorwegian orogeny, we used the mean compositions of Al-aegirine (clinopyroxene 4;

614 see above) computed at pressures of 2.5, 5 and 8 kbars to represent general greenschist as well  
615 as lower amphibolite facies conditions (Fig. 12b). The calculated equilibrium temperatures  
616 range from 350 to 650°C with  $a_{\text{SiO}_2}$  values from 0.55 to 0.15, with both T and  $a_{\text{SiO}_2}$  decreasing  
617 with decreasing pressure. Comparing these estimates with proposed moderate to high  
618 greenschist facies conditions, temperatures from 400-550°C at pressure of 5 kbar with an  $a_{\text{SiO}_2}$   
619 range between 0.25 and 0.4 are considered as most reasonable. These values are similar to  
620 those estimated for late magmatic conditions. This might indicate that  $a_{\text{SiO}_2}$  during the  
621 metamorphic stage did not essentially change, suggesting that fluid-rock interaction during  
622 this stage was largely (locally) rock-buffered and that the responsible fluids were not  
623 especially SiO<sub>2</sub>-rich, as there is no late-stage quartz formed as might be expected from fluids  
624 derived from granitic country rocks.

625 .

## 626 CONCLUSIONS

627

628 Even after metamorphic overprint and deformation, it remains possible to identify primary  
629 magmatic textures and compositions in EGM and clinopyroxene from peralkaline rocks and  
630 to find evidence for primary magmatic layering of the intrusion with continuous enrichment  
631 of REE, LREE and Mn from the outer to the inner subunit. The multi-stage evolution of such  
632 rocks can be reconstructed using clinopyroxene and EGM, as both minerals cover a large  
633 stability range (in term of P and T) and are able to adapt their composition to changing  
634 conditions, thereby recording the magmatic and metamorphic history of the complex.

635 The minerals investigated probably crystallized from a subsolvus syenite and  
636 crystallization took place during a relatively extensive temperature interval from about 700°C  
637 to 450°C with a correspondent decrease in  $a_{\text{SiO}_2}$  from 0.6 to about 0.3. Initially, Zr-aegirine

638 crystallized at relatively low peralkalinity and reducing conditions. Due to an increase in  $a_{\text{Nd}_2\text{O}_3}$   
639 and at moderate- to- high  $f_{\text{O}_2}$  and  $a_{\text{H}_2\text{O}}$ , Zr-aegirine was destabilized and crystallization  
640 continued forming aegirine (2) of sensu stricto compositions and catapleiite as the main Zr-  
641 bearing silicate mineral. Due to the absence of sodalite and the very early formation of  
642 catapleiite,  $a_{\text{HCl}}$  further increased and REE and HFSE were enriched in the residual melt,  
643 promoting EGM crystallization. Crystallization of EGM seems not necessarily restricted to  
644 specific conditions, forming even at very low  $a_{\text{HCl}}$  and at relatively high  $a_{\text{H}_2\text{O}}$ , but to the  
645 availability of specific elements in the melt.

646         During the Sveconorwegian/Grenvillian orogeny, the Norra Kärr rocks were deformed  
647 and folded at presumably moderate greenschist facies conditions (Fig. 12) and late EGM and  
648 Al-aegirine formed at 400°C - 550°C and in a  $a_{\text{SiO}_2}$  range of 0.25-0.4 at pressures not  
649 exceeding 5 kbar. We suggest that during this deformation event EGM interacted with  
650 metamorphic fluids, either changing its composition or being partly-destabilized to catapleiite  
651 and secondary REE-bearing minerals. During the whole history of the complex,  $a_{\text{SiO}_2}$  remains  
652 very similar, indicating very little interaction with the surrounding granitic rocks. This  
653 assumption is confirmed by the relatively-restricted occurrence of natrolite alteration,  
654 preferably occupying the more central part of the intrusion.

655

## 656 **ACKNOWLEDGEMENTS**

657

658 Research for this study was supported by and benefited from numerous discussions with Max  
659 Frenzel, Axel Sjöqvist, Johan Berg and Magnus Leijd. Bernhard Schulz assisted with the  
660 classification of meta-nepheline syenites and Kai Bachmann performed the high resolution  
661 BSE images. The authors are deeply grateful for their support. Furthermore, the authors are

662 indebted to Tasman Metals Ltd. for supporting field work, providing rock samples, geological  
663 maps and cross sections. The manuscript was improved by the reviews of Roger Mitchell and  
664 Stefano Salvi. Their comments and suggestions are greatly appreciated. The authors thank A.  
665 Skelton for the editorial handling.

666

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Fig.1. (a) Simplified geological map of Fennoscandia illustrating major lithological units (modified after Högdahl et al., 2004). Red box is enlarged in (b), where the red ellipse resembles the Norra Kärr complex. The white arrow indicates the direction of decreasing regional deformation during the Sveconorwegian orogeny. PZ-Protogine Zone, SFDZ - Sveconorwegian Frontal Deformation Zone (simplified after Sönderlund et al., 2002; Wahlgren et al., 1994).

Fig.2. (a) Simplified geological map of the Norra Kärr complex illustrating major lithological units (modified after Tasman Metals Ltd). The deformation is well visualized in W-E cross sections (b) J and (c) P. The sampling locations as given in Table 1 are marked as black crosses.

Fig.3. Mineral Liberation Analyzer (MLA) classified-color images (EDX-GXMAP mode) of the different subunits of Norra Kärr. Illustrated are characteristic textures of a) the incipient alignment and stretching of minerals and b) the macroscopic scale folding and alignment of minerals in the inner subunit; (c) poikilitic EGM in a fine grained matrix of the outer subunit; (d) homogeneous matrix of the mid subunit; (e)-(h) thoroughly preserved euhedral to subhedral EGM and clinopyroxene in leucocratic schlieren of the mid subunit. Nepheline is partly to strongly altered to natrolite. Areas marked as red boxes are enlarged in Figs 4, 5 and 6.

Fig.4. (a) Alignment of minerals as well as (b) and (c) microscopic- to macroscopic scale folding as common features of the inner subunit; EGM crystals in (a) are slightly stretched. (d) Folding of the fluoro-leakeite-aegirine meta-nepheline syenite rock. (e) Fine-grained meta-nepheline syenite with a band of mid- to coarse-grained nepheline and eudialyte. The nepheline crystals show alteration. (f) Microcline  $\delta$ -clast, (g) catapleiite and (h) EGM (type (4)) porphyroclasts are characteristic for the outer subunit. EGM contain inclusions of catapleiite, feldspar, nepheline and clinopyroxene. Catapleiite (Cat) forms tails and replaces EGM mainly at the rim. EGM are surrounded by columnar rinkite-group minerals (Rink). (i) Medium-grained schlieren of the mid subunit with feldspar, aegirine and eudialyte show no particular alignment, aegirine crystals (CPX) form radiating aggregates. (j) Coarse-grained microcline and extensively altered nepheline from schlieren of the mid subunit. (k) EGM (3) overgrow and/or replace the compositionally different EGM (1) in schlieren of the mid subunit; visible white spots represent EPMA measurement locations.

Fig.5. Compositional and textural characteristics of clinopyroxene from the Norra Kärr rocks. (a) Euhedral clinopyroxene from the mid unit (sample 5408). The resorbed Zr-aegirine core is irregularly rimmed by aegirine s.s. rich in catapleiite inclusions. Both are overgrown by alternating sectors of inclusion-free titanite-aegirine and aegirine s.s. Anhedral Al-aegirine rims irregularly the crystal. (b) Subhedral clinopyroxene from the mid subunit exhibiting well

defined Ti-rich sectors overgrown by Al-aegirine. In the center a large catapleiite is enclosed. (c) Subhedral to anhedral clinopyroxene of the inner subunit; (d) Anhedral clinopyroxene from the outer subunit. See also detailed descriptions in the main text.

Fig.6. Compositional and textural characteristics of EGM from the Norra Kärr rocks. At the top of the diagram, textural varieties 1-4 from all EGM investigated are collated considering the #Mn and the REEY<sub>apfu</sub>. Representative grains of the different units are shown in (a)-(d). In the mid subunit (a) single EGM crystal (sample 5618) shows sectors (1) being Zr-rich, but REE-poor, overgrown by an irregular flamy area (3) of high BSE contrast, rich in REE and Y. Highest Y is detected along rims and cracks (4). The crystal is closely associated with catapleiite (Cat) and britholite group minerals (Brit). Red box is enlarged in Fig. 4k. (b) Subhedral EGM featuring sector (1) and oscillatory (2) zoning, as well as irregular flamy textures (3) and poikilitic Y-rich areas (4). (c) EGM crystals from the inner subunit commonly show deformation features like stretching and alignment and are high in #Mn and REE. (d) Poikilitic EGM porphyroclast from the outer unit showing LREE depletion and Y enrichments at the rim. Catapleiite (Cat) and rinkite group minerals (Rink) are visible as white areas in the Zr and Ce maps. Visible spots in (a) and (b) represent EPMA (white) and LA-ICP-MS (dark) measurement locations.

Fig.7. Compositional (a) and textural (b-f) characteristics of catapleiite from the Norra Kärr rocks. (b) A single euhedral Ca-catapleiite crystal (Ca-Cat) and (c) Ca-catapleiite inclusions (Ca-Cat) in clinopyroxene, aegirine s.s. (CPX) (Fig. 4a) from schlieren of the mid subunit, sample 5408; (d) and (e) show characteristic catapleiite (Cat) and EGM textures in the mid subunit, sample 5618. Zones in radial aegirine s.s. (CPX), rich in catapleiite inclusions (Cat) are remarkably well defined. Catapleiite occurs enclosed in EGM, between and within EGM crystals and at the grain boundaries between clinopyroxene and EGM. (f) Catapleiite aggregates (Cat) from the outer subunit, sample 5611. Visible spots in (b) and (c) represent EPMA (white) and LA-ICP-MS (dark) measurement locations.

Fig.8. (a) Classification diagram for Ca-Na and Na pyroxenes with the endmembers Jadeite (Jd) – Aegirine (Ae) – Quad (Q) modified after Morimoto et al. (1988) and Rock (1990). The marked area is enlarged in (b) showing that pyroxene compositions from Norra Kärr include aegirine s.s., zirconian aegirine, calcian aegirine, titanian aegirine and aluminian aegirine.

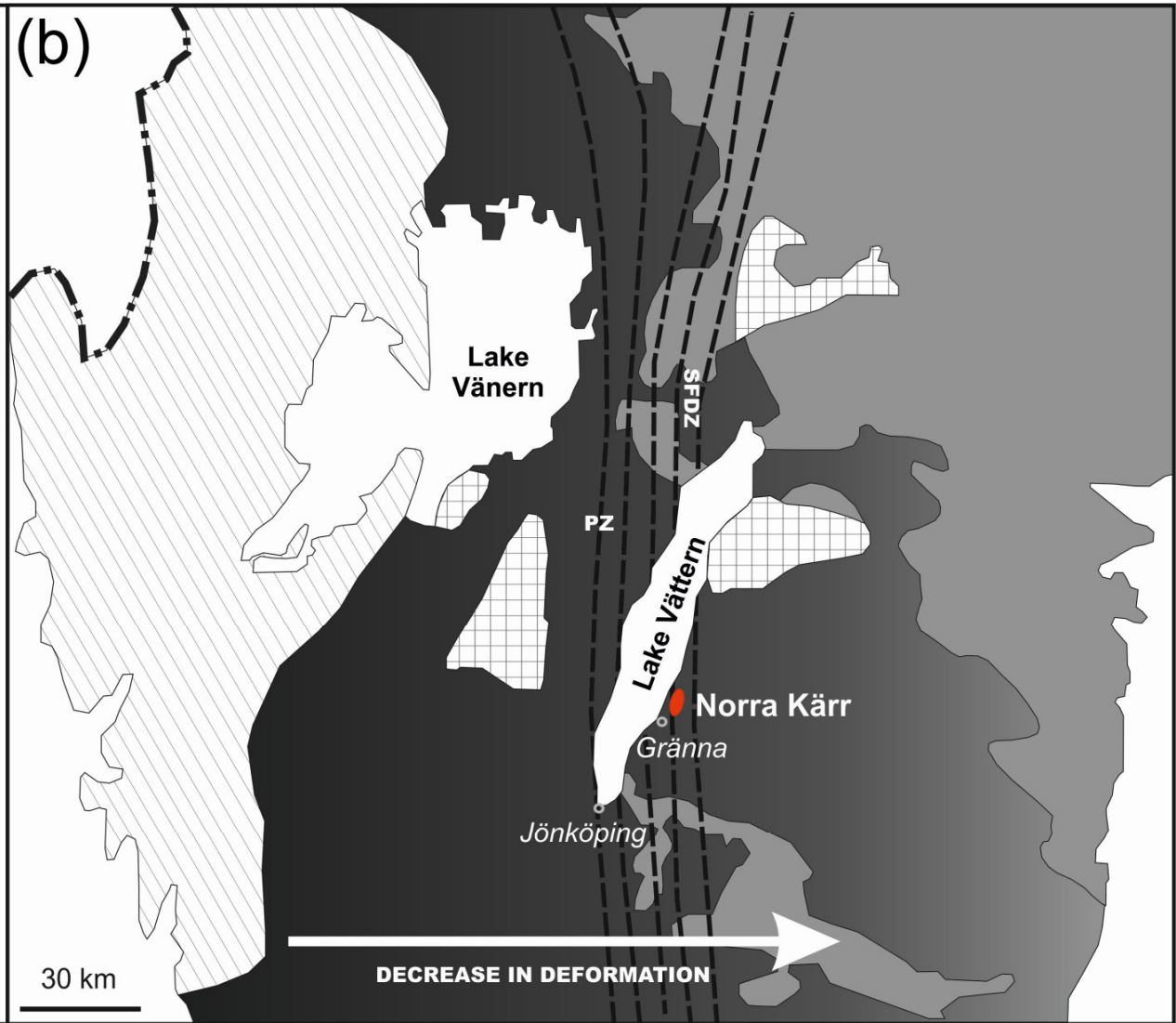
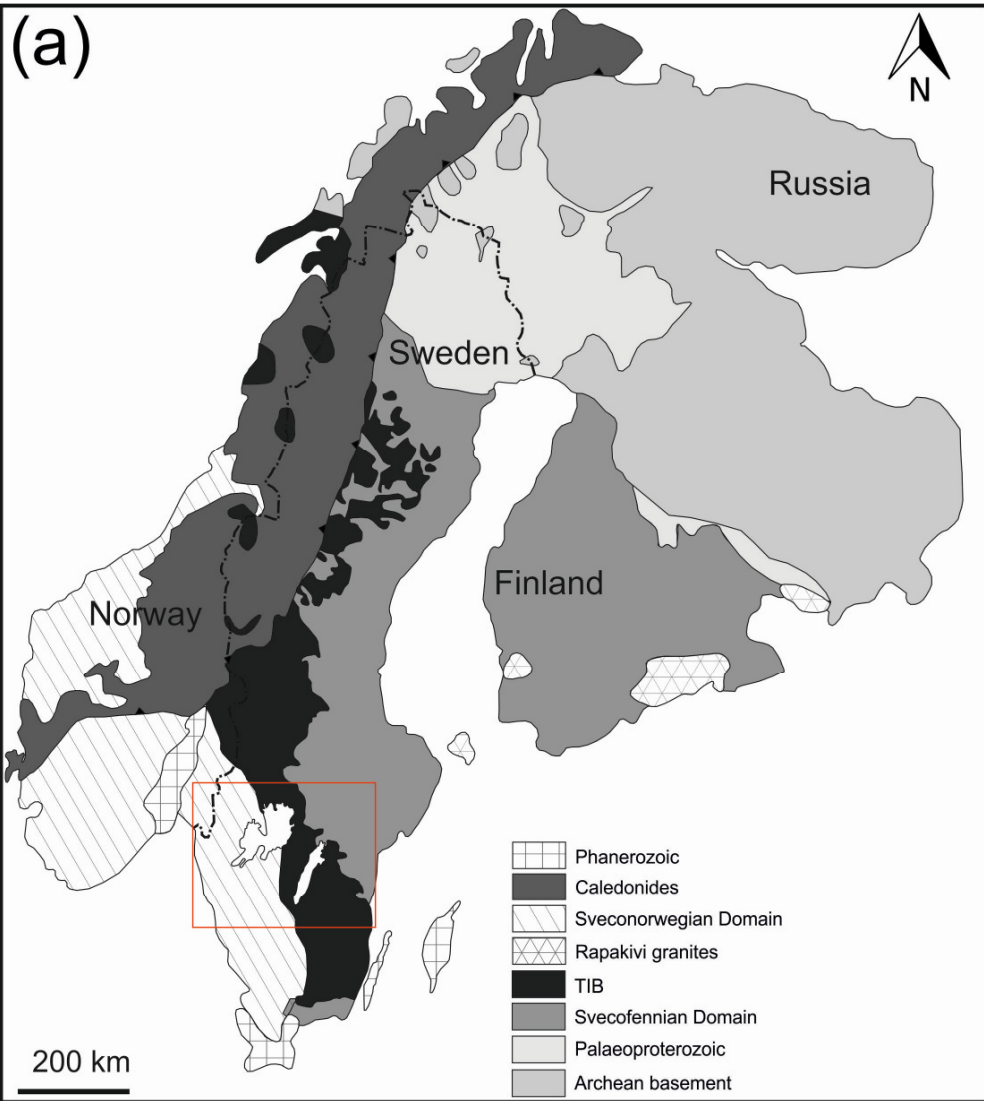
Fig.9. Distribution of (a) major and (b) minor elements in EGM according to the relative sampling location within the fold structure. Samples from the discovery outcrop are displayed for comparison. Colour coding displays EGM textural varieties 1-4 as defined in Fig. 5.

Fig.10. Diagram of the system nepheline-kalsilite-silica (wt.%) with isotherms showing the limit of nepheline solid solution at the indicated temperatures (after Hamilton 1961). The majority of analyses plot within the Morosewicz (M) – Buerger (B) convergence field (after Tilley, 1954). Most of the analyses indicate  $T \leq 500^{\circ}\text{C}$ . Only two analyses indicate  $T \geq 700^{\circ}\text{C}$ .

Fig.11. Suggested primary magmatic layering of the intrusion with upwards enrichment of REE, LREE and #Mn, preserved within the present fold structure.

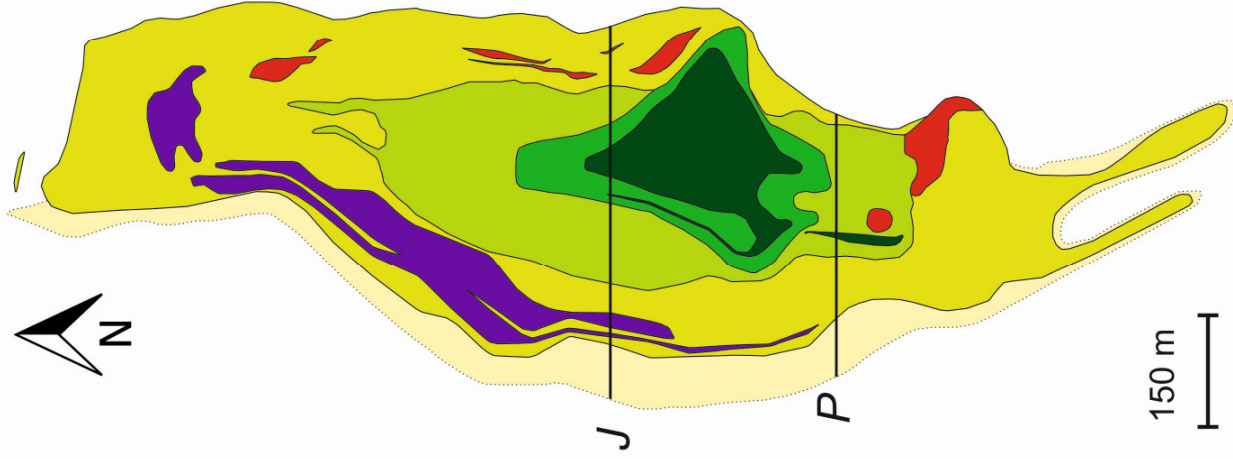
Fig.12. Two silica activity ( $a_{\text{SiO}_2}$ ) - temperature diagrams calculated based on equilibria (4)-(6). Different sets of conditions (different lines in (b)) are calculated due to variable activities of nepheline ( $a_{\text{Ne}}$ ) and jadeite ( $a_{\text{Jd}}$ ), while the activity of albite ( $a_{\text{Ab}}$ ) is 1 for all reactions. (a) Magmatic conditions are estimated for 1-2 kbar using early- and late-magmatic clinopyroxene compositions, respectively. (b) The metamorphic evolution of the Norra Kärr complex is illustrated for pressures representing general greenschist (grey area) to lower amphibolite facies conditions, considering metamorphic clinopyroxene textures and compositions. Details are given in the main text.

Electronic Appendix 1. Selected scans of core samples. (a) Folded fluoro-leakeite-aegirine meta-nepheline syenite. (b) Catapleiite-and EGM-bearing aegirine meta-nepheline syenite with 5 % pegmatoidal schlieren shows folding of the rock. (c) Catapleiite-and EGM-bearing aegirine meta-nepheline syenite with 30 to 50 % pegmatoidal schlieren with small scale folding of the schlieren, minerals show hematization. (d) Foliated fine- to- medium-grained aegirine leuco-meta-nepheline syenite. (e) Coarse-grained schlieren with microcline, albite and EGM, fine-grained euhedral aegirines form patchy aggregates.



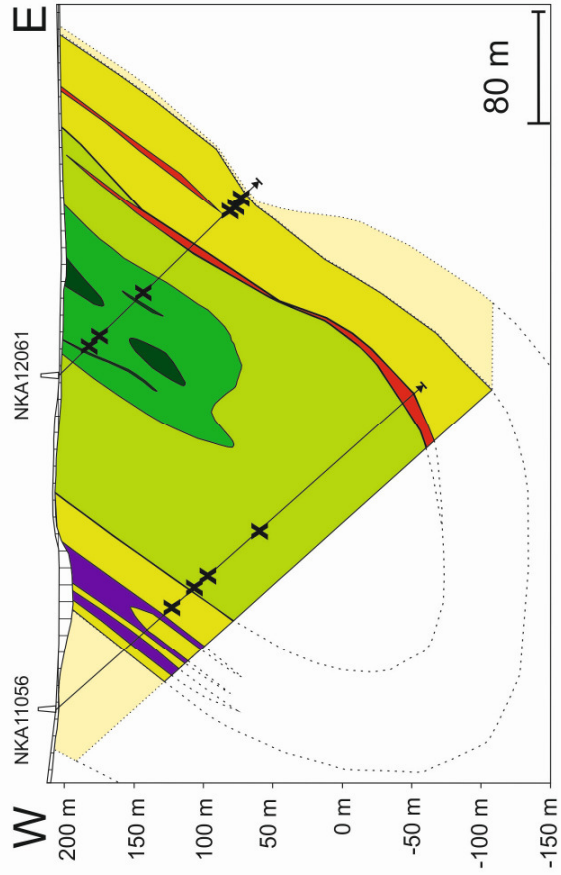


(a)



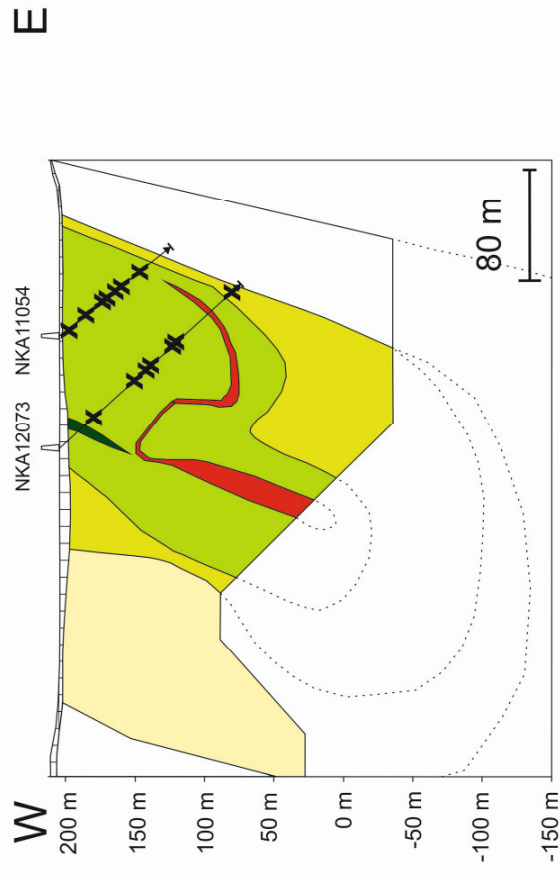
(b)

Section J



(c)

Section P

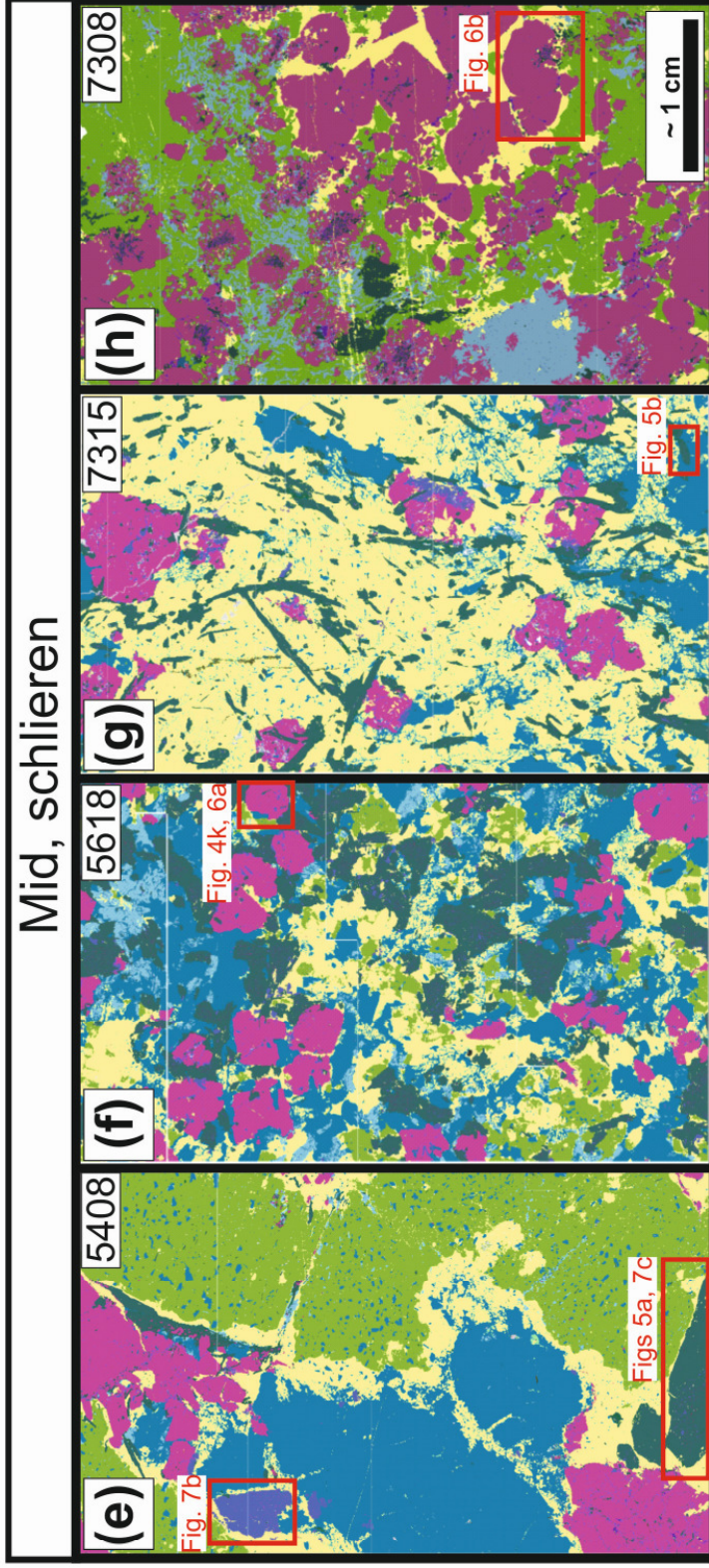
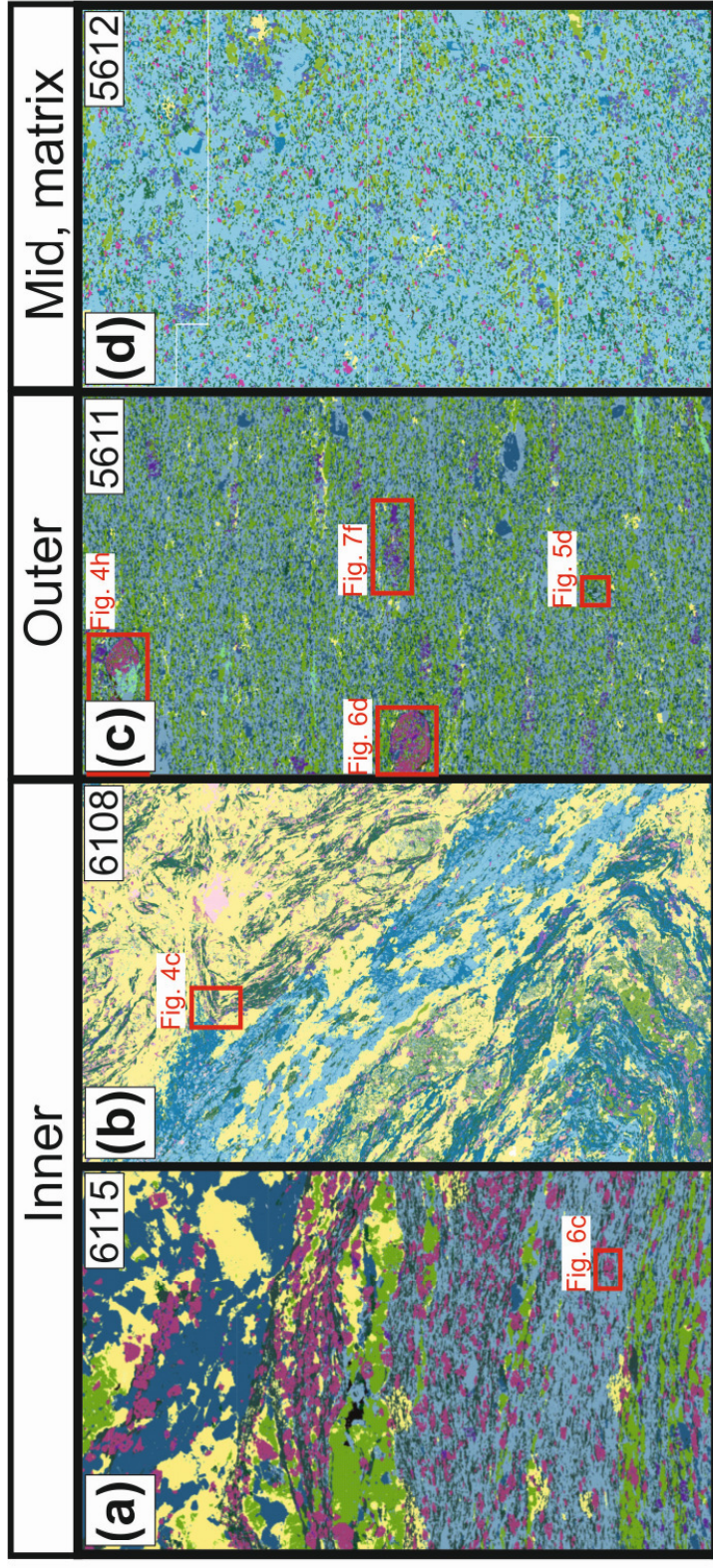


### Legend

- Outer subunit: Catapleite-bearing aegirine meta-nepheline syenite
- Mid subunit: Pegmatite-bearing aegirine meta-nepheline syenite
- Inner subunit: Aegirine leuco-meta-nepheline syenite
- Discovery: Arfvedsonite-aegirine meta-nepheline syenite
- Fluoro-leakeite-aegirine meta-nepheline syenite
- Aegirine-amphibole-biotite meta-nepheline syenite

- Overburden
- Granitic- to- syenitic host rock
- Host rock, fentitized
- Drill core
- Approximate sampling location



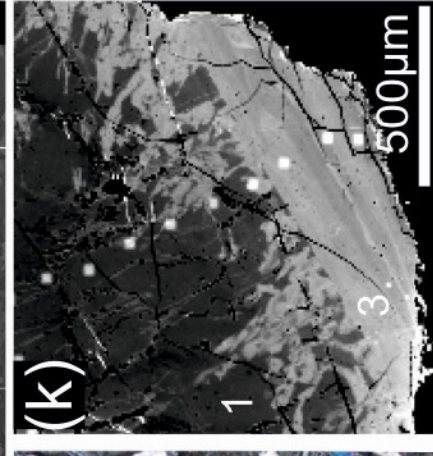
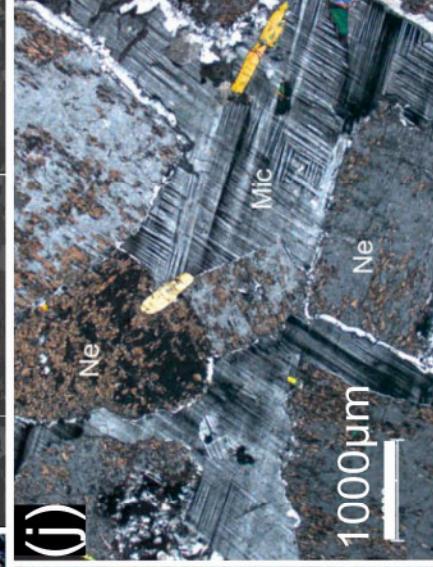
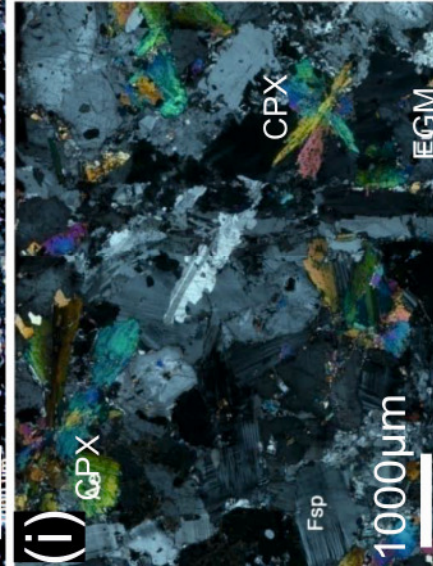
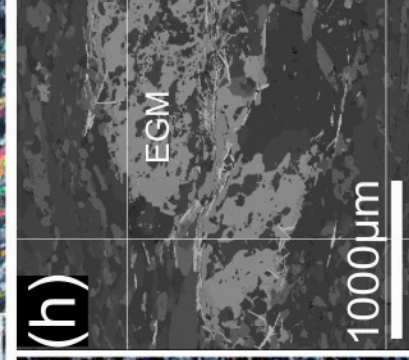
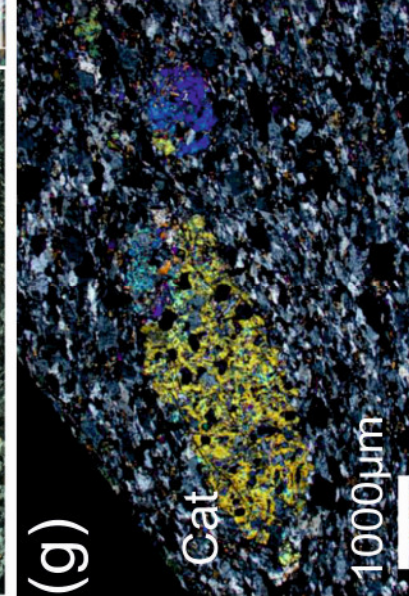
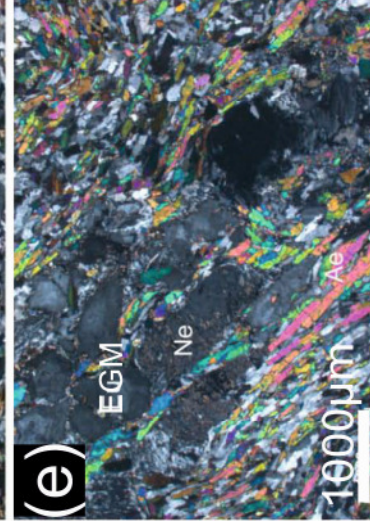
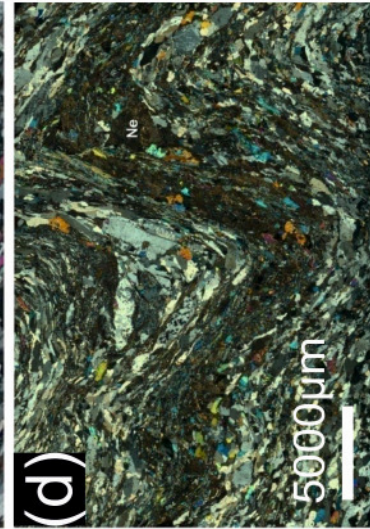
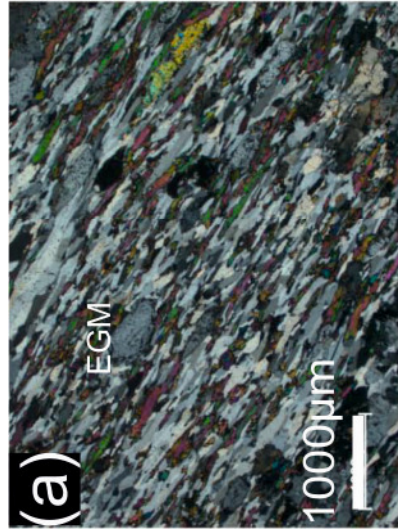


### Legend

- Aegirine
- Catapleiite
- Eudialyte group minerals

- Albite
- Microcline

- Nepheline
- Natrolite



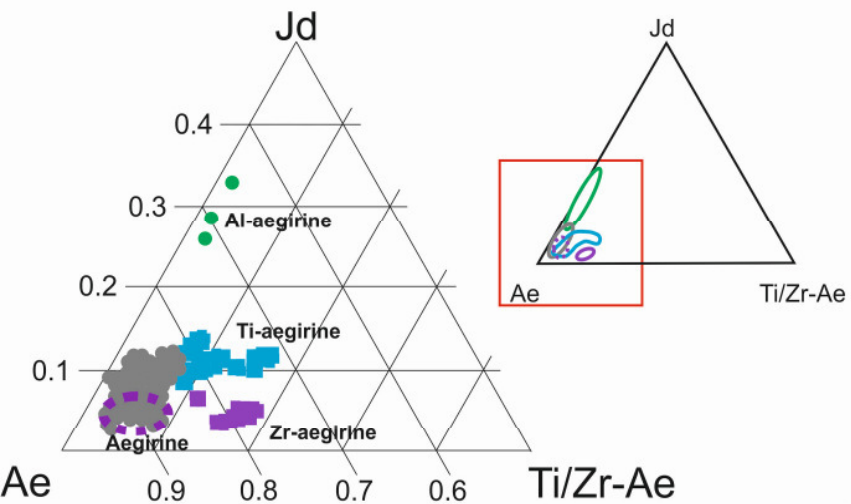
## Mid

## Inner

## Outer

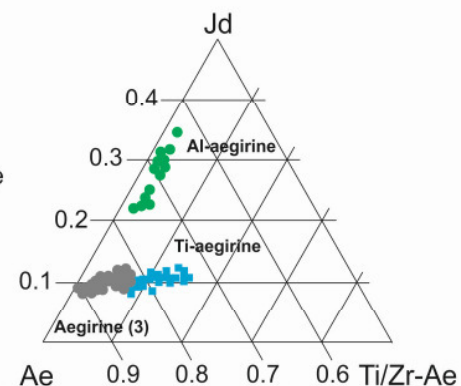
(a)

NKA5408



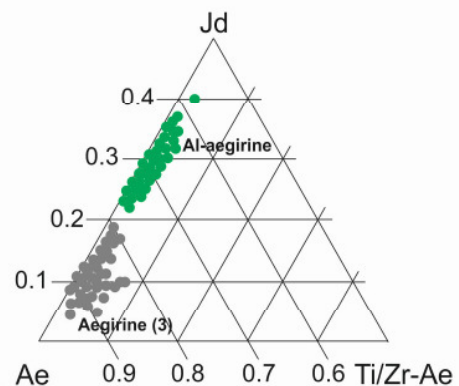
(b)

NKA7315



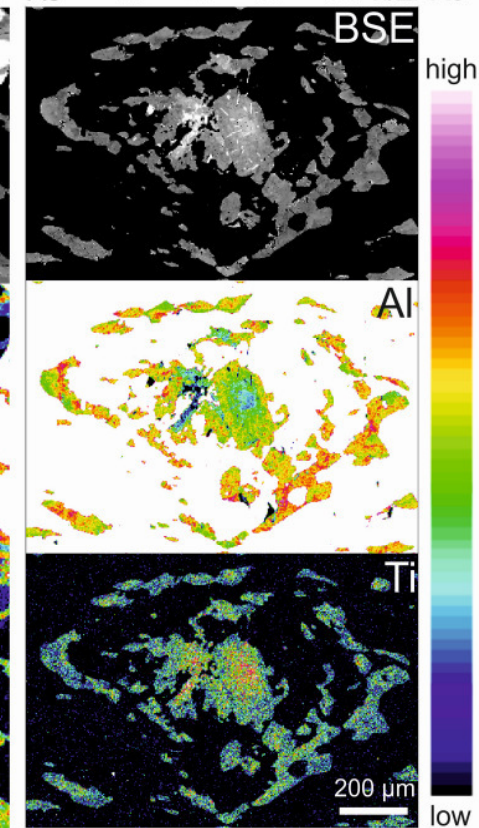
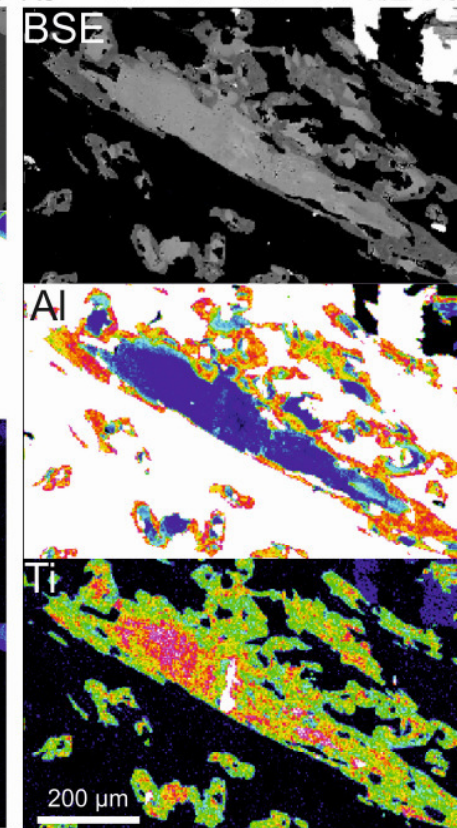
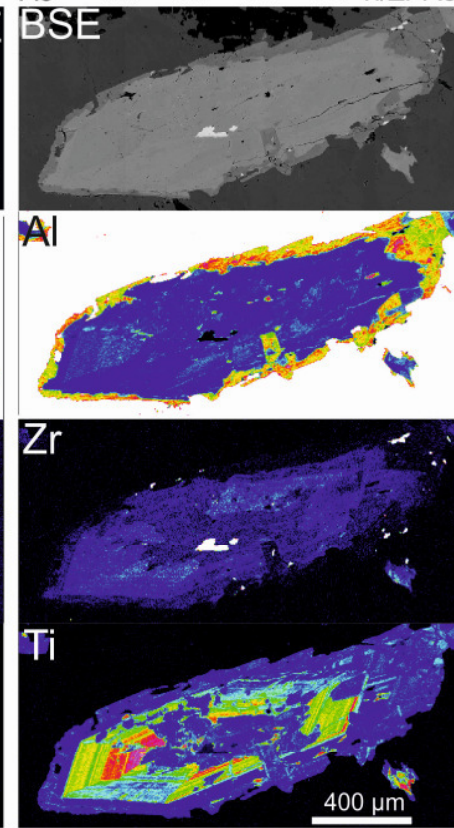
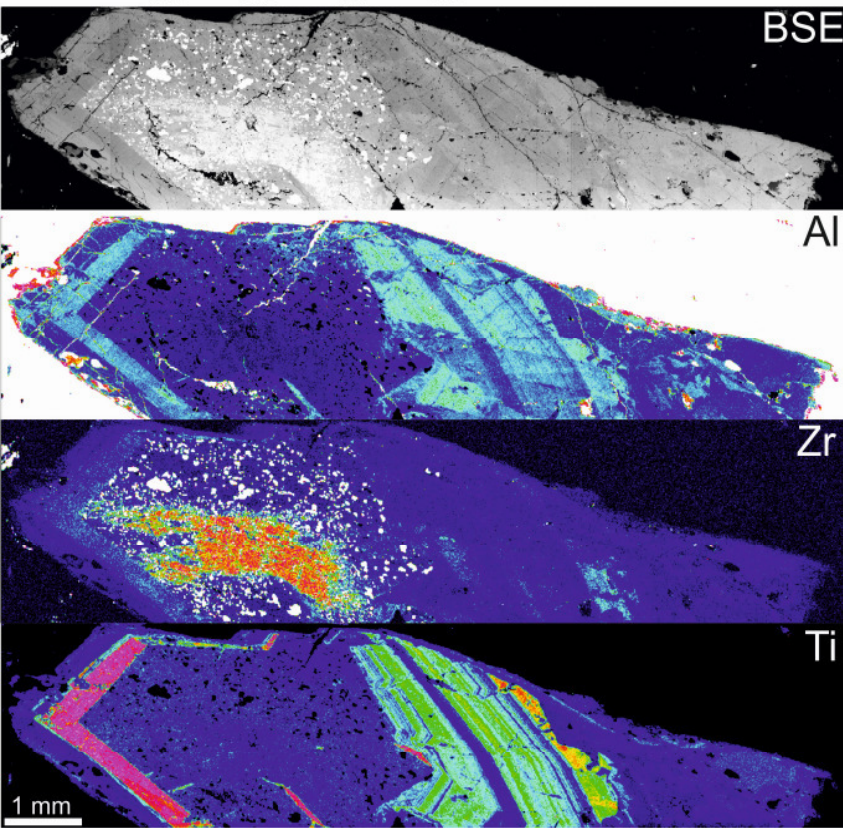
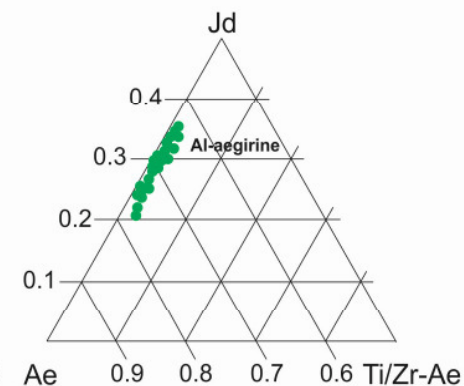
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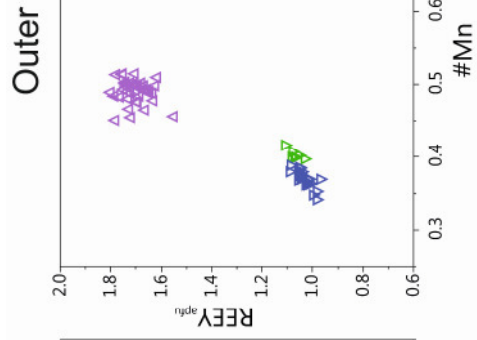
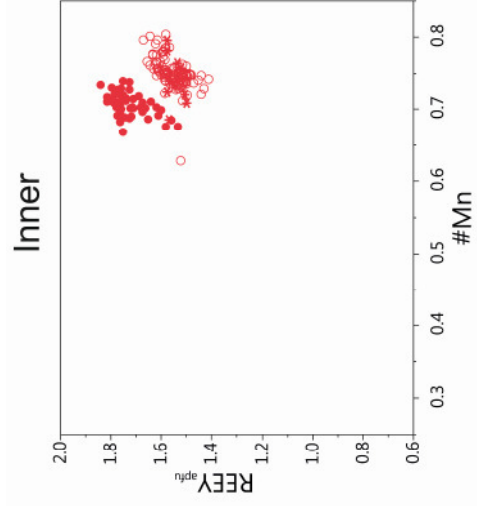
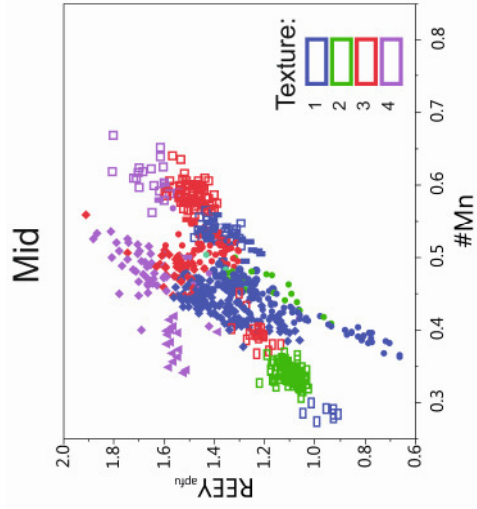
NKA6105



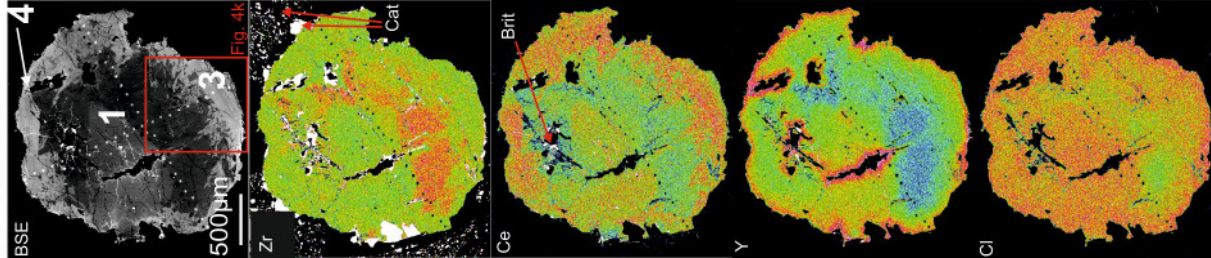
(d)

NKA5611

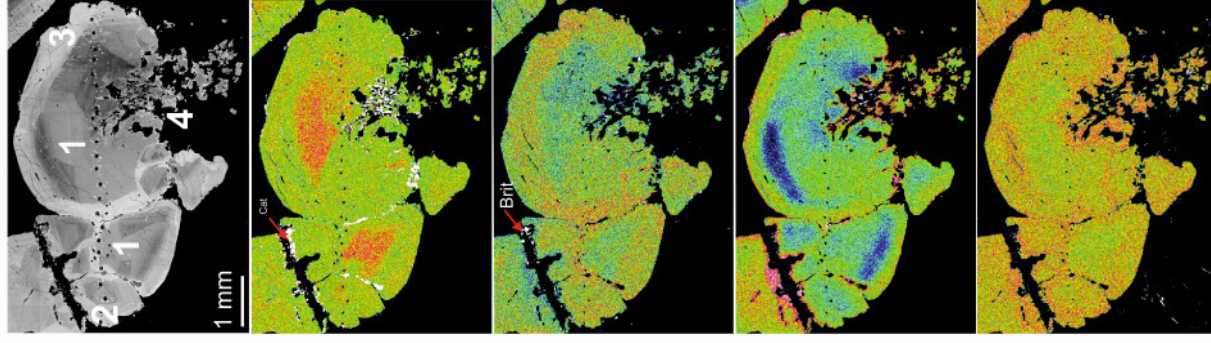




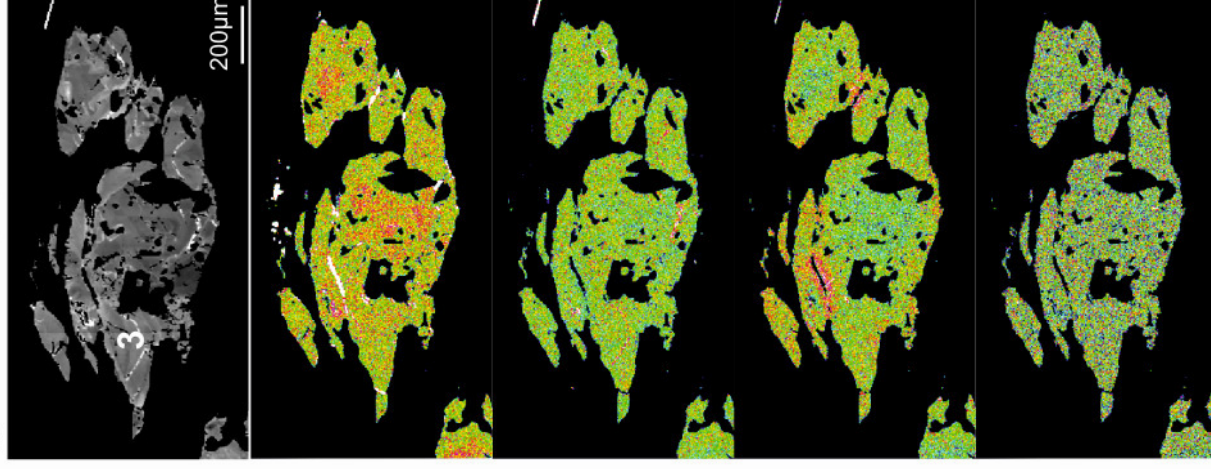
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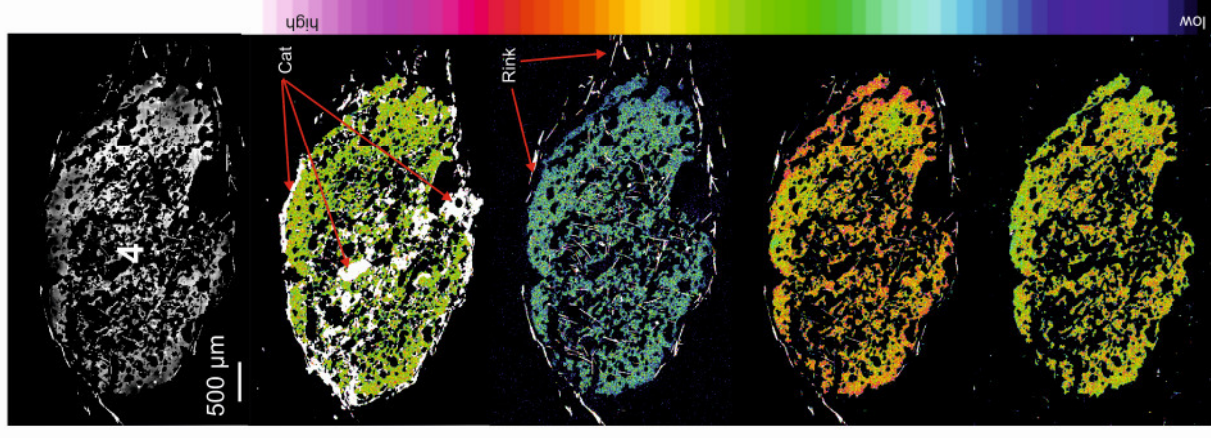
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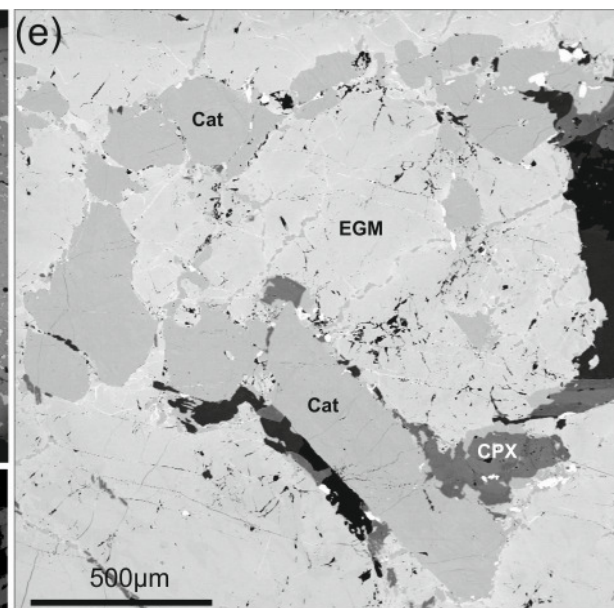
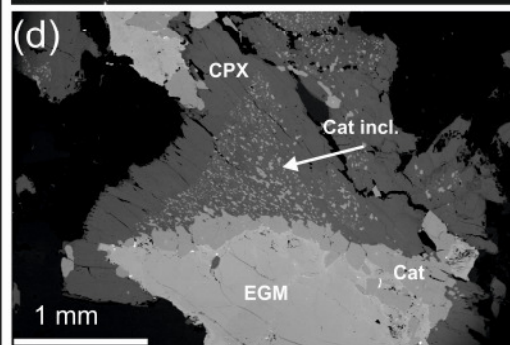
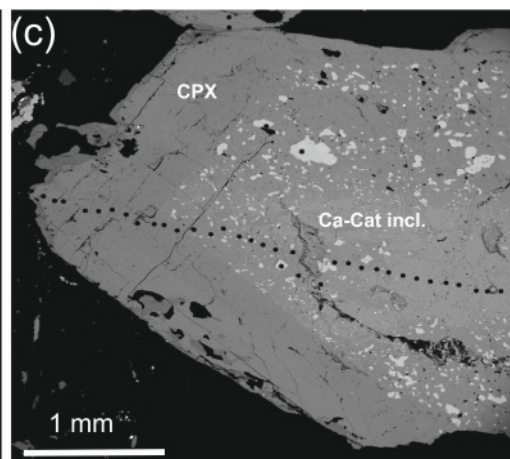
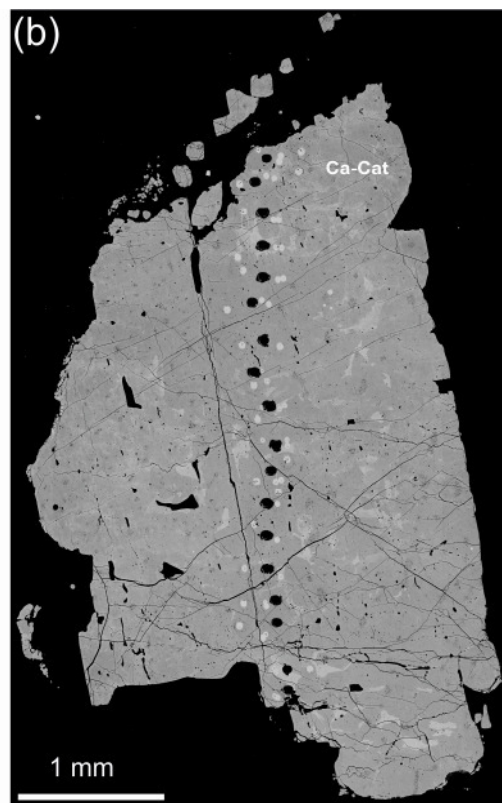
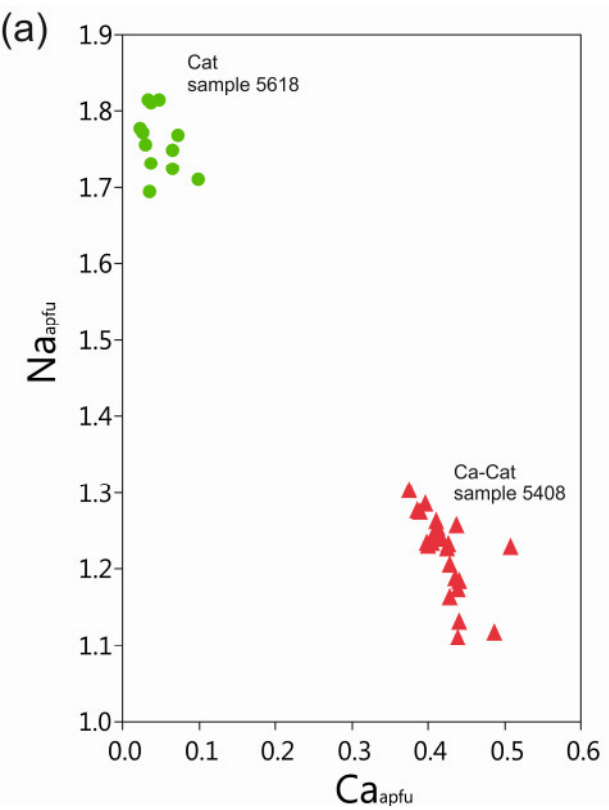


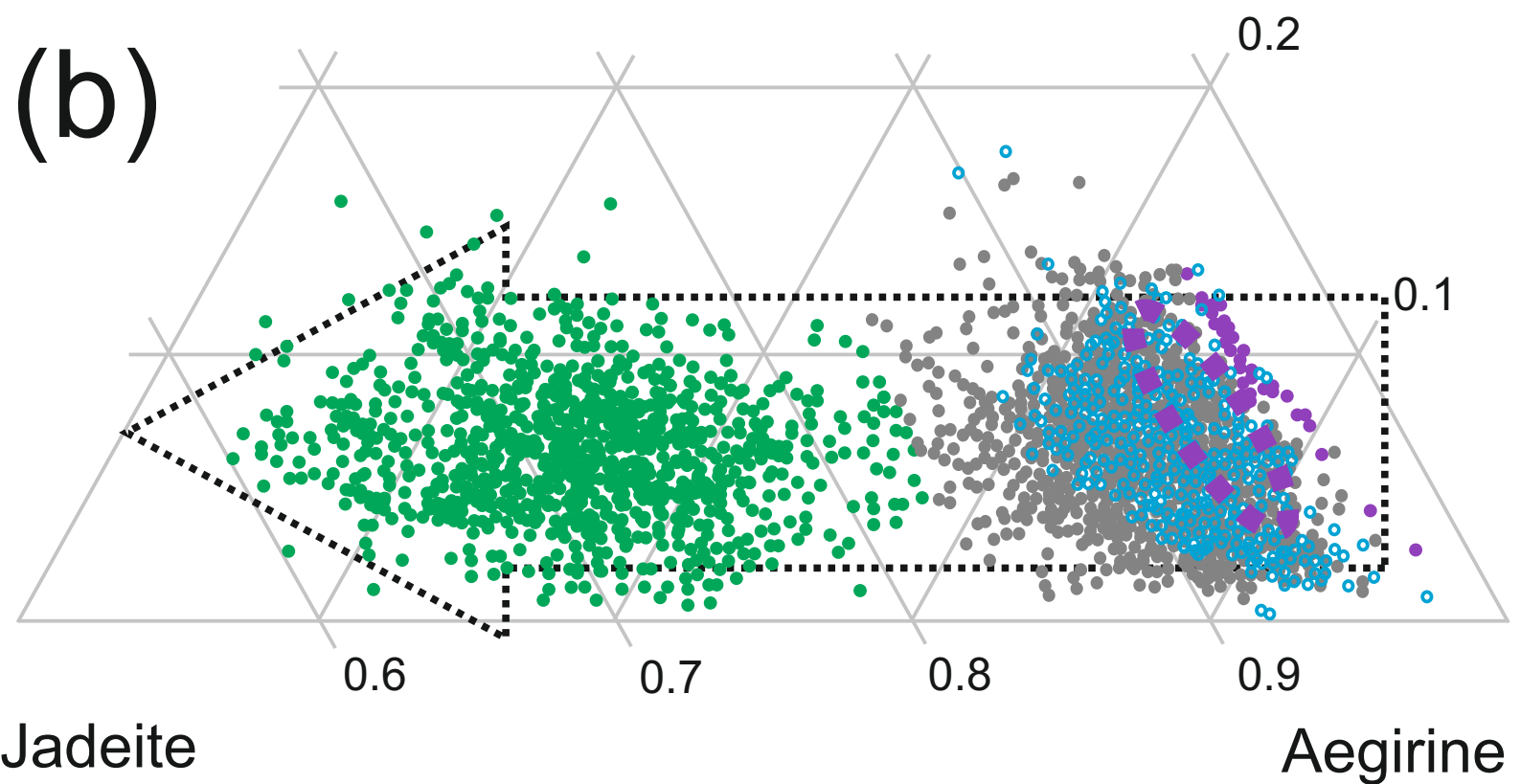
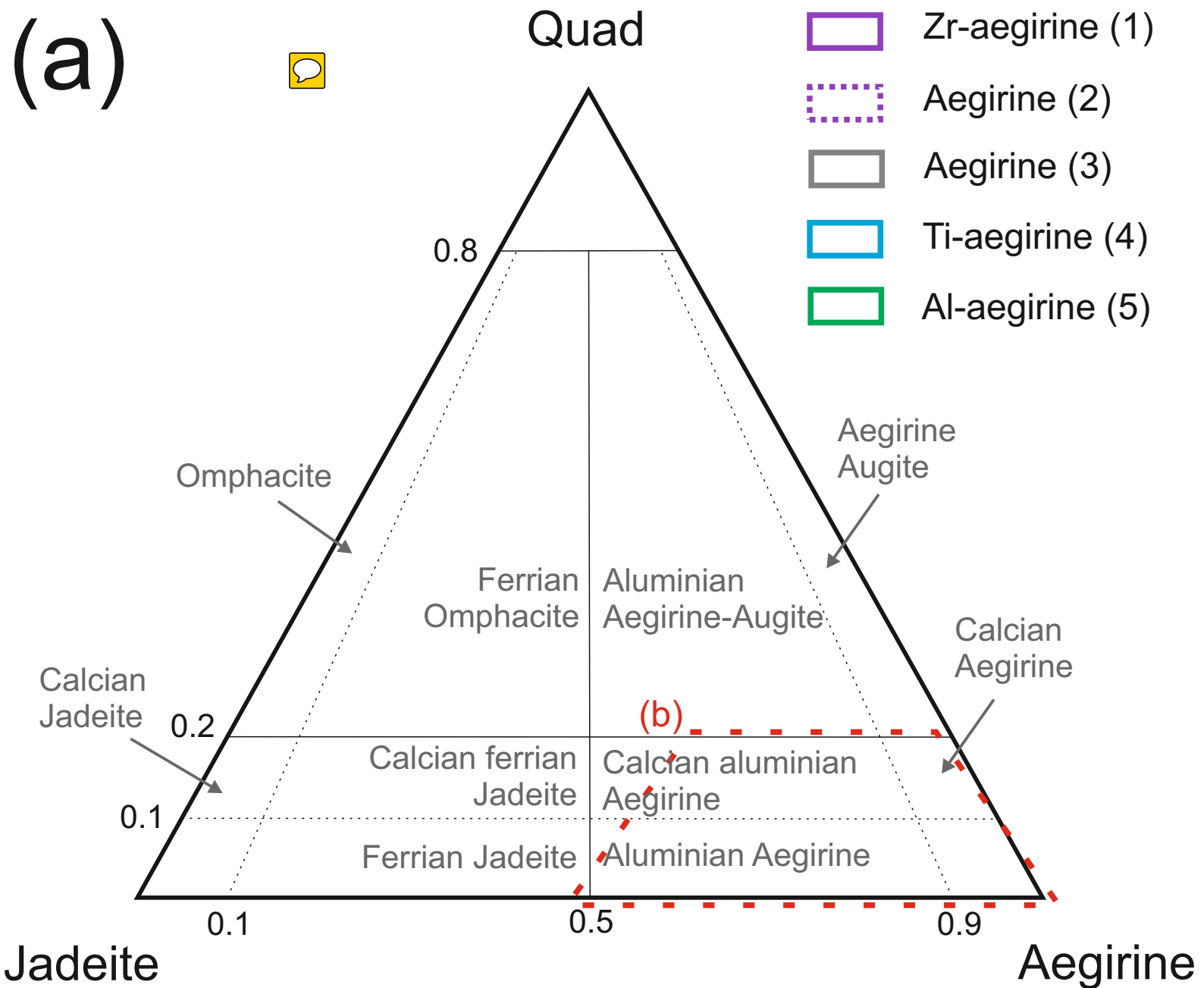
(c) NKA6115



(d) NKA5611

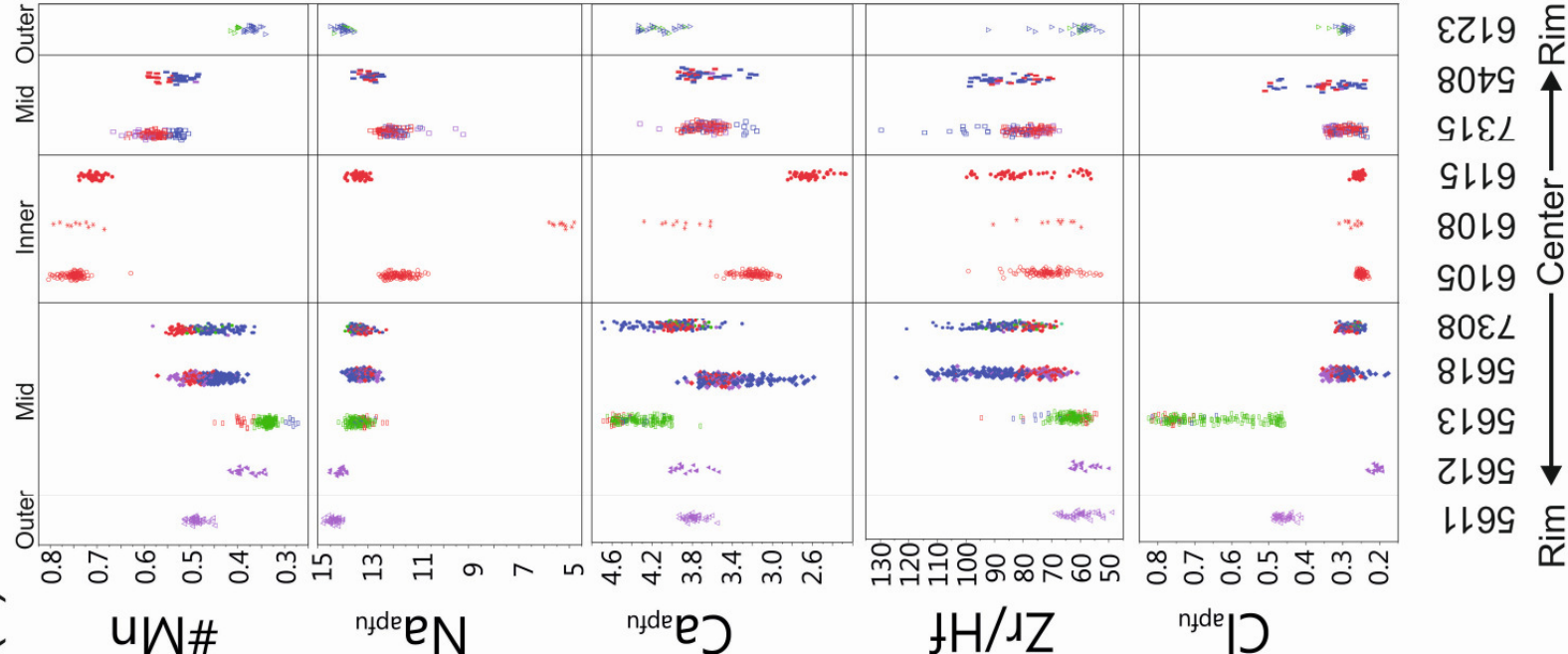




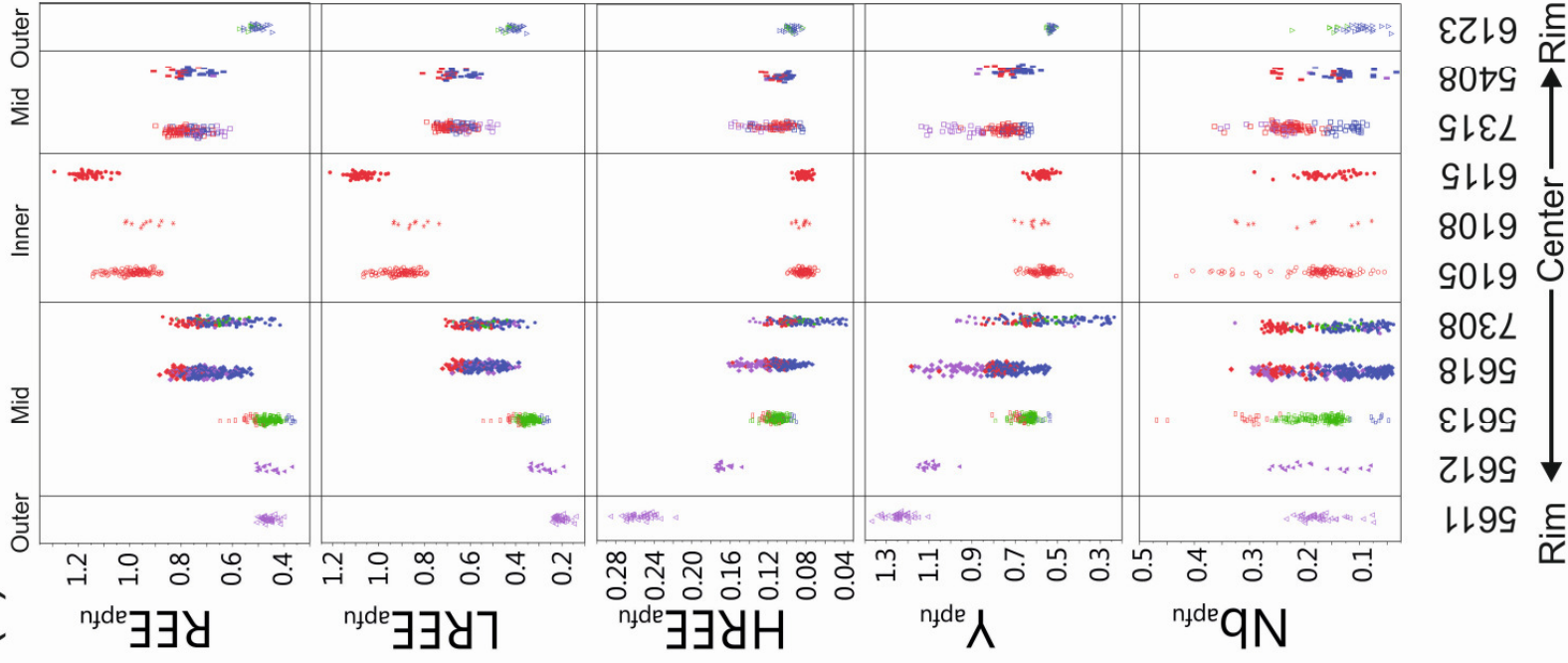


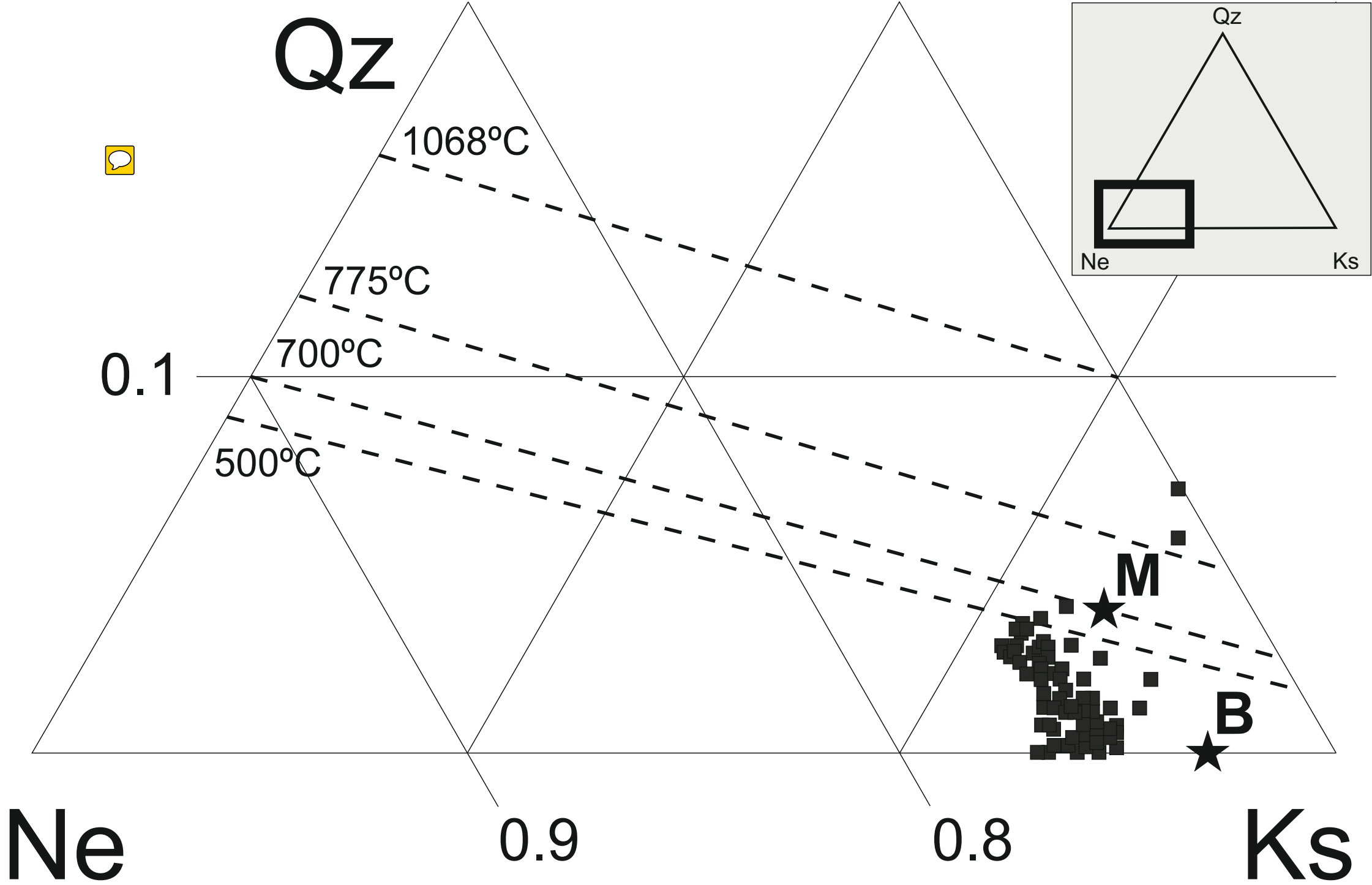


(a)



(b)







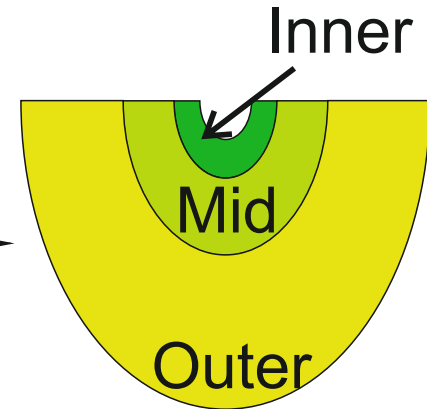


# Magmatic structure



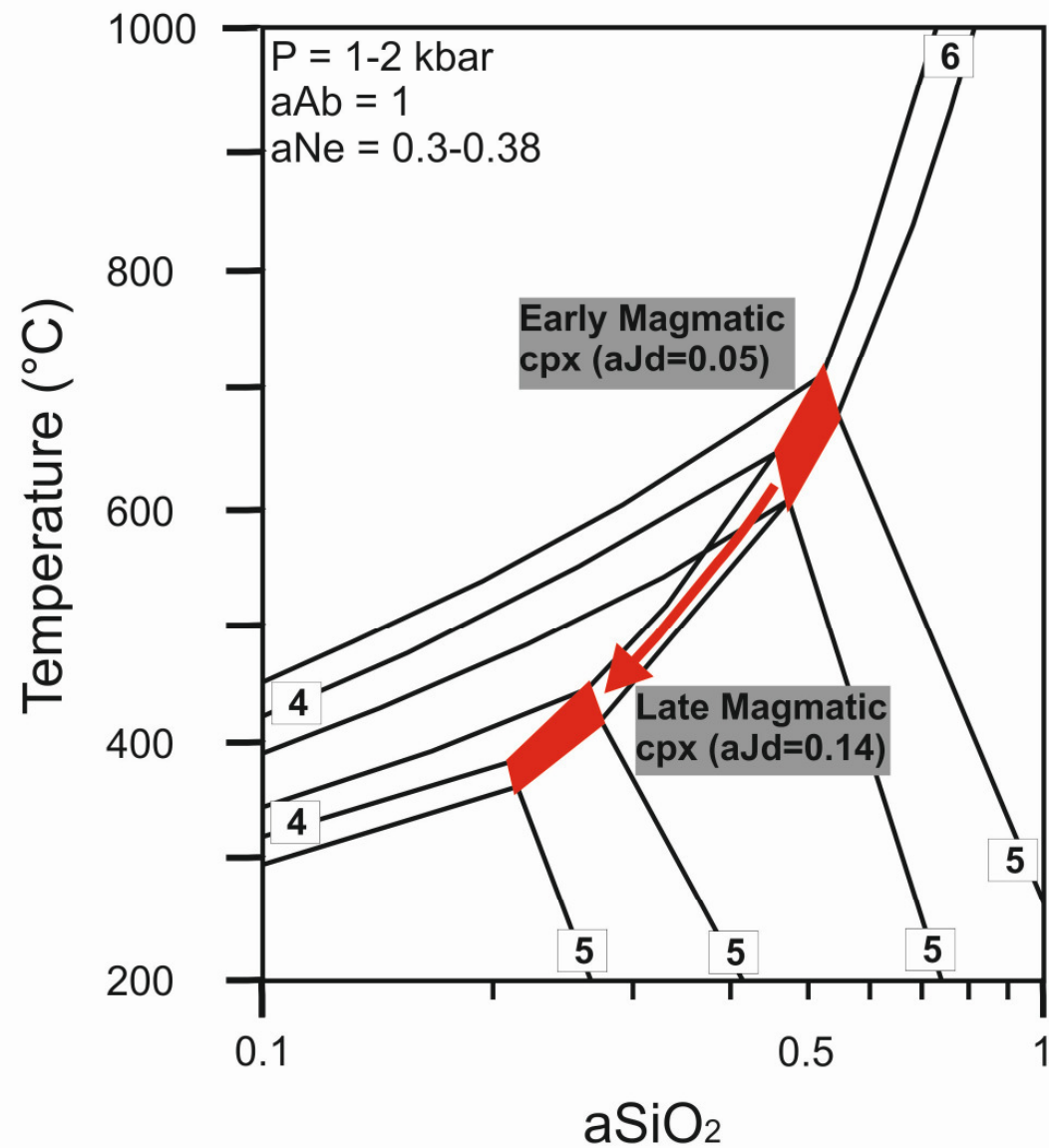
Sveconorwegian orogeny

# Deformed structure

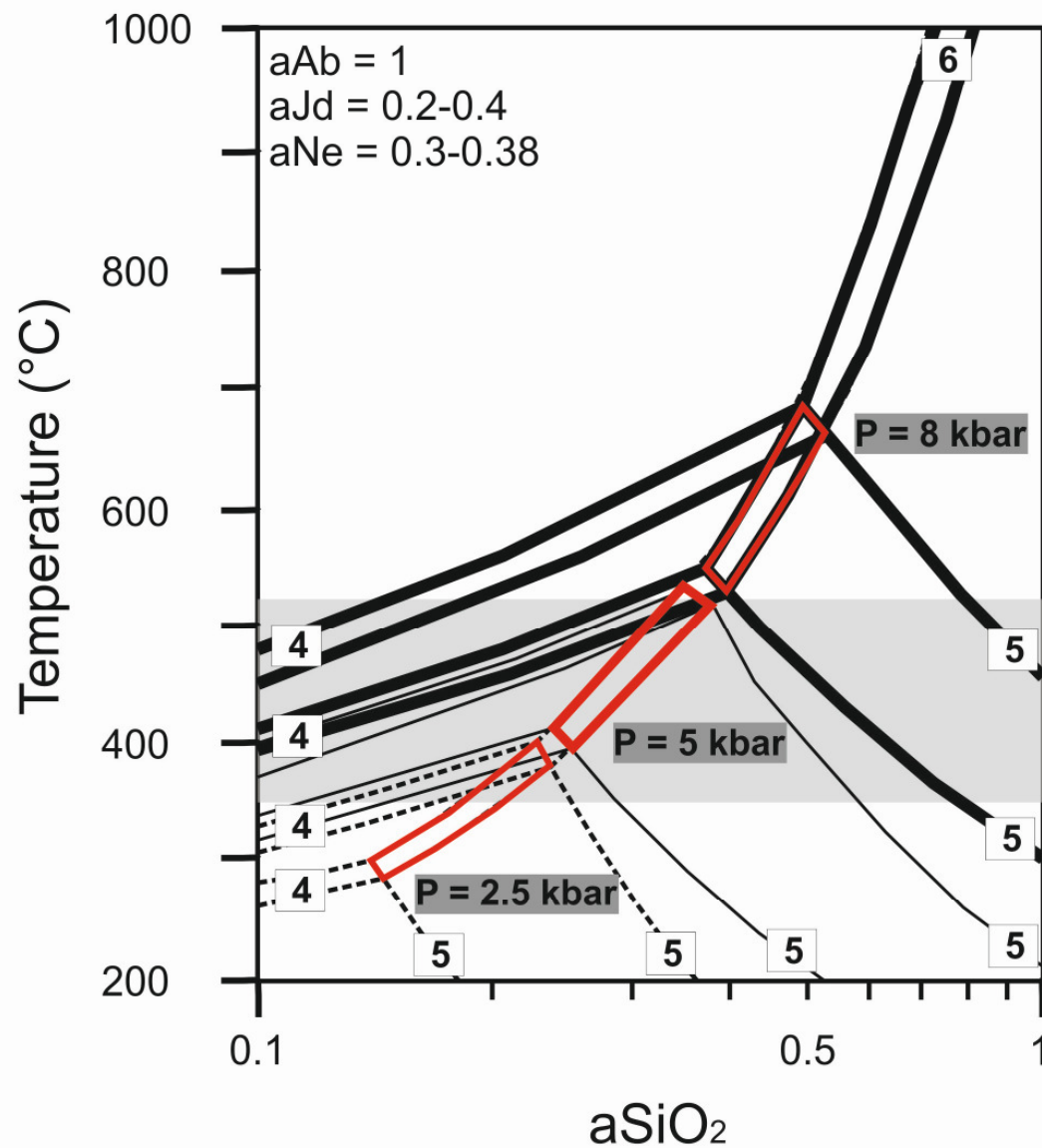




# (a) Magmatic evolution



# (b) Metamorphic conditions



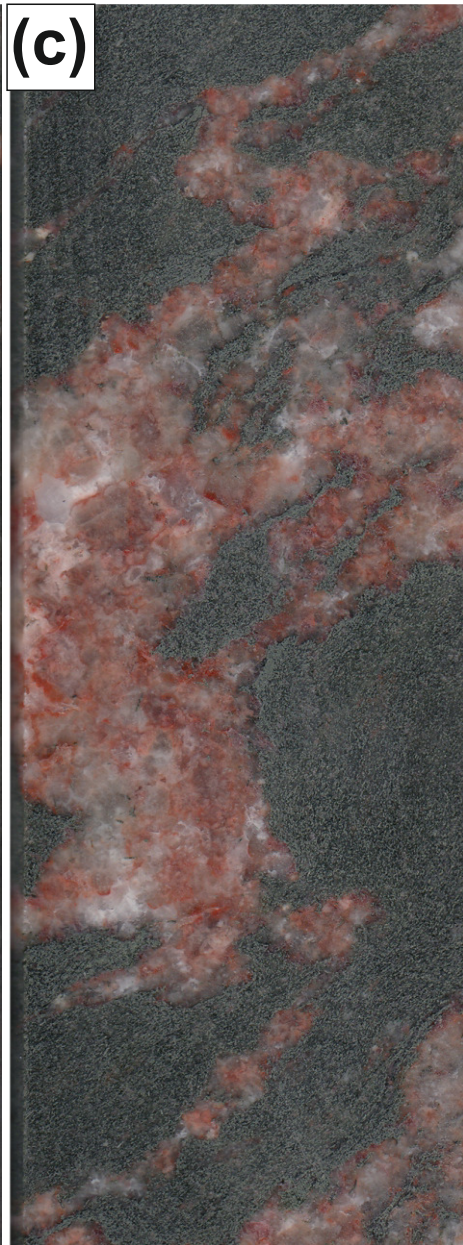


Table 1: List of investigated samples

Sample	Drill core	Position	Subunit	Analytical method (mineral)
5403	NKA 11054 P7	16.66-17.00m	Mid	MLA
5405	NKA 11054 P7	25.08-25.29m	Mid	MLA
5406	NKA 11054 P7	34.31-34.60m	Mid	MLA
5407-1	NKA 11054 P7	37.43-37.86m	Mid	MLA
5407-2	NKA 11054 P7	37.43-37.86m	Mid	MLA
5408	NKA 11054 P7	42.44-42.71m	Mid	MLA, EPMA (cpx, EGM, Cat)
5409	NKA 11054 P7	45.85-46.20m	Mid	MLA
5411	NKA 11054 P7	54.56-54.78m	Mid	MLA
5611	NKA 11056 J1	110.20-110.34m	Outer	MLA, EPMA (cpx, EGM)
5612	NKA 11056 J1	119.60-119.90m	Mid	MLA, EPMA (cpx, EGM)
5613	NKA 11056 J1	126.00-126.25m	Mid	MLA, EPMA (cpx, EGM)
5618	NKA 11056 J1	198.60-198.86m	Mid	MLA, EPMA (cpx, EGM, Cat)
5631	NKA 11056 J1	350.20-350.40m	Outer	MLA, EPMA (cpx)
6105	NKA 12061 J7	29.85-30.02m	Inner	MLA, EPMA (cpx, EGM)
6108	NKA 12061 J7	43.44-43.55m	Inner	MLA, EPMA (cpx, EGM)
6115	NKA 12061 J7	83.36-83.59m	Inner	MLA, EPMA (cpx, EGM)
6123	NKA 12061 J7	163.61-163.80m	Outer	MLA, EPMA (cpx, EGM)
6124-1	NKA 12061 J7	182.30-182.56m	Outer	MLA, EPMA (cpx)
6124-2	NKA 12061 J7	182.30-182.56m	Outer	MLA
6125	NKA 12061 J7	189.90-190.30m	Outer	MLA
7305-1	NKA 12073 P5	~ 22.94m	Mid	MLA
7305-2	NKA 12073 P5	~26.50m	Mid	MLA
7308	NKA 12073 P5	52.81-52.97m	Mid	MLA, EPMA (cpx, EGM)
7310	NKA 12073 P5	64.37-64.51m	Mid	MLA
7311	NKA 12073 P5	71.35-71.43m	Mid	MLA
7314	NKA 12073 P5	89.44-89.57m	Mid	MLA
7315	NKA 12073 P5	91.50-91.61m	Mid	MLA, EPMA (cpx, EGM)
7320	NKA 12073 P5	156.31-156.44m	Outer	MLA

MLA = Mineral Liberation Analyzer; EPMA = Electron Probe Microanalyzer; cpx = clinopyroxene; EGM = Eudialyte-group minerals; Cat = catapleiite

Table 2: Modal mineralogy of subunits of the Norra Kärr complex

wt. %	Inner N=3			Mid (Matrix) N=1	Mid (Schlieren) N=17			Outer N=7		
	Mean	CI (+/-)	95% Interval Min Max		Mean	CI (+/-)	95% Interval Min Max	Mean	CI (+/-)	95% Interval Min Max
Albite	22.88	4.15	13.27 31.66	57.87	4.43	0.84	0.00 21.83	39.66	1.44	29.73 53.85
Microcline	16.80	3.05	15.21 19.03	4.41	25.13	1.85	2.98 58.98	13.17	0.43	8.35 15.83
Aegirine	17.60	3.19	15.24 19.69	19.42	18.20	1.23	2.86 34.03	13.47	0.55	9.42 17.49
Nepheline	9.39	1.70	4.83 12.72	10.30	8.84	1.14	0.02 32.98	20.15	1.61	3.17 29.14
Natrolite	22.11	4.01	12.12 36.86	2.57	27.12	1.47	12.12 51.24	8.16	1.72	0.39 27.19
Catapleiite	1.74	0.32	0.94 2.19	2.97	2.45	0.23	0.10 6.29	1.13	0.14	0.14 2.63
Eudialyte	7.25	1.32	1.11 14.43	2.19	13.39	1.15	2.70 35.39	3.81	0.37	0.38 6.25
Other REE minerals	1.80	0.33	0.07 4.96	0.25	0.14	0.01	0.04 0.43	0.11	0.03	0.00 0.46

CI = confidential interval

Table 3: Average compositions of the various clinopyroxene types of the Norra Kärr complex

wt.%	(1) Zr-aegirine				(2) Aegirine				(3) Aegirine				(4) Ti-aegirine				(5) Al-aegirine			
	Mean 38	CI	95% Interval		Mean 155	CI	95% Interval		Mean 3053	CI	95% Interval		Mean 541	CI	95% Interval		Mean 1350	CI	95% Interval	
			Min	Max			Min	Max			Min	Max			Min	Max			Min	Max
SiO <sub>2</sub>	50.82	0.11	50.25	51.31	51.54	0.06	50.69	52.20	51.49	0.02	50.41	52.43	51.44	0.05	50.13	52.24	52.74	0.03	51.60	53.83
TiO <sub>2</sub>	0.51	0.01	0.47	0.55	0.43	0.02	0.21	0.60	0.31	0.00	0.12	0.61	1.10	0.03	0.66	2.10	0.21	0.01	0.03	0.56
Al <sub>2</sub> O <sub>3</sub>	0.86	0.02	0.78	0.94	1.56	0.06	0.87	2.25	1.93	0.02	1.26	3.03	1.99	0.03	1.46	2.56	6.02	0.04	4.28	7.63
FeO	25.78	0.04	25.55	25.95	27.54	0.07	26.61	28.24	27.58	0.02	26.08	28.53	26.35	0.06	24.84	27.53	22.45	0.05	20.54	24.55
MnO	0.64	0.01	0.59	0.70	0.28	0.02	0.14	0.61	0.18	0.00	0.09	0.33	0.31	0.01	0.20	0.49	0.15	0.00	0.07	0.26
MgO	0.43	0.01	0.41	0.48	0.33	0.02	0.17	0.49	0.25	0.00	0.12	0.40	0.35	0.01	0.24	0.48	0.17	0.00	0.08	0.35
ZrO <sub>2</sub>	3.20	0.02	3.11	3.28	0.44	0.03	0.15	0.94	0.29	0.01	0.05	0.92	0.53	0.03	0.08	1.69	0.08	0.00	0.02	0.20
CaO	1.61	0.03	1.50	1.77	1.02	0.04	0.61	1.57	0.81	0.01	0.34	1.53	0.60	0.02	0.36	1.01	0.74	0.01	0.32	1.31
Na <sub>2</sub> O	11.84	0.05	11.54	12.13	12.32	0.04	11.73	12.95	12.43	0.01	11.86	12.90	12.52	0.02	12.11	12.86	12.76	0.01	12.20	13.26
K <sub>2</sub> O	0.00	0.00	0.00	0.02	0.00	0.00	0.00	0.02	0.01	0.00	0.00	0.03	0.01	0.00	0.00	0.02	0.01	0.00	0.00	0.05
Total	95.69	0.14	94.98	96.39	95.45	0.08	94.48	96.35	95.27	0.02	93.78	96.44	95.19	0.06	93.52	96.56	95.33	0.04	93.87	96.50
Fe <sub>2</sub> O <sub>3</sub>	23.42	0.32	22.03	25.41	27.32	0.20	24.94	30.09	27.64	0.05	24.98	29.97	25.91	0.16	22.49	29.51	21.96	0.08	18.76	24.68
FeO	4.70	0.29	2.98	5.97	2.96	0.17	0.79	5.23	2.70	0.04	0.38	4.97	3.03	0.11	0.38	5.26	2.69	0.07	0.42	5.07
Ferric Total	98.04	0.14	97.31	98.94	98.19	0.08	97.22	99.19	98.04	0.02	96.59	99.22	97.79	0.06	96.37	99.00	97.53	0.04	96.04	98.73
Formulae based on 4 cations and 6 oxygens																				
Si	2.01	0.00	1.99	2.02	2.01	0.00	1.99	2.03	2.01	0.00	1.98	2.03	2.01	0.00	1.98	2.03	2.01	0.00	1.99	2.04
Al	0.04	0.00	0.04	0.04	0.07	0.00	0.04	0.10	0.09	0.00	0.06	0.14	0.09	0.00	0.07	0.12	0.27	0.00	0.19	0.34
Ti	0.02	0.00	0.01	0.02	0.01	0.00	0.01	0.02	0.01	0.00	0.00	0.02	0.03	0.00	0.02	0.06	0.01	0.00	0.00	0.02
Fe <sup>3+</sup>	0.70	0.01	0.66	0.75	0.80	0.01	0.73	0.88	0.81	0.00	0.73	0.88	0.76	0.00	0.66	0.87	0.63	0.00	0.54	0.71
Mg	0.03	0.00	0.02	0.03	0.02	0.00	0.01	0.03	0.01	0.00	0.01	0.02	0.02	0.00	0.01	0.03	0.01	0.00	0.00	0.02
Fe <sup>2+</sup>	0.16	0.01	0.10	0.20	0.10	0.01	0.03	0.17	0.09	0.00	0.01	0.16	0.10	0.00	0.01	0.17	0.09	0.00	0.01	0.16
Mn	0.02	0.00	0.02	0.02	0.01	0.00	0.00	0.02	0.01	0.00	0.00	0.01	0.01	0.00	0.01	0.02	0.00	0.00	0.00	0.01
Zr	0.06	0.00	0.06	0.06	0.01	0.00	0.00	0.02	0.01	0.00	0.00	0.02	0.01	0.00	0.00	0.03	0.00	0.00	0.00	0.00
Ca	0.07	0.00	0.06	0.08	0.04	0.00	0.03	0.07	0.03	0.00	0.01	0.06	0.02	0.00	0.01	0.04	0.03	0.00	0.01	0.05
Na	0.91	0.00	0.89	0.92	0.93	0.00	0.89	0.97	0.94	0.00	0.90	0.97	0.95	0.00	0.92	0.97	0.94	0.00	0.91	0.98
K	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
χ Ae	78.43	0.38	76.98	81.31	87.86	0.27	84.67	92.03	87.77	0.08	82.58	92.23	81.70	0.38	72.70	90.23	68.92	0.20	61.53	76.54
χ Jd	4.39	0.16	3.23	4.93	7.70	0.28	4.43	10.92	9.32	0.08	5.79	14.84	9.51	0.15	5.83	12.74	29.53	0.21	21.23	37.41

x Ti-Zr-Ae 17.18 0.25 15.47 18.11 4.44 0.21 1.87 7.32 2.91 0.05 0.71 6.37 8.79 0.30 3.00 16.35 1.55 0.05 0.19 3.86

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CI = confidential interval

Table 4: Average compositions of the various types of eudialyte-group minerals observed in the Norra Kärr rocks

	(1) Sector zoning				(2) Oscillatory zoning				(3) Irregular replacement textures				(4) Rims			
	Mean	CI	95% Interval		Mean	CI	95% Interval		Mean	CI	95% Interval		Mean	CI	95% Interval	
	321	(+/-)	Min	Max	180	(+/-)	Min	Max	374	(+/-)	Min	Max	139	(+/-)	Min	Max
SiO <sub>2</sub>	51.56	0.05	50.59	52.36	51.09	0.04	50.49	51.57	51.20	0.08	49.95	52.55	50.91	0.11	49.69	51.85
TiO <sub>2</sub>	0.16	0.00	0.08	0.22	0.09	0.00	0.06	0.14	0.16	0.00	0.07	0.21	0.12	0.01	0.04	0.20
ZrO <sub>2</sub>	11.92	0.06	11.35	13.65	11.67	0.06	11.06	12.40	11.53	0.02	11.12	11.99	11.38	0.05	11.03	11.83
HfO <sub>2</sub>	0.24	0.00	0.18	0.35	0.31	0.00	0.24	0.35	0.27	0.00	0.21	0.35	0.29	0.01	0.21	0.37
Al <sub>2</sub> O <sub>3</sub>	0.25	0.01	0.16	0.41	0.28	0.01	0.19	0.39	0.20	0.00	0.13	0.28	0.32	0.03	0.13	0.59
Nb <sub>2</sub> O <sub>5</sub>	0.50	0.02	0.21	0.85	0.74	0.02	0.47	1.07	0.94	0.03	0.44	1.54	0.89	0.05	0.30	1.36
La <sub>2</sub> O <sub>3</sub>	0.48	0.01	0.26	0.68	0.41	0.01	0.30	0.61	0.89	0.03	0.50	1.63	0.44	0.03	0.18	0.73
Ce <sub>2</sub> O <sub>3</sub>	1.32	0.02	0.79	1.63	0.95	0.03	0.74	1.45	2.01	0.05	1.07	2.86	0.98	0.07	0.40	1.63
Nd <sub>2</sub> O <sub>3</sub> +Pr <sub>2</sub> O <sub>3</sub> +Sm <sub>2</sub> O <sub>3</sub>	1.16	0.02	0.67	1.42	0.70	0.03	0.50	1.21	1.33	0.03	0.67	1.95	0.83	0.06	0.28	1.37
Tb <sub>2</sub> O <sub>3</sub> +Gd <sub>2</sub> O <sub>3</sub> +Dy <sub>2</sub> O <sub>3</sub>	0.69	0.01	0.32	0.88	0.68	0.01	0.56	0.79	0.67	0.01	0.48	0.89	1.03	0.03	0.76	1.29
Er <sub>2</sub> O <sub>3</sub> +Yb <sub>2</sub> O <sub>3</sub>	0.51	0.01	0.24	0.67	0.66	0.01	0.41	0.78	0.53	0.01	0.36	0.82	1.14	0.09	0.54	2.10
Y <sub>2</sub> O <sub>3</sub>	2.41	0.06	1.02	3.06	2.34	0.03	1.90	2.67	2.45	0.04	1.90	3.12	3.96	0.09	3.19	4.89
FeO	2.84	0.04	2.12	3.83	3.35	0.04	2.68	3.76	2.01	0.06	1.27	3.26	2.21	0.06	1.60	2.87
MnO	2.23	0.03	1.57	2.63	1.83	0.03	1.61	2.43	3.37	0.09	2.07	4.69	2.13	0.06	1.52	2.76
CaO	7.04	0.08	5.71	8.38	7.91	0.06	7.16	8.47	6.47	0.10	4.57	8.36	6.86	0.05	6.32	7.39
Na <sub>2</sub> O	13.42	0.06	12.33	14.48	13.73	0.05	13.11	14.35	12.61	0.13	7.73	13.74	13.58	0.16	11.31	14.92
K <sub>2</sub> O	0.46	0.01	0.19	0.63	0.35	0.01	0.21	0.51	0.50	0.01	0.26	0.95	0.46	0.02	0.28	0.73
Cl	0.36	0.01	0.29	0.59	0.68	0.03	0.31	0.93	0.35	0.01	0.28	0.88	0.41	0.02	0.23	0.56
O=Cl	0.08	0.00	0.07	0.13	0.15	0.01	0.07	0.21	0.08	0.00	0.06	0.20	0.09	0.00	0.05	0.13
Total, corrected	97.45	0.07	96.24	98.36	97.62	0.07	96.67	98.57	97.40	0.07	95.61	98.64	97.85	0.12	96.24	98.89
TREO total	4.16	0.06	2.79	4.89	3.40	0.05	2.98	4.45	5.43	0.09	3.76	6.97	4.41	0.08	3.52	5.22

Formula based on  $\Sigma(\text{Si Al Zr Ti Hf Nb})$  normalized to 29 apfu

Si	25.75	0.02	25.36	25.91	25.72	0.01	25.57	25.86	25.75	0.01	25.63	25.85	25.72	0.01	25.60	25.84
Ti	0.06	0.00	0.03	0.08	0.03	0.00	0.02	0.05	0.06	0.00	0.03	0.08	0.04	0.00	0.02	0.08
Zr	2.90	0.01	2.79	3.27	2.87	0.01	2.73	3.03	2.83	0.01	2.74	2.96	2.80	0.01	2.74	2.90
Hf	0.03	0.00	0.03	0.05	0.04	0.00	0.03	0.05	0.04	0.00	0.03	0.05	0.04	0.00	0.03	0.05
Al	0.15	0.00	0.09	0.23	0.17	0.00	0.12	0.23	0.12	0.00	0.07	0.16	0.19	0.02	0.08	0.35
Nb	0.11	0.00	0.05	0.19	0.17	0.01	0.11	0.25	0.21	0.01	0.10	0.35	0.20	0.01	0.07	0.31
La	0.09	0.00	0.05	0.12	0.08	0.00	0.06	0.11	0.17	0.01	0.09	0.30	0.08	0.01	0.03	0.14
Ce	0.24	0.00	0.14	0.30	0.18	0.00	0.14	0.27	0.37	0.01	0.20	0.54	0.18	0.01	0.07	0.30
Nd+Pr+Sm	0.21	0.00	0.12	0.26	0.13	0.00	0.09	0.22	0.24	0.01	0.12	0.36	0.15	0.01	0.05	0.25



Tb+Gd+Dy	0.06	0.00	0.03	0.07	0.06	0.00	0.05	0.07	0.06	0.00	0.04	0.07	0.09	0.00	0.06	0.11
Er+Yb	0.04	0.00	0.02	0.05	0.05	0.00	0.03	0.06	0.04	0.00	0.03	0.06	0.09	0.01	0.04	0.17
Y	0.64	0.01	0.27	0.81	0.63	0.01	0.51	0.72	0.66	0.01	0.51	0.83	1.07	0.03	0.85	1.32
Fe	1.19	0.02	0.88	1.59	1.41	0.02	1.13	1.59	0.85	0.02	0.54	1.38	0.93	0.03	0.67	1.20
Mn	0.94	0.01	0.67	1.11	0.78	0.01	0.68	1.04	1.43	0.04	0.90	2.02	0.91	0.03	0.65	1.17
Ca	3.77	0.04	2.98	4.49	4.27	0.03	3.84	4.60	3.48	0.05	2.50	4.55	3.72	0.03	3.40	4.03
Na	12.99	0.05	11.90	14.00	13.40	0.04	12.84	13.92	12.30	0.14	7.37	13.57	13.31	0.18	11.04	14.62
K	0.29	0.01	0.12	0.40	0.22	0.01	0.14	0.33	0.32	0.01	0.17	0.59	0.30	0.01	0.18	0.47
Cl	0.31	0.01	0.24	0.50	0.58	0.03	0.27	0.80	0.30	0.01	0.24	0.76	0.35	0.01	0.20	0.49

CI = confidential interval

Table 5: Catapeliite compositions from the Norra Kärr complex

wt %	Ca-Catapleite sample 5408, n=24				Catapleite sample 5618, n=12			
	Mean	CI (+/-)	95% Interval Min Max		Mean	CI (+/-)	95% Interval Min Max	
SiO <sub>2</sub>	45.85	0.18	45.04	46.53	46.54	0.16	46.18	46.93
TiO <sub>2</sub>	0.02	0.00	0.01	0.03	0.03	0.01	0.02	0.05
ZrO <sub>2</sub>	29.07	0.09	28.61	29.39	29.92	0.37	27.80	29.56
HfO <sub>2</sub>	0.59	0.07	0.47	0.91	0.69	0.06	0.54	0.83
REEY <sub>2</sub> O <sub>3</sub>	0.41	0.03	0.30	0.56	0.46	0.05	0.31	0.58
FeO	0.20	0.13	0.04	0.95	1.10	0.55	0.10	2.80
CaO	5.99	0.18	5.41	6.99	0.68	0.20	0.35	1.30
Na <sub>2</sub> O	9.55	0.16	8.75	10.14	13.83	0.23	13.30	14.33
K <sub>2</sub> O	0.04	0.01	0.01	0.09	0.05	0.02	0.02	0.10
Total	91.71	0.31	90.63	93.14	92.30	0.38	91.66	93.27
H <sub>2</sub> O (calc)	7.44	0.02	7.37	7.49	7.76	0.02	7.70	7.78
Total (incl H <sub>2</sub> O)	99.15	0.31	98.10	100.56	100.06	0.39	99.37	101.08
Formulae based on 10 cations and 11 oxygens								
Si	3.02	0.00	3.00	3.04	3.06	0.01	3.05	3.07
Ti	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Zr	0.93	0.00	0.92	0.94	0.93	0.01	0.89	0.95
Hf	0.01	0.00	0.01	0.02	0.01	0.00	0.01	0.02
REEY	0.01	0.00	0.01	0.01	0.01	0.00	0.01	0.01
Fe	0.01	0.01	0.00	0.05	0.06	0.03	0.01	0.15
Ca	0.42	0.01	0.38	0.50	0.05	0.01	0.02	0.09
Na	1.22	0.02	1.11	1.29	1.76	0.03	1.70	1.82
K	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01
H (calc)	3.76	0.01	3.72	3.78	3.92	0.01	3.89	3.95

CI = confidential interval

Table 6: Average feldspar compositions in metasyenites from the Norra Kärr complex

wt %	Microcline n=338				Albite n=122			
	Mean	CI (+/-)	95% Interval Min Max		Mean	CI (+/-)	95% Interval Min Max	
SiO <sub>2</sub>	65.03	0.03	64.39	65.44	69.44	0.04	68.92	69.87
Al <sub>2</sub> O <sub>3</sub>	18.99	0.01	18.76	19.18	19.54	0.02	19.32	19.72
FeO	0.10	0.01	0.01	0.34	0.09	0.01	0.00	0.23
BaO	0.06	0.00	0.00	0.12	0.03	0.00	0.00	0.07
CaO	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01
Na <sub>2</sub> O	0.38	0.02	0.06	0.60	11.85	0.03	11.56	12.11
K <sub>2</sub> O	17.15	0.03	16.81	17.64	0.10	0.01	0.05	0.17
TOTAL	101.70	0.04	100.81	102.26	101.06	0.06	100.36	101.70
Formulae based on 32 oxygens								
Si	12.03	0.01	11.92	12.11	12.74	0.01	12.64	12.82
Al	4.14	0.00	4.09	4.18	4.22	0.00	4.18	4.26
Fe <sup>2+</sup>	0.02	0.00	0.00	0.05	0.01	0.00	0.00	0.03
Ca	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Na	0.14	0.01	0.02	0.21	4.22	0.01	4.11	4.31
K	4.05	0.01	3.97	4.16	0.02	0.00	0.01	0.04
Ba	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01
X <sub>An</sub>	0.01	0.00	0.00	0.05	0.02	0.00	0.00	0.06
X <sub>Ab</sub>	3.23	0.15	0.52	5.12	99.42	0.05	99.08	99.69
X <sub>Or</sub>	96.76	0.15	94.88	99.45	0.56	0.05	0.28	0.91

CI = confidential interval

Table 7: Average nepheline, natrolite and analcime compositions from Norra Kärr rocks

wt %	Nepheline N=108				Natrolite N=104				Analcime N=4			
	Mean	CI	95% Interval		Mean	CI	95% Interval		Mean	CI	95% Interval	
		(+/-)	Min	Max		(+/-)	Min	Max		(+/-)	Min	Max
SiO <sub>2</sub>	42.23	0.16	41.45	43.25	47.66	0.06	47.18	48.99	50.17	7.24	46.77	52.30
Al <sub>2</sub> O <sub>3</sub>	34.27	0.13	33.41	34.92	27.39	0.09	26.69	28.11	26.30	4.60	25.12	28.35
Fe <sub>2</sub> O <sub>3</sub>	0.12	0.02	0.04	0.33	0.13	0.04	0.00	0.44	0.13	0.23	0.04	0.23
BaO	0.04	0.00	0.00	0.09	0.03	0.00	0.00	0.07	0.04	0.07	0.01	0.06
CaO	0.02	0.02	0.00	0.27	0.01	0.01	0.00	0.06	0.03	0.03	0.02	0.03
Na <sub>2</sub> O	16.08	0.10	14.92	16.52	15.24	0.04	14.92	15.60	13.50	2.64	12.25	14.17
K <sub>2</sub> O	7.59	0.10	6.87	8.03	0.06	0.01	0.01	0.15	0.53	1.49	0.05	1.10
Total	100.35	0.09	99.66	101.07	90.53	0.12	89.52	91.66	90.71	5.05	88.51	92.52
	Formulae based on 16 oxygens				Formulae based on 10 oxygens				Formulae based on 6 oxygens			
Si	0.70	0.00	0.69	0.72	3.01	0.00	2.98	3.03	1.90	0.27	1.77	1.98
Al	0.67	0.00	0.66	0.69	2.04	0.01	1.99	2.09	1.17	0.21	1.12	1.27
Fe <sup>3+</sup>	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.02	0.00	0.01	0.00	0.01
Ca	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Na	0.52	0.00	0.48	0.53	1.87	0.01	1.83	1.91	0.99	0.19	0.90	1.04
K	0.16	0.00	0.15	0.17	0.00	0.00	0.00	0.01	0.03	0.07	0.00	0.05
B	-	-	-	-	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00

CI = confidential interval

Table 8: Reintegrated compositions of Zr-aegirine (based on the composition, frequency and molar volumes of catapleite and aegirine s.str.) compared with measured compositions of Zr-aegirine core from sample 5408

	Reintergrated		Measured
Vol %	1	2	
Cpx	89	84	
Catapleite	11	16	
SiO <sub>2</sub>	51.07	50.94	50.83
TiO <sub>2</sub>	0.51	0.49	0.50
Al <sub>2</sub> O <sub>3</sub>	1.13	1.10	0.85
FeO	25.93	25.15	25.78
MnO	0.34	0.33	0.64
ZrO <sub>2</sub>	2.11	2.96	3.21
CaO	1.20	1.17	1.63
Na <sub>2</sub> O	12.25	12.30	11.83
K <sub>2</sub> O	0.01	0.01	0.00
Total	94.55	94.46	95.27