

1 **Insights into post-magmatic metasomatism and Li circulation in**
2 **granitic systems from phosphate minerals of the Nanping No. 31**
3 **pegmatite (SE China)**

4

5 Can Rao^{a,*}, Rucheng Wang^b, Yueqing Yang^c, Frédéric Hatert^d, Qunke Xia^a, Xuege Yue^a,
6 Wumengyu Wang^a

7

8 ^a School of Earth Sciences, Zhejiang University, Hangzhou 310027, China

9 Corresponding author's E-mail: canrao@zju.edu.cn

10 ^b State Key Laboratory for Mineral Deposits Research, School of Earth Sciences and Engineering,
11 Nanjing University, Nanjing 210046, China

12 ^c Institute of Mineral resource, Chinese academy of Geological science, Beijing, 100037, China

13 ^d Laboratoire de Minéralogie, B18, Université de Liège, B-4000 Liège, Belgium

14

15

16

17

18

19

20

21

22

23

24

25 **ABSTRACT**

26 Phosphate minerals are key indicators for understanding the processes of diagenesis
27 and mineralization in granitic systems. More importantly, these minerals provide
28 constraints on the post-magmatic behaviour of Li in pegmatite systems that remain to
29 be explored, although Li from early-crystallized Li phosphates is known to leach into
30 hydrothermal fluids. In this study, we performed a detailed petrographic and
31 compositional analysis of phosphate minerals from the Nanping No. 31 pegmatite in
32 southeastern of China, which provides new evidence of Li circulation in granitic
33 systems. Primary phosphate minerals evolved from xenotime and monazite to Mn-rich
34 fluorapatite and then to triphylite and montebrasite, with large amounts of
35 montebrasite in the intermediate zones of the pegmatite, reflecting the increasing
36 activities of both Li and P in peraluminous granitic melts. Montebrasite experienced
37 stages of both high-temperature (360-273°C) and low-temperature (273-100°C)
38 hydrothermal alteration. The high-temperature hydrothermal alteration of
39 montebrasite by Fe- and Mg-rich fluids resulted in the formation of a series of Fe-,
40 Mg-bearing phosphates; the replacement of montebrasite under low-temperature
41 alteration generated amounts of Ca-, Sr-, Ba-bearing phosphate minerals, muscovite
42 and akdalaite (Al₂O₃)₄·H₂O. The formation of micro-networks of akdalaite from
43 montebrasite indicate the low mobility of Al in hydrothermal fluids. A four-stage
44 scenario of post-magmatic Li transport in the Nanping No. 31 pegmatite is proposed:
45 (1) Li derived from the breakdown of primary montebrasite was locally recrystallized
46 and produced secondary montebrasite (Mtb-1); (2) hydrothermal alteration by Fe- and
47 Mg-bearing fluids leached Li from montebrasite to form secondary triphylite and
48 simferite; (3) Li derived from the alteration of montebrasite by Sr-, Ca-rich fluids was
49 involved in the formation of secondary palermoite and bertossaite; and (4) the
50 replacement of secondary triphylite, montebrasite, palermoite and bertossaite by later
51 phases (apatites, ludlamite, anapaite, augelite and fine-grained muscovite) reflect the
52 leaching of Li back into hydrothermal fluids. The re-enrichment of Li during the
53 post-magmatic stage most likely increased the solubility of Ta in the hydrothermal
54 fluids of the granitic pegmatite.

55 **Keywords:** phosphate minerals; montebrasite; hydrothermal alteration; akdalaite; Li
56 circulation; Nanping pegmatite.

57

58 **1. Introduction**

59 Montebrasite, triphylite, apatites and other phosphate phases are common
60 constituents of phosphorus-bearing pegmatites (Moore, 1973; Shigley and Brown
61 1985). Due to their variable crystallochemical properties and complex paragenesis,
62 these minerals are sensitive to different geochemical conditions in granitic pegmatites
63 (Fransolet, 2007); they thus behave as key indicators of pegmatite differentiation and
64 can be used to better understand the detailed magmatic-hydrothermal processes of
65 pegmatites (Shigley and Brown, 1985). Amblygonite-montebrasite $\text{LiAlPO}_4(\text{F},\text{OH})$
66 and triphylite-lithiophilite $\text{Li}(\text{Fe},\text{Mn})\text{PO}_4$ group minerals, which are common
67 magmatic Li phases in pegmatites, are often intensively altered to secondary phases
68 during the post-magmatic stages of pegmatite development (e.g., London and Burt,
69 1982; Fransolet et al., 1986; Baldwin et al., 2000; Roda-Robles et al., 2004; Nizamoff,
70 2006; Vignola et al., 2008; Hatert et al., 2011; Baijot et al., 2012; Galliski et al., 2012;
71 Shirose and Uehara, 2014). Hydrothermal fluids at the late stage of pegmatite
72 crystallization are also well known to affect spodumene or other Li phases (e.g.,
73 London and Burt, 1982; Rao et al., 2012). These metasomatic alterations involve leaching
74 Li from early Li phases into hydrothermal fluids. However, due to the lack of a
75 reliable record of the post-magmatic transport of Li, the circulation of Li in pegmatitic
76 systems is poorly understood. Meanwhile, increasing demand for pegmatitic Li
77 resources (Grosjean et al., 2012; Linnen et al., 2012) and advances in the
78 measurement of Li isotopes (Tomascak et al., 2002; Elliott et al., 2004, 2006; Halama
79 et al., 2009; Pogge von Strandmann and Henderson, 2015; Magna et al., 2016)
80 provide the impetus and mechanisms for understanding the entire Li history of
81 granitic systems.

82 The Nanping No. 31 pegmatite (Fujian Province, southeastern China) is a highly
83 evolved and mineralized granitic pegmatite (Rao et al., 2009). It is extremely enriched

84 in Nb, Ta, Sn, Be, and Li and hosts abundant phosphate minerals, thus recording many
85 species and their complex paragenesis. Previous studies of the Nanping No. 31
86 pegmatite mainly focused on its geology and petrology (Li et al., 1983; Yang et al.,
87 1987), Nb-Ta-Sn-bearing minerals (Yang et al., 1989, 2006; Wang et al., 1999; Rao et
88 al., 2009), and the behaviour of its hydrothermal alterations (Yang et al., 1994, 1995;
89 Rao et al., 2011, 2012, 2014a). Three new mineral species, namely, nanpingite (Yang
90 et al., 1988), strontiohurlbutite (Rao et al., 2014b) and minjiangite (Rao et al., 2015),
91 were first discovered in the Nanping No. 31 pegmatite. In the present study, we
92 investigate the petrologic characteristics and evolution of phosphate minerals and
93 reveal the complex metasomatic processes associated with montebrasite from the
94 Nanping No. 31 pegmatite. Based on these data, we constrain the magmatic and
95 post-magmatic behaviour of Li in the Nanping No. 31 pegmatite. Furthermore, we
96 provide direct evidence of the immobile geochemical behaviour of Al in the
97 hydrothermal fluids, and discuss the influence of Li circulation on the solubilities of
98 Ta in hydrothermal fluids.

99

100 **2. Geological setting**

101 The Nanping pegmatite zone is located in the northeastern region of the Cathaysia
102 Block in southeastern China (Fig. 1). In this pegmatite zone, the exposed strata can be
103 grouped into the cover and the basement. The cover consists of sedimentary rocks that
104 were deposited between the Late Devonian and the Late Jurassic; the basement is
105 composed of Meso- and Neo-Proterozoic schists and granulites of the Xiafeng fabric
106 in the Wanqun rock suite, which was intruded by most of the Nanping pegmatite
107 dykes (Yang et al., 1987). Folds and faults are well developed in the basement. The
108 Nanping synclinorium is the primary tectonic unit in this field and is further divided
109 into groups of sub-folds with axes oriented to the northeast and from south to north, in
110 which the latter group overlaps the former. The occurrence, shape and distribution of
111 the Nanping pegmatite dykes are mainly controlled by these sub-folds. Different
112 levels of deformation, caused by faults running south to north, northeast, and

113 northwest, have been found in both the basement and cover rocks. The Hercynian and
114 Yanshanian (Jurassic) granites represent the main magmatic rocks found in this field,
115 both of which are characterized by alkaline, peraluminous rocks with high Si contents
116 and low Ti, Fe, Mg and Ca concentrations (Yang et al., 1987). According to the
117 LA-ICPMS U-Pb dating of columbite-(Fe) and zircon, the emplacement of the Nanping
118 pegmatites occurred at 387 Ma (Tang et al., 2017).

119

120 **3. Petrographic characteristics of the Nanping pegmatites**

121 The Nanping pegmatite zone includes approximately 500 pegmatite dykes with an
122 outcrop area of approximately 250 km². These dykes commonly form lenticular
123 bodies with a general north-northeast strike, are generally 10 to 100 m long (with a
124 maximum length of 600 m) in the direction of strike and are approximately 2 to 10 m
125 wide (with a maximum of 32 m) (see Fig. 2). Based on their different degrees of
126 fractionation, Yang et al. (1987) divided these pegmatites into four groups:
127 muscovite-orthoclase-albite (tabular) pegmatite (type I), muscovite-albite-orthoclase
128 pegmatite (type II), muscovite-orthoclase-albite (fine-grained) pegmatite (type III),
129 and muscovite-albite-spodumene pegmatite (type IV). The type I pegmatite mainly
130 contains 19 vol.% K-feldspar, 35 vol.% albite, 39 vol.% quartz, and 5.9 vol.%
131 muscovite; the type II pegmatite is mainly composed of 27.5 vol.% K-feldspar, 26
132 vol.% albite, 29.5 vol.% quartz, and 15.5 vol.% muscovite; the type III pegmatite
133 consists of 13 vol.% K-feldspar, 48 vol.% albite, 26 vol.% quartz, 10 vol.% muscovite,
134 and < 5 vol.% spodumene; and the type IV pegmatite is characterized by 4.01 vol.%
135 K-feldspar, 39.98 vol.% albite, 29.95 vol.% quartz, 8.68 vol.% muscovite, 10.92
136 vol.% spodumene, and 1.02 vol.% montebrasite. From the type I pegmatites to the
137 type IV pegmatites, the Nanping pegmatite varies from the barren to spodumene
138 subtypes. The No. 31 pegmatite is a type IV pegmatite and is one of the most highly
139 evolved and well-mineralized pegmatites in the Nanping pegmatite zone (Yang et al.,
140 1987).

141 The Nanping No. 31 pegmatite dyke is located at longitude E 118°06', latitude N

142 26°40', it is a typical LCT-type pegmatite with strong mineralization of Nb, Ta, Sn,
143 Be, and Li (Rao et al., 2009). It is 5 to 6 m in width, 300 to 600 m in length, and
144 extends to a depth of 90 m. The internal textures observed in the quarry 515 m below
145 the surface define five discontinuous mineral-textural zones. From the outermost zone
146 inward (Yang et al., 1987; Rao et al., 2009), they are **Zone I** (the
147 quartz-albite-muscovite zone), which is mainly composed of medium-grained quartz,
148 muscovite and fine-grained albite, with small amounts of accessory minerals such as
149 cassiterite, columbite-tantalite, zircon, beryl, hurlbutite, phenakite, apatite, and
150 strontiohurlbutite; **Zone II** (the saccharoidal albite ± muscovite zone), which is
151 characterized by dominant saccharoidal albite with greenish muscovite and can be
152 further subdivided into subzone IIa (saccharoidal albite > 90 vol%) and subzone IIb
153 (greenish muscovite with 10 vol% quartz and 30 vol% albite), in which
154 Nb-Ta-Sn-bearing oxide minerals such as cassiterite, columbite-tantalite, wodginite,
155 tapiolite-(Fe) are mainly concentrated at the boundary between the two subzones (Rao
156 et al., 2009) and Be-bearing minerals (beryl, phenakite, hydroxyherderite, hurlbutite,
157 euclase and strontiohurlbutite) have also been found (Rao et al., 2011, 2014b); **Zone**
158 **III** (the albite-quartz-spodumene zone), which is mainly composed of platy crystals of
159 albite, coarse-grained quartz and spodumene, although columbite-tantalite,
160 wodginite-group minerals, tapiolite, microlite, zircon, cassiterite, apatite, montebrasite
161 and beryl are also present in this zone; **Zone IV** (the quartz-spodumene-montebrasite
162 zone), which mainly contains coarse-grained quartz and spodumene and montebrasite
163 crystals, as well as accessory minerals such as columbite-group minerals,
164 wodginite-group minerals, tapiolite-(Fe), microlite, beryl, cassiterite, pollucite,
165 minjiangite, lazulite, kulanite and strontiohurlbutite; and **Zone V** (the blocky
166 quartz-K-feldspar zone) is the pegmatite core and mainly contains blocky quartz and
167 K-feldspar crystals with a small amount of accessory minerals.

168

169 **4. Analytical methods**

170 In this study, a suite of rock samples was collected from the mine quarry excavated

171 515 m below the surface in the Nanping No. 31 pegmatite dyke. A total of 31
172 phosphate mineral species were identified based on their petrologic and chemical
173 features (Table 1). The chemical compositions of these minerals were obtained by
174 using a JEOL JXA-8100M electron microprobe to perform wavelength-dispersion
175 spectrometry (WDS mode, with settings of 15 kV, 20 nA, and a beam diameter of 1
176 μm). Measurements were performed in the State Key Laboratory for Mineral Deposits
177 Research at Nanjing University. Element peaks and backgrounds were measured with
178 counting times of 10 s and 5 s, respectively. The following standards were used: albite
179 (Si), hornblende (Na, K, Mg, Al, Ca, and Fe), fluorapatite (P and F), MnTiO_4 (Mn and
180 Ti), synthetic $\text{Ba}_3(\text{PO}_4)_2$ (Ba), and synthetic SrSO_4 (Sr). A computer program that
181 implements the ZAF method was used for all data reduction. The contents of Li_2O ,
182 BeO and H_2O in phosphate minerals were estimated by stoichiometry.

183 Powder XRD patterns were collected from finely ground samples spread on glass
184 slides using a RIGAKU D/max Rapid IIR micro-diffractometer ($\text{CuK}\alpha$, $\lambda = 1.54056$
185 \AA). These measurements were performed in the School of Earth Sciences and
186 Info-physics at Central South University in China. The micro-diffractometer was
187 operated with a Gandolfi-like motion, under conditions of 48 kV and 25 mA, using a
188 0.05 mm diameter collimator; the total exposure time was 2 hr. Structural models of
189 fluorapatite and akdalaite (see below) were used to refine the powder XRD data.

190

191 **5. Mineralogy of phosphate minerals**

192 **5.1. *Montebrasite* $\text{LiAl}(\text{PO}_4)(\text{OH},\text{F})$**

193 Montebrasite occurs as one of the main rock-forming minerals in the Nanping No.
194 31 pegmatite. According to petrographic relationships, montebrasite occurs as both
195 primary and secondary phases in different zones of this pegmatite. Primary
196 montebrasite is distributed with variable abundance and different occurrences in zones
197 I, III and IV (Table 1). In zone I, montebrasite occurs as discrete crystals up to one
198 millimeter long between quartz and fine-grained albite; some crystals are replaced by
199 late fine-grained muscovite and fluorapatite (Fig. 3a). Montebrasite from zone III

200 forms anhedral crystals up to 10 cm long, and is interstitial to coarse albite or located
201 between quartz and spodumene. In some cases, montebrasite crystals have been
202 progressively altered to lazulite, palermoite and fluorapatite (Fig. 3b). Massive
203 montebrasite crystals mainly appear in zone IV; they are generally 20 to 40 cm long
204 and are closely associated with massive quartz, spodumene, beryl, Nb-Ta-bearing
205 oxides, cassiterite, and muscovite. Chemically, primary montebrasite has low F
206 contents, which decrease from the outer zones of the pegmatite moving inward (the
207 average F content in montebrasite is 1.59 wt.% in zone I and 1.27 wt.% in zone IV,
208 Table 2).

209 Secondary montebrasite mainly occurs in zones III and IV, in which three different
210 types of secondary montebrasite were distinguished based on their petrological
211 features and paragenetic relationships. The most common occurrence is represented
212 by secondary montebrasite (Mtb-1), which forms small crystals, up to 500 μm in
213 diameter, along the fractures or rims of massive montebrasite in zones III and IV (e.g.
214 Fig. 4a). The second occurrence is characterized by montebrasite assemblages (Mtb-2)
215 that are up to 400 μm in diameter and form along the cleavage planes and/or rims of
216 lazulite crystals in zone IV (Fig. 4b). In some cases, this type of secondary
217 montebrasite occurs in the cleavage planes of lazulite, forming up to 40 μm long
218 veinlets (Fig. 4c). Irregular assemblages or veinlets of montebrasite (Mtb-3) were
219 observed in the fractures and rims associated with palermoite or bertossaite in zone IV
220 (Fig. 4d). The results of electron microprobe analyses indicate that the chemical
221 compositions of secondary montebrasite are similar, as they contain very low F
222 contents (with an average value of 0.10 wt.% F), which are indicative of almost
223 end-member montebrasite (Table 2).

224 5.2. Apatites $\text{Ca}_5(\text{PO}_4)_3(\text{F},\text{OH})$

225 Apatite group minerals occur as primary and secondary phases in different zones
226 of the Nanping No. 31 pegmatite. Primary apatites are more abundant, but secondary
227 apatites are more mineralogically diverse. Primary apatites generally form light green
228 euhedral to subhedral or granular crystals up to 20 centimeters long; the sizes of the
229 apatite crystals increase from the outer zones inward. Secondary apatites generally

230 occur as the alteration products of primary and/or early phosphate minerals and were
231 deposited during the different hydrothermal stages of the Nanping No. 31 pegmatite
232 (Fig. 3 and Fig. 4). They generally form irregular aggregates and/or veinlets, which
233 are intergrown with secondary quartz and fine-grained muscovite among the alteration
234 assemblages of montebrasite. In some cases, Sr-rich apatites replace early apatites,
235 forming veinlets, rims or mosaic-like phases in the hydrothermal alteration mineral
236 assemblages formed from montebrasite. Electron microprobe analyses show that
237 primary apatites are enriched in F and Mn (up to 4.79 wt.% F and 5.42 wt.% MnO,
238 respectively), which correspond to Mn-rich fluorapatite (Table 3 and Fig. 5). In
239 contrast, secondary apatites are enriched in Sr (with an average value of 1.67 wt.%
240 SrO, but ranging up to 12.8 wt.% SrO) and are classified as hydroxy-fluorapatite
241 (Table 3). The variation of SrO and MnO/FeO in secondary apatites is shown in
242 Figure 5.

243 **5.3. Triphylite $LiFe(PO_4)$**

244 Both primary and secondary triphylite have been found in the Nanping No. 31
245 pegmatite (Rao et al., 2014a). Primary triphylite forms small granular aggregates
246 approximately 80 to 200 μm in diameter in the rocks of zones III and IV and is
247 associated with lazulite, wagnerite, fluorarrojadite-(BaNa), palermoite, and
248 fluorapatite. Secondary triphylite occurs as veinlets or aggregates along the fractures
249 and rims of primary montebrasite (Fig. 4a and Fig. 6a) and quartz (Fig. 6b) in zone IV.
250 Triphylite veinlets are up to 1 mm wide and several cm long. They are associated with
251 fine-grained muscovite, fluorapatite, ludlamite, anapaite, wagnerite and
252 fluorarrojadite-(BaNa). Some Nb-Ta-bearing oxide (tantelite) crystals are observed in
253 the triphylite assemblage (Fig. 6a). The chemical features of triphylite have been
254 described in our previous study (Rao et al., 2014a).

255 **5.4. Wagnerite $Mg_2(PO_4)F$**

256 Wagnerite occurs in two different modes in the Nanping No. 31 pegmatite. It
257 occurs as one of the hydrothermal alteration products (denoted wag-1) of triphylite
258 (Fig. 6b, and Fig. 5a: Rao et al., 2014a) and as anhedral crystals (denoted wag-2)
259 along fractures developed in montebrasite in zone IV (Fig. 6c). The representative

260 compositions of wagnerite are given in [Table 4](#). Compared to the anhedral crystals of
261 wagnerite (wag-2), which contain, on average, 7.77 wt.% F, 5.61 wt.% FeO, 0.79
262 wt.% MnO and 45.29 wt.% MgO, irregular assemblages of wagnerite (wag-1) have
263 clearly higher concentrations of F, Mn and Fe (with average values of 11.08 wt.% F,
264 4.20 wt.% MnO and 11.24 wt.% FeO) but lower concentrations of Mg (with an
265 average of 37.74 wt.% MgO).

266 **5.5. Lazulite $MgAl_2(PO_4)_2(OH)_2$**

267 In hand specimens, lazulite appears as blue granular aggregates along the fractures
268 of primary montebrasite. Based on their petrological textures and chemical
269 compositions, three different types of lazulite were distinguished. The first type of
270 lazulite (denoted laz-1) is characterized by subhedral to euhedral crystals up to 150
271 μm long, which are included in the secondary montebrasite assemblage from zone IV
272 ([Fig. 4b](#)). They are closely associated with palermoite, bertossaite, augelite,
273 fluorapatite and fluorarrojadite-(BaNa). In back-scattered electron (BSE) images, they
274 are weakly heterogeneous, consisting of intergrown brighter and darker areas; the
275 brighter areas are slightly enriched in Fe compared to the darker areas. The second
276 type of lazulite (denoted laz-2) occurs as anhedral grains, irregular assemblages
277 around primary montebrasite crystals ([Figs 3b, 4c and 6d](#)), or along fractures in the
278 central portions of granular aggregates of triphylite ([Fig. 5a: Rao et al., 2014a](#)).
279 Wagnerite, fluorarrojadite-(BaNa), fluorapatite, palermoite and bertossaite were also
280 observed in these assemblages. Lazulite veinlets (denoted laz-3) represent the third
281 type of lazulite, which occurs along cleavage faces or fractures in montebrasite in
282 zone IV ([Fig. 6e](#)). These veinlets are approximately 1 μm wide but up to hundreds of
283 μm long and are rimmed by fine-grained muscovite and palermoite.

284 The results of electron microprobe analyses indicate that the P content of lazulites
285 is slightly less than the ideal value, but their Al content is close to or slightly higher
286 than the ideal value ([Table 4](#)). Lazulite-1 has relatively high Fe concentrations but low
287 Mg concentrations, with contents of approximately 11.09 wt. % FeO and 6.9 wt. %
288 MgO, which are indicative of Fe-rich lazulite; the chemical compositions of lazulite-2
289 and lazulite-3 are similar ([Table 4](#)). The MgO contents of these lazulites range from

290 9.10 to 11.77 wt. % (with an average value of 10.96 wt. %). Their FeO contents range
291 from 2.30 to 6.81 wt. % (with an average value of 4.04 wt. %), with minor
292 concentration of K, Na, Mn, Ti, Sr, Ba, Si, and Ca but no F.

293 **5.6. Palermoite-bertossaite $SrLi_2Al_4(PO_4)_4(OH)_4 - CaLi_2Al_4(PO_4)_4(OH)_4$**

294 Palermoite and bertossaite are end-members of a Sr-Ca isomorphous series with an
295 ideal chemical formula of $Li_2(Sr,Ca)Al_4(PO_4)_4(OH)_4$, which occur as secondary
296 phases of montebrasite in zones III and IV of the Nanping No. 31 pegmatite.
297 Palermoite commonly forms fine grains or nodules up to 600 μm wide, and is
298 distributed among fine-grained muscovite. Lazulite, secondary montebrasite, kulanite,
299 minjiangite, goyazite and fluorapatite are also observed in these assemblages. In some
300 cases, irregular assemblages, approximately 1 to 2 cm in diameter, are observed
301 around montebrasite crystals in zone IV; additionally, palermoite veinlets are located
302 in the rims of primary montebrasite in zone III (Fig. 3b) or as intergrowths with
303 fine-grained muscovite in the fractures developed in primary montebrasite in zone IV
304 (Fig. 6e and Fig. 7b)

305 Bertossaite formed slightly later than palermoite in the Nanping No. 31 pegmatite.
306 It generally occurs as irregular assemblages around palermoite (Fig. 7a) or forms
307 veinlets crosscutting montebrasite (Fig. 7c). Its associated minerals include kulanite,
308 goyazite, hydroxylapatite, and fine-grained muscovite. Bertossaite is also observed as
309 irregular assemblages up to 2 cm in diameter or intergrown with secondary
310 montebrasite (mtb-3) among the alteration products of primary montebrasite in zone
311 IV (Fig. 4d). In some cases, bertossaite crystals were observed to be altered to augelite
312 (Fig. 7d). Electron microprobe analyses results indicate that palermoite and
313 bertossaite are members of a continuous isomorphous series of (Sr, Ca); they contain
314 0.19 to 13.10 wt% SrO and 0.83 to 8.89 wt% CaO (Table 4).

315 **5.7. Kulanite $BaFe^{2+}_2Al_2(PO_4)_3(OH)_3$**

316 Kulanite generally occurs as small grains distributed among the hydrothermal
317 alteration products of massive montebrasite in zone IV. It occurs along the fractures
318 developed in lazulite, or forms veinlets up to 1 centimeter long within the assemblage
319 of bertossaite + montebrasite (Fig. 7c). Fluorapatite, goyazite, secondary montebrasite,

320 palermoite, and fine-grained muscovite are found in this assemblage. Chemically,
321 kulanite contains 15.70 to 16.99 wt.% Al₂O₃, 33.62 to 35.41 wt.% P₂O₅, 22.01 to
322 25.52 wt.% BaO, 6.59 to 12.95 wt.% FeO, 5.20 to 12.78 wt.% MnO, 0.27 to 3.01
323 wt.% MgO, < 2.50 wt.% SrO, < 1.72 wt.% TiO₂, and minor K₂O, Na₂O, CaO, SiO₂
324 and F (Table 4).

325 **5.8. Simferite $Li(Mg,Fe^{3+},Mn^{3+})_2(PO_4)_2$**

326 Simferite occurs as small orange grains or irregular assemblages along the rims of
327 primary montebrasite crystals in zone IV (Fig. 6f). The sizes of the simferite grains
328 range from 50 to 100 μm in diameter. Its associated minerals contain secondary
329 montebrasite, fluorapatite, ludlamite and fine-grained muscovite. The results of
330 electron microprobe analyses indicate that simferite contains average
331 values of 50.86 wt.% P₂O₅, 18.66 wt.% MgO, 14.72 wt.% FeO (16.34 wt.% Fe₂O₃),
332 and 5.38 wt.% MnO (5.97 wt.% Mn₂O₃), as well as minor amounts of K₂O, Na₂O,
333 CaO, SrO, BaO, SiO₂, and F (< 1 wt.% in total).

334 **5.9. Accessory phosphates**

335 *Goyazite* SrAl₃(PO₄)₂(OH)₅(H₂O) is a relatively late secondary phosphate phase
336 in the Nanping No. 31 pegmatite. It occurs as veinlets up to hundreds of μm long that
337 fill the interstices between albite crystals in zone II, as well as fractures developed in
338 bertossaite in zone IV. Its associated minerals are hydroxylapatite, kulanite,
339 palermoite, secondary montebrasite and fine grained muscovite. Goyazite contains
340 13.92 to 20.19 wt.% SrO and up to 3.51 wt.% BaO (Table 4).

341 *Augelite* Al₂(PO₄)(OH)₃ occurs as one of the hydrothermal alteration products of
342 montebrasite in the Nanping No. 31 pegmatite. It forms euhedral to subhedral crystals,
343 and occurs, together with secondary montebrasite, palermoite, and fluorapatite,
344 around primary montebrasite in zone IV. However, augelite also occurs as one of the
345 alteration products of bertossaite in zone IV (Fig. 7d); some remnants of bertossaite
346 are sporadically distributed in the augelite assemblages. Chemically, augelite contains
347 approximately 36 wt. % P₂O₅ and 50 wt. % Al₂O₃, with low concentrations of minor
348 elements (total < 1 wt. %), suggesting that it is an end-member phase.

349 *Beryllonite* NaBePO₄ was detected by microprobe analysis in one thin section
350 from zone I. It forms anhedral crystals up to 30 μm long between quartz and
351 muscovite, and is associated with strontiohurlbutite, hydroxylapatite, and
352 hydroxylherderite. Chemically, beryllonite contains 57.06 wt.% P₂O₅ and 20.11 wt.%
353 Na₂O, as well as small amounts of K₂O, FeO, MnO, MgO, CaO, SrO, BaO, and SiO₂
354 (< 1 wt.% in total).

355 *Phosphoferrite* occurs as irregular assemblages among the alteration products of
356 montebrasite. Its chemical composition data indicate that phosphoferrite has an
357 empirical formula corresponding to (Fe²⁺_{2.74}Mn_{0.26}Mg_{0.03})_{Σ3.03}(PO₄)₂·3(H₂O).
358

359 **6. Alteration assemblages of montebrasite**

360 Both primary montebrasite and secondary montebrasite (Mtb-1) in the Nanping
361 No. 31 pegmatite were strongly affected by late hydrothermal fluids, which produced
362 different secondary phases. Here, five distinct types of alteration assemblages of
363 montebrasite were distinguished and are described below.

364 ***Triphylite ± wagnerite + fluorapatite assemblage.*** This alteration assemblage of
365 montebrasite mainly consists of triphylite, wagnerite and fluorapatite in zone IV. This
366 type of triphylite is restricted to primary montebrasite; it occurs as fine-grained
367 crystals along fractures within montebrasite. Wagnerite typically surrounds crystals of
368 triphylite (Fig. 6b). In some cases, montebrasite has been replaced by wagnerite (Fig.
369 6c). Fluorapatite, muscovite and fluorarrodite are also observed in this assemblage.

370 ***Lazulite + palermoite + fluorapatite assemblage.*** Lazulite is one of the most
371 common alteration products of montebrasite in the Nanping No. 31 pegmatite. It
372 generally occurs as irregular aggregates (Fig. 6d) or veinlets (Fig. 6e) along the rims
373 or fractures of montebrasite. Palermoite formed later than lazulite; it occurs as veinlets
374 surrounding the aggregates of lazulite + montebrasite in zone III (Fig. 3b), or the rims
375 of the alteration products of montebrasite in zone IV (Fig. 6e), or as irregular
376 aggregates crosscutting the early alteration products of montebrasite (Fig. 6d). In this

377 assemblage, fluorapatite is the youngest phase, as it crosscuts both lazulite and
378 palermoite.

379 ***Palermoite + bertossaite + kulanite + fluorapatite assemblage.*** In the Nanping
380 No. 31 pegmatite, both palermoite and bertossaite occur as alteration products of
381 montebrasite along the rims and fractures of primary montebrasite (Fig. 4d, Fig. 7a
382 and Fig. 7c). Palermoite veinlets are commonly observed among the alteration
383 products of montebrasite (Fig. 3b, Fig. 6e and Fig. 7b). In addition, palermoite is
384 generally replaced by bertossaite along its rims and fractures (Fig. 7a and Fig. 7d).
385 Kulanite, fluorapatite, goyazite and minjiangite also occur in this assemblage.

386 ***Simferite + fluorapatite assemblage.*** In addition to triphylite, palermoite and
387 bertossaite, simferite is another secondary Li phosphate formed from montebrasite. It
388 occurs as small orange grains among the fine-grained muscovite formed from the
389 primary montebrasite in zone IV (Fig. 6f). Fluorapatite generally surrounds the orange
390 grains or occurs as irregular aggregates in this assemblage.

391 ***Fluorapatite + akdalaite assemblage.*** Apatites are common phases produced by
392 the hydrothermal alteration of montebrasite. They commonly occur as veinlets
393 crosscutting earlier phases, or as irregular aggregates along the rims and fractures of
394 the alteration products of montebrasite. In zone IV, some montebrasite crystals are
395 notably rimmed by veinlets that are approximately 50 to 200 μm wide (Fig. 8a and
396 Fig. 8b). Under crossed polarizers, they are isotropically black (Fig. 8a). The results
397 of electron microprobe analyses indicate that these veinlets are mainly composed of
398 CaO, Al₂O₃, P₂O₅ and F. However, scanning electron microscope (SEM) analyses
399 indicate that they are composed of brighter and darker regions, which form
400 micron-scale networks (Fig. 8c and Fig. 8d). The powder X-ray diffraction patterns
401 observed by in-situ micro-diffraction analyses was shown in Figure 9, which appear to
402 display two differently indexed X-ray diffraction patterns. The brighter phase, which
403 has six distinct lines [d in $\text{\AA}(I)(hkl)$]: 2.803(100)(211), 2.776(81)(112), 2.706(29)(300),
404 1.936(34)(222), 1.837 (39)(213) and 1.797(14)(321), corresponds to fluorapatite with
405 $a = 9.370 (2) \text{\AA}$, $c = 6.877 (2) \text{\AA}$, and $V = 522.9 \text{\AA}^3$. The darker phase has eight distinct
406 lines [d in $\text{\AA}(I)(hkl)$]: 2.502(34)(006), 2.317(52)(412), 2.114(94)(324), 1.862(20)(334),

407 1.681(24)(523), 1.418(100)(445), 1.394(56)(329) and 1.215(51)(20 12) and unit cell
408 parameters of $a = 12.889(9) \text{ \AA}$, $c = 14.949(7) \text{ \AA}$, and $V = 2150.83 \text{ \AA}^3$, which is
409 indicative of akdalaite, $(\text{Al}_2\text{O}_3)_4 \cdot \text{H}_2\text{O}$. These patterns and peak data are in agreement
410 with those of fluorapatite from ICDD pdf No. 15-0876 and akdalaite from ICDD pdf
411 No. 25-0017, respectively.

412

413 **7. Discussion**

414 **7.1. Genetic sequence of primary phosphates**

415 Phosphate minerals are common accessory phases in granitic bodies such as the
416 Beauvoir granite of France (Charoy, 1999), the Tanco pegmatite of Canada (Černá et
417 al., 1972), the Cañada pegmatite of Spain (Roda-Robles et al., 2004), and the Palermo
418 #2 pegmatite in the U.S.A. (Nizamoff, 2006). Montebasite-amblygonite and
419 lithiophilite-triptychite commonly occur as subhedral crystals or rounded nodules
420 embedded in rock-forming minerals within different zones in pegmatites (e.g.,
421 London and Burt, 1982; Galliski et al., 2012). The magmatic crystallization and
422 evolution of these phosphates are believed to be controlled by their solubilities and the
423 incompatible behaviour of P in granitic melts (Roda-Robles et al., 2004; Roda-Robles
424 et al., 2012). The low solubilities of phosphates in granitic melts could result in their
425 crystallization, while the incompatible behaviour of P can enhance the evolution of
426 phosphates and cause the abundance of phosphates to increase.

427 Table 1 shows the distribution and crystallization sequence of primary phosphate
428 minerals in the Nanping No. 31 pegmatite. REE-bearing phosphates (xenotime and
429 monazite) occur in the border zone (zone I) of the pegmatite (Yang et al., 1987),
430 indicating that they were formed by early magmatic crystallization. This geochemical
431 trend can be attributed to the low solubilities of xenotime and monazite in
432 peraluminous granitic melts (Montel, 1986; Wolf and London, 1995). Similar
433 occurrences of xenotime and monazite have been found in the Palermo #2 pegmatite
434 in New Hampshire, U.S.A. (Nizamoff, 2006). Primary fluorapatite, which occurs as
435 euhedral to subhedral or granular crystals, is associated with rock-forming minerals

436 such as quartz, albite and muscovite in different zones and is enriched in Mn and F
437 (Table 3 and Fig. 5), suggesting that it was formed by early magmatic crystallization.
438 This feature is in agreement with previous studies of fluorapatite from the Silbergrube
439 Aplite (SA) of the Hagendorf-Pleystein pegmatite district (Dill et al., 2008).
440 Roda-Robles et al. (2004) suggested that Mn-rich fluorapatite is formed after the
441 crystallization of xenotime. The crystallization of triphylite and montebrasite, as well
442 as spodumene, which mainly occur in zones III and IV, represents the magmatic Li
443 stage of the Nanping No. 31 pegmatite. London and Burt (1982) suggested that
444 primary Li phosphates appear to crystallize after spodumene. Triphylite generally
445 crystallizes shortly before the magmatic Li stage (Ginsburg, 1960). In the Nanping No.
446 31 pegmatite, small montebrasite crystals occur in the out zone (zone I) of the
447 pegmatite (Fig. 3a), whereas spodumene and triphylite are only found in the
448 intermediate pegmatite zones (zones III and IV) (Rao et al., 2012, 2014a). Therefore,
449 spodumene and triphylite likely began to crystallize somewhat later, but finished
450 crystallizing before the formation of montebrasite. Montebrasite crystals are massive
451 and abundant in the exocontact zone of the pegmatite core (zone IV), thus reflecting
452 the incompatible behaviour of P in granitic melts. The observed decrease in F content
453 in primary montebrasite from the outer zones inward (Table 2) is consistent with
454 magmatic differentiation.

455

456 ***7.2. Hydrothermal alteration of montebrasite***

457 In granitic pegmatites, montebrasite is generally affected by post-magmatic fluids,
458 which can initiate various metasomatic and/or dissolution-crystallization processes
459 and generate different sequences of metasomatic alterations (e.g., London and Burt,
460 1982; Baldwin et al., 2000; Galliski et al., 2012; Shirose and Uehara, 2014).
461 Montebrasite from the White Picacho pegmatites was gradually replaced by low-F
462 montebrasite, hydroxylapatite + crandallite, then by hydroxylapatite + muscovite +
463 brazilianite + augelite + scorzalite + kulanite + wyllieite, and finally by muscovite and
464 carbonate + apatites (London and Burt, 1982). Baldwin et al. (2000) suggested that
465 natromontebrasite, brazilianite, lazulite, scorzalite, goyazite, gorceixite, crandallite,

466 hydroxylapatite and muscovite are the typical alteration products of montebrasite.
467 [Nizamoff \(2006\)](#) further divided montebrasite alteration into high- and
468 low-temperature stages of metasomatic alteration, which occur under both oxidizing
469 and non-oxidizing conditions. In the Nanping No. 31 pegmatite, the textural
470 relationships of alteration assemblages ([Fig. 3](#), [Fig. 4](#), [Fig. 6](#), [Fig. 7](#) and [Fig. 8](#))
471 indicate that multiple periods of hydrothermal alteration of montebrasite occurred.
472 [Yang et al. \(1987, 1994\)](#) suggested that the emplacement temperature of the Nanping
473 No. 31 pegmatite is 420-360°C, with the end magmatic crystallization temperature
474 (273 °C). Therefore, the metasomatic alteration of montebrasite can be split into
475 high-temperature (360 to 273 °C) and low-temperature (273-100 °C) alteration stages
476 ([Fig. 10](#)).

477 During the high-temperature alteration stage, the transformation of montebrasite
478 to low-F montebrasite could have been the first alteration process of primary
479 montebrasite to occur ([London and Burt, 1982](#); [Baldwin et al., 2000](#)). This process is
480 attributed to the exchange of OH for F and occurs after the completion of magmatic
481 crystallization due to the increasing $\alpha_{\text{H}_2\text{O}}$ in post-magmatic fluids ([Galliski et al.,](#)
482 [2012](#)). Then, Li-leaching and/or $\text{Fe}^{2+}/\text{Mg}^{2+}$ cation exchange processes, caused by Fe-
483 and Mg-rich fluids, generate Fe/Mg phosphate minerals such as lazulite, wagnerite,
484 triphylite, and simferite ([Fig. 10](#)), which suggests that early-stage hydrothermal fluids
485 have relatively high activities of Mg and F. This feature is in good agreement with
486 previous studies of fluid inclusions in the Nanping pegmatite ([Yang et al., 1994](#)). The
487 alteration of montebrasite to lazulite and wagnerite involves Li leaching and
488 $\text{Fe}^{2+}/\text{Mg}^{2+}$ cation exchanges. The $\text{Fe}^{2+}/\text{Mg}^{2+}$ -rich hydrothermal fluids were possibly
489 derived from the hydrothermal alteration of pre-existing pegmatite minerals such as
490 primary triphylite ([Rao et al., 2014a](#)) The presence of triphylite veinlets in fractures
491 developed within primary montebrasite ([Fig. 4a](#) and [\(Fig. 6a\)](#)), and the occurrence of
492 simferite aggregates around primary montebrasite ([Fig. 6f](#)), reflect the dissolution of
493 montebrasite and reprecipitation of Li phases from hydrothermal fluids. The
494 occurrence of simferite, which is a typical Fe^{3+} -bearing phase, suggests that it
495 occurred under oxidizing conditions. The crystallization sequence of triphylite,

496 wagnerite and simferite suggests that the hydrothermal fluids were transformed from
497 non-oxidizing to oxidizing conditions. These Mg-/Fe-bearing phases, however, were
498 also affected by later hydrothermal fluids such as Ca-rich fluids, resulting in the
499 crystallization of ludlamite, phosphoferrite and anapaite.

500 Both primary montebrasite and secondary montebrasite (Mtb-1) during the
501 low-temperature alteration stage were subsequently affected by different Ca-, Sr-, and
502 Ba-rich hydrothermal fluids. These processes likely reflect Ca-, Sr-, and Ba-cation
503 exchanges and the K+H processes of montebrasite, which are responsible for the
504 formation of the diverse suite of secondary phosphates (Fig. 10). Apatites are the most
505 common and main phases among the hydrothermal alteration products of
506 montebrasite, indicating that Ca is dominant within hydrothermal fluids. The presence
507 of palermoite (Fig. 3b, Fig. 6e and Fig. 7b), strontiohurlbutite and goyazite, as well as
508 secondary Sr-rich apatites, reflect the replacement of montebrasite by different Sr-rich
509 fluids. Goyazite commonly crystallized as veinlets slightly later than palermoite and
510 strontiohurlbutite. The crystallization of bertossaite after palermoite (Fig. 7a and Fig.
511 7d), indicates that another exchange of Ca for Sr occurred in late hydrothermal fluids.
512 Minjiangite and fluorarrojadite-(BaNa) generally formed after the palermoite group
513 minerals in montebrasite alteration assemblages, reflecting replacements from Ba-rich
514 fluids that occurred after replacements from Sr-rich fluids. The presence of muscovite
515 as fine-grained assemblages or veinlets among alteration assemblages of montebrasite
516 corresponds to the K+H processes of montebrasite. Shirose and Uehara (2014)
517 suggested that this type of muscovite crystallized in acidic hydrothermal fluids
518 associated with pegmatites. Akdalaite was first observed as one of alteration products
519 of montebrasite. The micron-scale network structures (Fig. 8) reflect a typical process
520 of dissolution and re-precipitation (Putnis and Austrheim, 2010).

521

522 ***7.3. Li circulation in the Nanping No. 31 pegmatite***

523 In Li-bearing pegmatites, spodumene, montebrasite and triphylite group minerals
524 are the main carriers of Li (Charoy et al., 2001). Due to their instability in
525 hydrothermal fluids, these Li minerals are extensively altered in pegmatites (e.g.,

526 [London and Burt, 1982](#)). The exchange of Na for Li in spodumene is considered to
527 represent the albitization of pegmatite ([London and Burt, 1982](#); [Charoy et al., 2001](#)).
528 The Quensel-Mason sequence in triphylite group minerals involves the progressive
529 oxidation of Fe and Mn ([Fransolet, 2007](#)). These processes result in the leaching of Li
530 into hydrothermal fluids associated with pegmatites. [London and Burt \(1982\)](#)
531 suggested that some of the Li leached from the hydrothermal alteration of primary
532 Li-bearing minerals is subsequently incorporated into the fine-grained muscovite and
533 tourmaline at the exocontacts of pegmatites. In the Nanping No. 31 pegmatite, the
534 extreme Li enrichment of the magma induced the successive crystallization of
535 spodumene, triphylite and montebrasite. Spodumene and triphylite were strongly
536 affected by post-magmatic fluids ([Rao et al., 2012, 2014a](#)), but no secondary Li
537 phases were found; only late tourmaline contains up to 0.49 wt.% Li₂O ([Rao et al.,](#)
538 [2012](#)). Various secondary Li phases formed by the hydrothermal alteration of
539 montebrasite ([Fig. 10](#)), however, reflect the dissolution, transport and crystallization
540 of Li in the post-magmatic stages of the Nanping No. 31 pegmatite.

541 The occurrence and crystallization sequence of secondary Li phases suggest that
542 four Li circulations occurred in the Nanping No. 31 pegmatite. The transformation of
543 primary montebrasite to secondary montebrasite (Mtb-1) ([Fig. 4a](#)) could have been
544 the first stage of circulation of Li in the pegmatite. This process reflects the
545 dissolution and local recrystallization of Li-bearing minerals. During the
546 high-temperature alteration stage, primary montebrasite and secondary montebrasite
547 (Mtb-1) were replaced by lazulite, wagnerite, augelite and fluorapatite ([Fig. 10](#)),
548 resulting in the leaching of Li into hydrothermal fluids. These Li-rich fluids, however,
549 were likely responsible for the formation of secondary Li phases such as triphylite
550 ([Fig. 4a](#)) and simferite ([Fig. 6f](#)). Therefore, the dissolution of Li minerals by
551 Fe-/Mg-rich fluids, transport of Li and the crystallization of secondary Li minerals,
552 comprise the second stage of Li circulation. The presence of secondary montebrasite
553 (Mtb-2 and Mtb-3) ([Fig. 4b-d](#)), palermoite veinlets ([Fig. 3b](#), [Fig. 6e](#) and [Fig. 8b](#)) and
554 bertossaite veinlets ([Fig. 7a](#) and [Fig. 7c](#)) suggest a relatively late stage of Li
555 circulation in hydrothermal fluids. The replacement of bertossaite by augelite ([Fig.](#)

556 7d), and the replacement of secondary triphylite by late phases reflect the leaching of
557 Li from secondary Li minerals into the hydrothermal fluids. It must be noted that,
558 however, compared to the Li released from the primary Li minerals, large amounts of
559 Li were leached from the pegmatite by the hydrothermal fluids so far as to be carried
560 out of the pegmatite, only small amounts of Li were preserved in secondary
561 Li-bearing minerals.

562 In the Nanping No. 31 pegmatite, it must be noted that Nb-, Ta-, Sn- and
563 Be-bearing minerals increase with abundances from outer zones inward in parallel
564 with the enrichment of Li (Rao et al., 2009). These rare-metal-bearing minerals were
565 traditionally considered as magmatic phases in pegmatites such as the Tanco
566 pegmatite of Canada (Černá et al., 1972) and the Palermo #2 pegmatite in the U.S.A.
567 (Nizamoff, 2006). However, small tantalite crystals occur in the secondary triphylite
568 formed from primary montebrasite (Fig. 6a). This textural relationship suggests that
569 the crystallization of tantalite is related to the high-temperature alteration of
570 montebrasite by Fe-/Mg-rich hydrothermal fluids. Indeed, a network of veinlets of
571 secondary tantalite has been observed in the Nanping No. 31 pegmatite (Rao et al.,
572 2009), which is suggestive of the existence of Ta-rich fluids. In this case, the
573 associated spodumene was also strongly affected by late hydrothermal fluids,
574 resulting in the leaching of Li into the hydrothermal fluids. This implies that the
575 replacement of Li phases during the process of metasomatic alteration promotes the
576 enrichment of Ta in hydrothermal fluids but that the Ta crystallized before secondary
577 Li phases. Therefore, the enrichment of Li during the hydrothermal stages of the
578 pegmatite could possibly increase the solubility of Ta in hydrothermal fluids (Zaraisky
579 et al., 2010) .

580

581 ***7.4. Transport of Al in the hydrothermal fluids***

582 Aluminium generally behaves as an immobile element under hydrothermal
583 conditions (Carmichael, 1969; Salvi et al., 1998). During metasomatic processes,
584 therefore, Al is transferred from one phase to other phases (Kister et al., 2006). In the
585 Nanping No. 31 pegmatite, the hydrothermal alteration of montebrasite resulted in the

586 formation of various Al-bearing phosphate minerals, such as palermoite, bertossaite,
587 lazulite, kulanite, goyazite, crandallite, augelite and fluorarrojadite, indicating the
588 local transformation of Al-bearing phases by hydrothermal fluids. Although non-Al
589 phosphate phases, such as apatites, triphylite, wagnerite, and anapaite, are present in
590 the alteration assemblages, they are associated with fine-grained muscovite (Fig. 3a,
591 Fig. 6a, Fig. 6b and Fig. 6e). These features indicate that Al cannot be transported
592 long distances by late-stage hydrothermal fluids.

593 Akdalaite (Al_2O_3)₄·H₂O was first reported by Shpanovet et al. (1970); it occurs as
594 veinlets cutting through amesite-fluorite-muscovite rock and fluorite-magnetite-
595 diopside-vesuvianite skarn from Kazakhstan. After this initial description, no natural
596 akdalaite was reported. In the Nanping No. 31 pegmatite, however, akdalaite and
597 fluorapatite form micro-network phases as the alteration products of montebrasite (Fig.
598 8). This textural relationship supports the immobile geochemical behaviour of Al in
599 the hydrothermal fluids. In fact, corundum produced by spodumene alteration has also
600 been observed in the Nanping No. 31 pegmatite (Rao et al., 2012).

601

602 **8. Conclusions**

603 (1) Primary phosphate minerals from the Nanping No. 31 pegmatite evolved from
604 xenotime and monazite to Mn-rich fluorapatite and then to triphylite and
605 montebrasite, reflecting the increasing activities of both Li and P in peraluminous
606 granitic melts. The formation of secondary phosphate minerals is mainly related
607 to the metasomatic alteration of montebrasite.

608 (2) Montebrasite experienced high-temperature (360 to 273 °C) and low-temperature
609 (273-100 °C) hydrothermal alteration stages. The high-temperature hydrothermal
610 alteration of montebrasite mainly corresponds to Li-leaching and/or (Fe,Mg)²⁺
611 cation exchange processes, which produced a series of Fe-/Mg-bearing
612 phosphates. The low-temperature hydrothermal alteration of primary
613 montebrasite and secondary montebrasite (Mtb-1) is related to Ca-, Sr-, and
614 Ba-cation exchanges and K + H processes in the hydrothermal fluids, resulting in

615 the formation of Ca-, Sr-, Ba-bearing phosphate minerals, muscovite and
616 akdalaite. The hydrothermal fluids underwent a transformation from
617 non-oxidizing conditions to oxidizing conditions.

618 (3) Montebrasite, triphylite and spodumene, which are the magmatic carriers of Li in
619 the Nanping No. 31 pegmatite, were strongly affected by late hydrothermal fluids.
620 Four stages of the post-magmatic circulation of Li were proposed: first, secondary
621 montebrasite (Mtb-1) reflects the local circulation of Li; then, Li derived from the
622 high-temperature hydrothermal alteration of montebrasite by Fe-Mg-bearing
623 fluids is combined with Fe and Mg to form secondary Li-bearing phases such as
624 triphylite and simferite; then, Li derived from the low-temperature hydrothermal
625 alteration of montebrasite by Sr-Ca-rich fluids is involved in the formation of
626 secondary palermoite, bertossaite and montebrasite (Mtb-2 and Mtb-3); finally,
627 the replacement of secondary triphylite, montebrasite, palermoite and bertossaite
628 by later phases results in the leaching of Li back into hydrothermal fluids. The
629 re-enrichment of Li during the hydrothermal stages could have increased the
630 solubility of Ta in the hydrothermal fluids derived from the granitic pegmatite.

631 (4) Al derived from the hydrothermal alteration of montebrasite was combined with
632 other ions into secondary Al minerals such as lazulite, palermoite, bertossaite,
633 goyazite, and muscovite. The micro-networks of akdalaite $(\text{Al}_2\text{O}_3)_4 \cdot \text{H}_2\text{O}$, which
634 were first found as one of the secondary phases of montebrasite, clearly
635 demonstrate the low mobility of Al in hydrothermal fluids.

636

637 **Acknowledgements** This paper benefitted from critical comments of H.G. Dill, P.
638 Tomascak, and Editor F. Pirajno. Their careful comments and helpful suggestions
639 improved greatly our manuscript. Financial support for the research was provided by
640 NSF of China (Grant No. 41472036) and the Fundamental Research Funds for the
641 Central Universities.

642

643 **References**

- 644 Bajjot, M., Hatert, F., Philippo, S., 2012. Mineralogy and geochemistry of phosphates and silicates
645 in the Sapucaia pegmatite, Minas Gerais, Brazil: genetic implications. *Can. Mineral.* 50,
646 1531–1554.
- 647 Baldwin, J.R., Hill, P.G., Von Knorring, O., Oliver, G.J.H., 2000. Exotic aluminum phosphates,
648 natromon-tebrasite, brazilianite, goyazite, gorceixite and crandallite from rare-element
649 pegmatites in Namibia. *Mineral .Mag.* 64, 1147–1164.
- 650 Carmichael, D.M., 1969. On the mechanism of prograde metamorphic reactions in quartz-bearing
651 pelitic rocks. *Contrib. Mineral. Petrol.* 20, 244–267.
- 652 Černá, I., Černý, P., Ferguson, R.B., 1972. The Tanco pegmatite at Bernic Lake, Manitoba; III,
653 Amblygonite-montebbrasite. *Can. Mineral.* 11, 643–659.
- 654 Černý, P., Ercit, T.S., 2005. The classification of granitic pegmatites revised. *Can. Mineral.* 43,
655 2005–2026.
- 656 Charoy, B., 1999. Beryllium speciation in evolved granitic magmas: phosphates versus silicates.
657 *Eur. J. Mineral.* 11, 135–148.
- 658 Charoy, B., Noronha, F., Lima, A., 2001. Spodumene-petalite-eucryptite mutual relationships and
659 pattern of alteration in Li-rich aplite-pegmatite dikes from northern Portugal. *Can. Mineral.*
660 39, 729–746.
- 661 Dill, H.G., Weber, B., Gerdes, A., Melcher, F., 2008. The Fe-Mn phosphate aplite ‘Silbergrube’
662 near Waidhaus, Germany: epithermal phosphate mineralization in the Hagendorf-Pleystein
663 pegmatite province. *Mineral. Mag.* 72(5), 1119–1144.
- 664 Elliott, T., Jeffcoate, A., Bouman, C., 2004. The terrestrial Li isotope cycle: light-weight
665 constraints on mantle convection. *Earth Planet Sci. Lett.* 220, 231–245.
- 666 Elliott, T., Thomas, A., Jeffcoate, A., Niu, Y., 2006. Lithium isotope evidence for
667 subduction-enriched mantle in the source of midocean-ridge basalts. *Nature* 443, 565–568.
- 668 Fransolet, A.M., 2007. Phosphate associations in the granitic pegmatites: the relevant significance
669 of these accessory minerals. *Granitic Pegmatites: The State of the Art - International*
670 *Symposium.* 06th – 12th May 2007, Porto, Portugal.

671 Fransolet, A.M., Keller, P., Fontan, F., 1986. The phosphate mineral associations of the
672 Tsaobismund pegmatite, Namibia. *Contrib. Mineral. Petrol.* 92, 502–517.

673 Galliski, M.Á., Černý, P., Márquez-Zavalía, M.F., Chapman, R., 2012. An association of
674 secondary Al-Li-Be-Ca-Sr phosphates in the San Elías pegmatite, San Luis, Argentina. *Can.*
675 *Mineral.* 50, 933–942.

676 Ginsburg, A.I., 1960. Specific geochemical features of the pegmatitic process. *Proc 21st Int Geol*
677 *Congress (Norden)* 17, 111–121.

678 Grosjean, C., Miranda, P.H., Perrin, M., Poggi, P., 2012. Assessment of world lithium resources
679 and consequences of their geographic distribution on the expected development of the electric
680 vehicle industry. *Renew Sustain Energy Rev.* 16, 1735–1744.

681 Halama, R., Savov, I.P., Rudnick, R.L., McDonough, W.F., 2009. Insights into Li and Li isotope
682 cycling and sub-arc metasomatism from veined mantle xenoliths, Kamchatka. *Contrib.*
683 *Mineral. Petrol.* 158, 197–222, DOI 10.1007/s00410-009-0378-5.

684 Hatert, F., Ottolini, L., Schmid-Beurmann, P., 2011. Experimental investigation of the alluauditet
685 triphylite assemblage, and development of the Na-in-triphylite geothermometer: applications
686 to natural pegmatite phosphates. *Contrib. Mineral. Petrol.* 161, 531–546.

687 Kister, P., Laverret, E., Quirt, D., Cuney, M., Patrier Mas, P., Beaufort, D., Bruneton, P., 2006.
688 Mineralogy and geochemistry of the host-rock alterations associated with the Shea Creek
689 unconformity-type uranium deposits (Athabasca Basin, Saskatchewan, Canada). Part 2.
690 Regional-scale spatial distribution of the Athabasca Group sandstone matrix minerals. *Clay*
691 *Clay Min.* 54(3), 295–313.

692 Li, Z.X., Li, X.H., Wartho, J.A., Clark, C., Li, W.X., Zhang, C.L., Bao, C.M., 2010. Magmatic and
693 metamorphic events during the early Paleozoic Wuyi-Yunkai orogeny, southeastern south
694 China: new age constraints and pressure-temperature conditions. *Geol. Soc. Am. Bull.* 122,
695 772–793.

696 Linnen, R.L., Van Lichtervelde, M., Černý, P., 2012. Granitic pegmatites as sources of strategic
697 metals. *Elements* 8, 275–280.

698 London, D., Burt, D.M., 1982. Alteration of spodumene and lithiophilite in pegmatites of the
699 White Picacho district, Arizona. *Am. Mineral.* 67, 97–113.

700 Magma, T., Novák, M., Cempírik, J., Janoušek, V., Ullmann, C.V. and Wiechert, U. 2016.
701 Crystallographic control on lithium isotope fractionation in Archean to Cenozoic
702 lithium-cesium-tantalum pegmatites. *Geology* 44, 655-658.

703 Montel, J.M., 1986. Experimental determination of the solubility of Ce-monazite in
704 $\text{SiO}_2\text{-Al}_2\text{O}_3\text{-K}_2\text{O-Na}_2\text{O}$ melts at 800°C, 2 kbar, under H_2O -saturated conditions. *Geol.* 14,
705 659–662.

706 Moore, P.B., 1973. Pegmatite phosphates: descriptive mineralogy and crystal chemistry. *Mineral.*
707 *Rec.* 4, 103–130.

708 Nizamoff, J., 2006. The mineralogy, geochemistry and phosphate paragenesis of the Palermo #2
709 Pegmatite, North Groton, New Hampshire. University of New Orleans, Unpublished Master
710 Thesis.

711 Pogge von Strandmann, P.A.E., Henderson, G.M., 2015. The Li isotope response to mountain
712 uplift. *Geol.* 43(1), 67–70.

713 Putnis, A., Austrheim, H., 2010. Fluid-induced processes: metasomatism and metamorphism.
714 *Geofluids.* 10, 254–269.

715 Rao, C., Hatert, F., Wang, R.C., Gu, X.P., Dal Bo, F., Dong, C.W., 2015. Minjiangite,
716 $\text{BaBe}_2(\text{PO}_4)_2$, a new mineral from Nanping No. 31 pegmatite, Fujian Province, southeastern
717 China. *Mineral. Mag.* 79(5), 1195–1202.

718 Rao, C., Wang, R.C., Hatert, F., Baijot, M., 2014a. Hydrothermal transformations of triphylite
719 from the Nanping No. 31 pegmatite dyke, southeastern China, *Eur. J. Mineral.* 26, 179–188.

720 Rao, C., Wang, R.C., Hatert, F., Gu, X.P., Ottolini, L., Hu, H., Dong, C.W., Dal Bo, F., Baijot, M.,
721 2014b. Strontiohurlbutite, $\text{SrBe}_2(\text{PO}_4)_2$, a new mineral from Nanping No. 31 pegmatite,
722 Fujian Province, Southeastern China. *Am. Mineral.* 99, 494–499.

723 Rao, C., Wang, R.C., Hu, H., 2011. Paragenetic assemblages of beryllium silicates and phosphates
724 from the Nanping no. 31 granitic pegmatite dyke, Fujian province, southeastern china. *Can.*
725 *Mineral.* 49, 1175–1187.

726 Rao, C., Wang, R.C., Hu, H., Zhang, W.L., 2009. Complex internal texture in oxide minerals from
727 the Nanping No. 31 dyke of granitic pegmatite, Fujian Province, southeastern China. *Can.*
728 *Mineral.* 47, 1195–1212.

729 Rao, C., Wang, R.C., Zhang, A.C., Hu, H., 2012. The corundum + tourmaline nodules related to
730 hydrothermal alteration of spodumene in the Nanping No. 31 pegmatite dyke, Fujian
731 province, southeastern China. *Can. Mineral.* 50, 1623–1635.

732 Roda-Robles, E., Galliski, M.A., Roquet, M.B., Hatert, F., Paeseval, P.D., 2012. Phosphate
733 nodules containing two distinct assemblages in the Cema Granitic Pegmatite, San Luis
734 Province, Argentina: Paragenesis, composition and significance. *Can. Mineral.* 50(4),
735 913–931.

736 Roda-Robles, E., Pesquera, A., Fontan, F., Keller, P., 2004. Phosphate mineral associations in the
737 Cañada pegmatite (Salamanca, Spain): paragenetic relationships, chemical compositions, and
738 implications for pegmatite evolution. *Am. Mineral.* 89, 110–125.

739 Roda-Robles, E., Vieira, R., Pesquera, A., Lima, A., 2010. Chemical variations and significance of
740 phosphates from the Fregeneda-Almendra pegmatite field, Central Iberian Zone (Spain and
741 Portugal). *Miner. Petrol.* 100, 23–34.

742 Salvi, S., Pokrovski, G.S., Schott, J., 1998. Experimental investigation of aluminium-silica
743 aqueous complexing at 300°C. *Chem. Geol.* 151, 51–67.

744 Shigley, J.E., Brown Jr, G.E., 1985. Occurrence and alteration of phosphate minerals at the Stewart
745 pegmatite, Pala district, San Diego Co., California. *Am. Mineral.* 70, 395–408.

746 Shirose, Y., Uehara, S., 2014. Secondary phosphates in montebbrasite and amblygonite from
747 Nagatare, Fukuoka Prefecture, Japan. *J Mineral. Petrol. Sci.* 109(2). 103–108.

748 Shpanov, E.P., Sidorenko, G.A., Stolyarova, T.I., 1970. Akdalaite, a new hydrous modification of
749 alumina. *Zap Vses Mineral Obshch* 99, 333–339 (in Russian).

750 Tang, Y., Zhao, J.Y., Zhang, H., Cai, D.W., Lv, Z.H., Liu, Y.L., Zhang, X., 2017. Precise
751 columbite-(Fe) and zircon U-Pb dating of the Nanping No. 31 pegmatite vein in northeastern
752 Cathaysia Block, SE China. *Ore Geol. Rev.* 83, 300-311.

753 Vignola, P., Diella, V., Oppizzi, P., Tiepolo, M., Weiss, S., 2008. Phosphate assemblages from the
754 Brissago granitic pegmatite, Western Southern Alps, Switzerland. *Can. Mineral.* 46, 635–650.

755 Wang, W.Y., Yang, Y.Q., Chen, C.H., Zhu, J.H., 1999. Study on the Nb and Ta-minerals from the
756 granitic pegmatites in Nanping, Fujian Province. *Geol. Fujian* 3, 113–134 (in Chinese with
757 English abstr.).

758 Wang, Y.J., Zhang, A.M., Fan, W.M., Zhao, G.C., Zhang, G.W., Zhang, Y.Z., Zhang, F.F., Li, S.Z.,
759 2011. Kwanghsian crustal anatexis within the eastern South China Block: Geochemical, zircon
760 U-Pb geochronological and Hf isotopic fingerprints from the gneissoid granites of Wugong
761 and Wuyi-Yunkai Domains. *Lithos* 127, 239–260.

762 Wolf, B., London, D., 1995. Incongruent dissolution of REE- and Sr-rich apatite in peraluminous
763 granitic liquids: differential apatite, monazite, and xenotime solubilities during anatexis. *Am.*
764 *Mineral.* 80, 765–775.

765 Yang, Y.Q., Wang, W.Y., Chen, C.H., Zhu, J.H., 2006. Study on the cassiterite from the Granitic
766 pegmatites in Nanping, Fujian Province. *Geol. Fujian.* 25(2), 75–81.

767 Yang, Y.Q., Ni, Y.X., Guo, Y.Q., Qiu, N.M., Chen, C.H., Cai, C.F., Zhang, Y.P., Liu, J.B., Chen,
768 Y.X., 1987. Rock-forming characteristics of the Xikeng granitic pegmatites in Fujian
769 Province. *Mineral. Deposits.* 6, 10–21 (in Chinese with English abstract).

770 Yang, Y.Q., Ni, Y.X., Wang, L.B., Wang, W.Y., Zhang, Y.P., Chen, C.H., 1988. Nanpingite, a new
771 cesium mineral. *Acta Petrol Mineral Sinica.* 7(1), 49–58 (in Chinese with English abstr.).

772 Yang, Y.Q., Wang, W.Y., Lin, G.X., Zhu, J.H., Chen, C.H., 2003. Study on the potash feldspars
773 from the Granitic Pegmatites in Nanping, Fujian Province. *Geol. Fujian.* 22, 1–12 (Chinese
774 with English abstract).

775 Yang, Y.Q., Wang, W.Y., Ni, Y.X., Chen, C.H., Zhu, J.H., 1994. Phosphate minerals and their
776 geochemical evolution of granitic pegmatite in Nanping, Fujian Province. *Geol. Fujian.* 13,
777 215–226 (in Chinese with English abstract).

778 Yang, Y.Q., Wang, W.Y., Ni, Y.X., Chen, C.H., Zhu, J.H., 1995. A study on montebrasite in
779 Nanping granitic pegmatite. *Geol. Fujian.* 14, 8–21 (in Chinese with English abstract).

780 Yang, Y.Q., Wang, W.Y., Ni, Y.X., Guo, Y.Q., Zhang, Y.P., Chen, C.H., 1989. Nb, Ta-minerals in
781 Nanping pegmatites and their geological evolution. *Bulletin of the Institute of Mineral*
782 *Deposits, Chinese Academy of Geological Sciences.* 1989(1), 55–68.

783 Zaraisky, G.P., Korzhinskaya, V., Kotova, N., 2010. Experimental studies of Ta₂O₅ and
784 columbite–tantalite solubility in fluoride solutions from 300 to 550°C and 50 to 100 MPa.
785 *Mineral. Petrol.* 99, 287–300.

786

787

788 **Figure captions:**

789 **Fig. 1.** Geological map showing the distribution of the Nanping pegmatite zone, the
790 Early Paleozoic granitic and mafic plutons in the South China Block (modified after [Li](#)
791 [et al., 2010](#) and [Wang et al., 2011](#)).

792

793 **Fig. 2.** Location and simplified geological map of the Nanping pegmatite zone. (modified after
794 [Yang et al., 1987](#)).

795

796 **Fig. 3.** Backscattered-electron (BSE) images of primary montebrasite and its alteration
797 assemblages. (a) Montebrasite with fine grained muscovite and fluorapatite assemblage between
798 quartz and albite in zone I. (b) Montebrasite with lazulite, palermoite and fluorapatite assemblage
799 interstitial to coarse albite in zone III. Abbr.: Mtb - montebrasite, Apt - apatites, Ms - muscovite,
800 Ab - albite, Laz-2 - secondary lazulite, Plm - palermoite.

801

802 **Fig. 4.** Photomicrographs and BSE images of secondary montebrasite in zone IV. (a) Primary
803 montebrasite (Mtb) with secondary montebrasite (Mtb-1) and triphylite veinlet. (b) Secondary
804 montebrasite (Mtb-2) replaced Fe-rich lazulite (Laz-1) among the palermoite assemblage. (c)
805 Secondary montebrasite (Mtb-2) within the cleavages of lazulite crystals (Laz-2). (d) Secondary
806 montebrasite (Mtb-3) among bertossaite assemblage. Abbr.: Mtb - primary montebrasite, Mtb-1, 2
807 and 3 - secondary montebrasite, Trp - triphylite, Laz-1,2 - secondary lazulite, Plm - palermoite,
808 Apt - apatites, Bts - bertossaite, Ajt - arrojadite.

809

810 **Fig. 5.** Triangular Ca-Sr-(Mn+Fe) diagram showing the atomic proportions of compositional
811 variations for apatite group minerals.

812

813 **Fig. 6.** Backscattered-electron (BSE) images of the primary montebrasite and its alteration
814 assemblages in zone IV. (a) Montebrasite with triphylite + ludlamite + fluorapatite veins. (b)
815 Triphylite + wagnerite + fluorapatite + fluorarrojadite assemblages along the fracture of quartz. (c)
816 Montebrasite with wagnerite + fluorapatite assemblage. (d) Montebrasite with lazulite +
817 palermoite + fluorapatite assemblage. (e) Montebrasite with lazulite + palermoite + muscovite

818 assemblage. (f) Montebrasite with simferite + fluorapatite + muscovite assemblage. Abbr.: Mtb -
819 montebrasite, Trp - triphylite, Apt - apatites, Wag-1,2 - secondary wagnerite, Ms - muscovite, Lud
820 - ludlamite, Ftn - tantalite-(Fe), Ajt - fluorarrojadite, Laz-2,3 - secondary lazulite, Plm - palermoite,
821 Smf -simferite.

822

823 **Fig. 7.** Backscattered-electron (BSE) images of alteration assemblages of montebrasite in zone IV.
824 (a) Palermoite + bertossaite + fluorapatite assemblage in the fracture of montebrasite. (b)
825 Palermoite + muscovite veinlets within montebrasite. (c) Bertossaite + kulanite assemblage in the
826 fractures of montebrasite. (d) Palermoite + bertossaite + augelite + fluorapatite assemblage nearly
827 massive montebrasite. Abbr.: Mtb - montebrasite, Apt - apatites, Plm - palermoite, Bts - bertossaite,
828 Kln - kulanite, Agl - augelite.

829

830 **Fig. 8.** Photomicrograph, BSE and SEM images of fluorapatite + akdalaite assemblages around
831 primary montebrasite in zone IV. The photomicrograph (a) and BSE image (b) of fluorapatite +
832 akdalaite assemblages around the primary montebrasite. (c) and (d) SEM images of fluorapatite +
833 akdalaite assemblage, the brighter region: fluorapatite, the darker region: akdalaite. Abbr.: Mtb -
834 primary montebrasite, Apt - apatites, Ab - albite, Akd - akdalaite, Ms - muscovite.

835

836 **Fig. 9.** XRD pattern of of fluorapatite + akdalaite assemblage form the Nanping No. 31
837 pegmatite.

838

839 **Fig. 10.** General hydrothermal alteration sequence of montebrasite form the Nanping No. 31
840 pegmatite.

841

842

843

844

845

846

847

848 **Table captions:**

849

850 **Table 1**

851 List of the phosphate minerals and their schematic sequence in the Nanping No. 31 pegmatite.

852

853 **Table 2**

854 The chemical composition of montebrasite in the Nanping No. 31 pegmatite.

855

856 **Table 3**

857 The chemical composition of apatite group minerals in the Nanping No. 31 pegmatite.

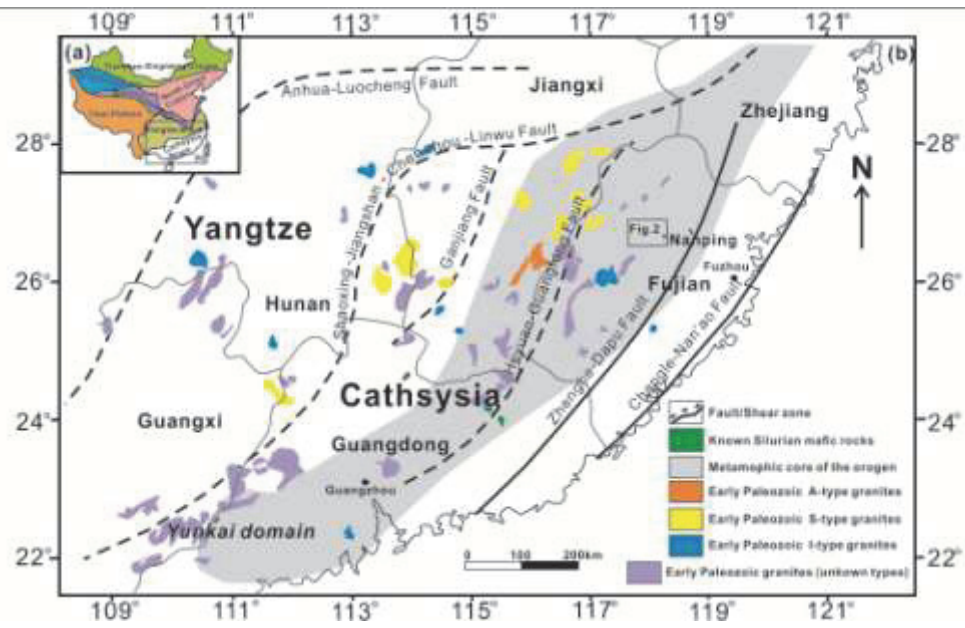
858

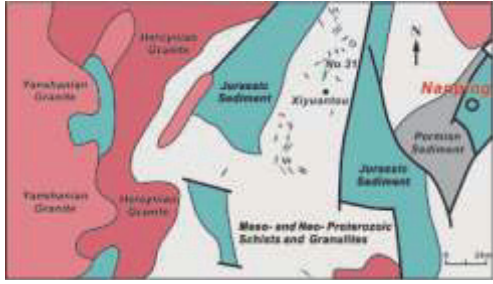
859 **Table 4**

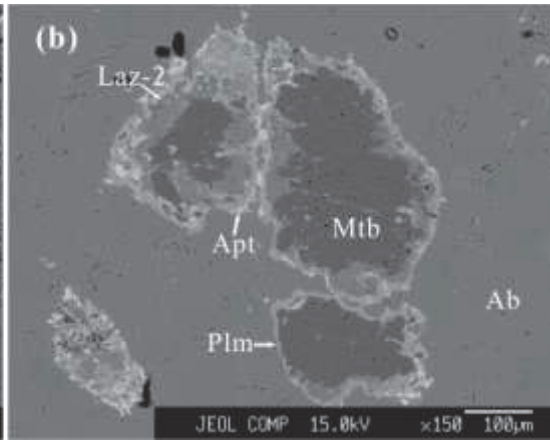
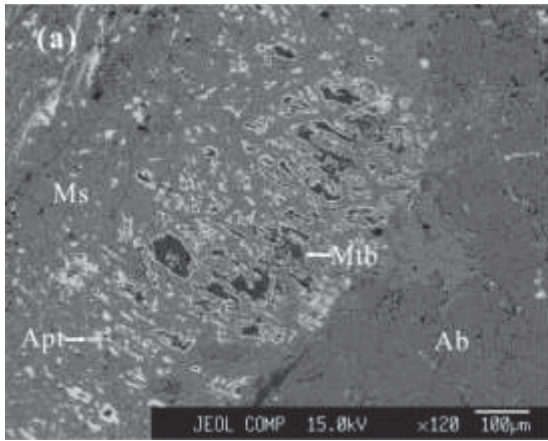
860 The chemical composition of phosphate minerals in the Nanping No. 31 pegmatite.

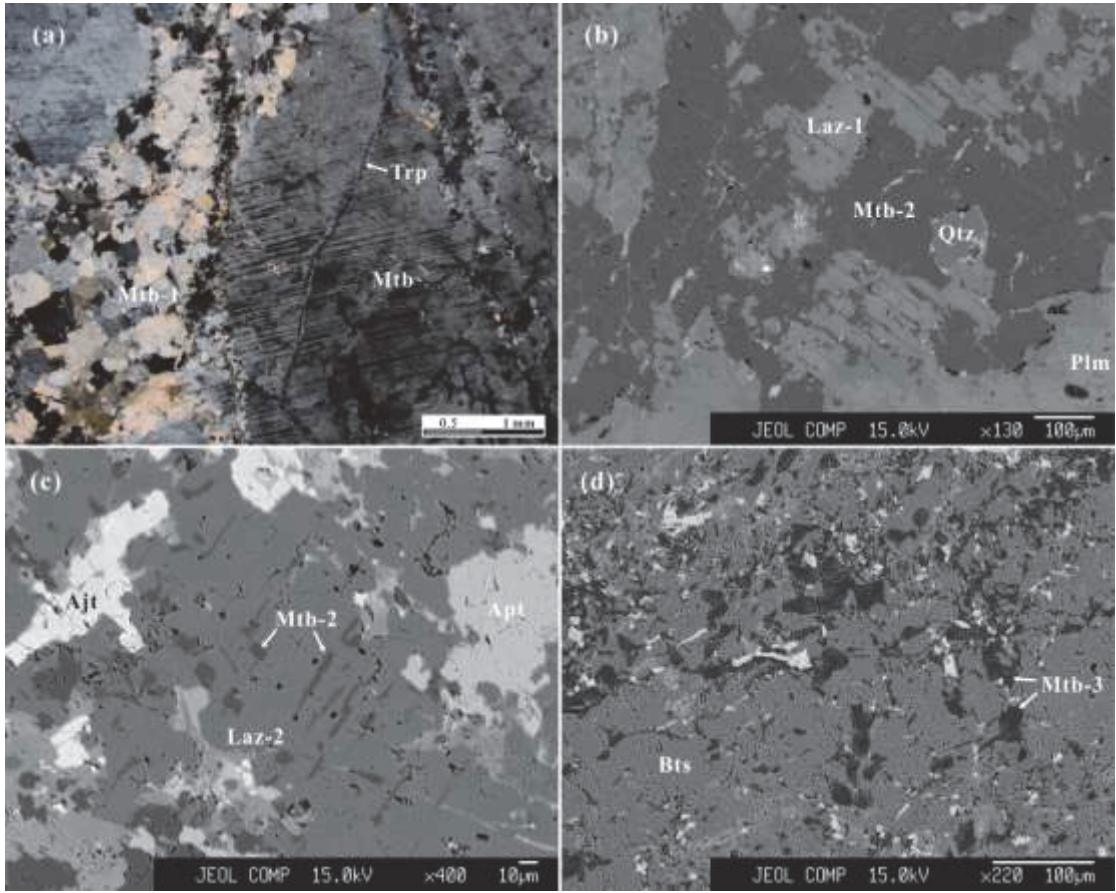
861

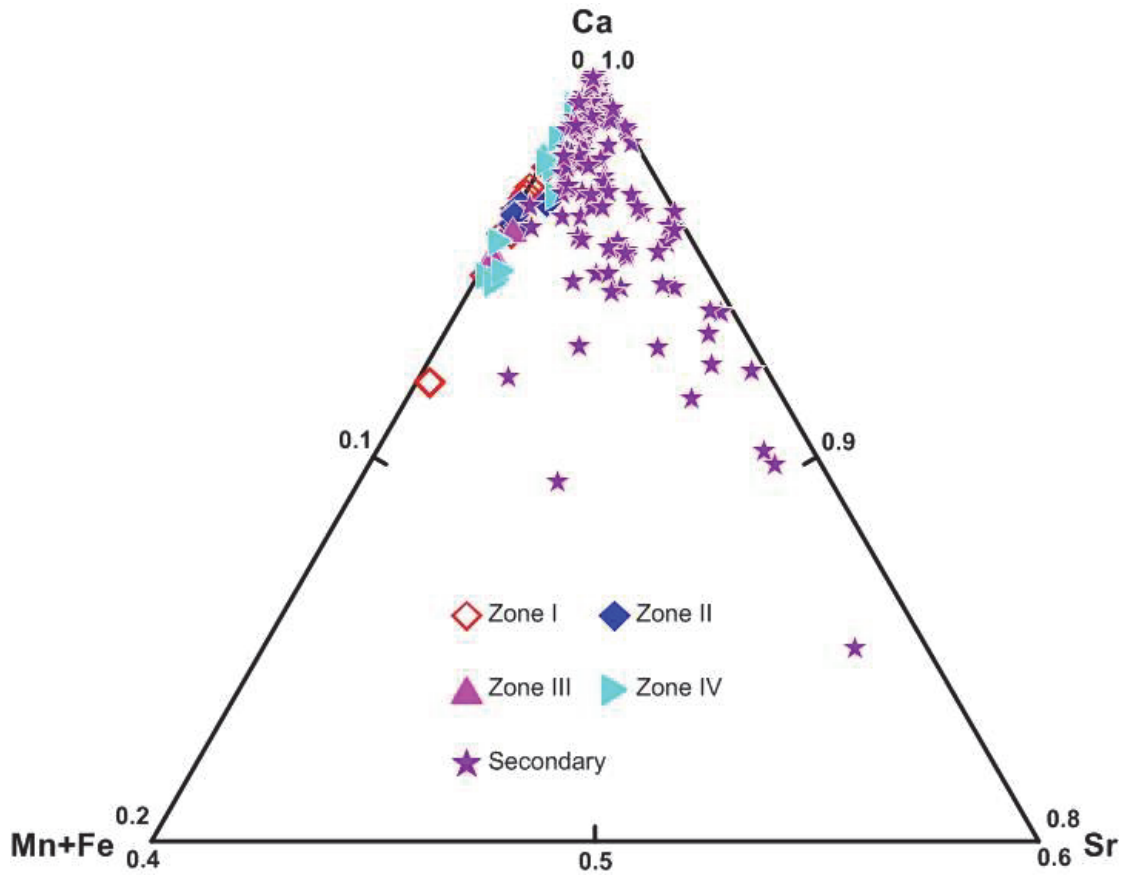
862

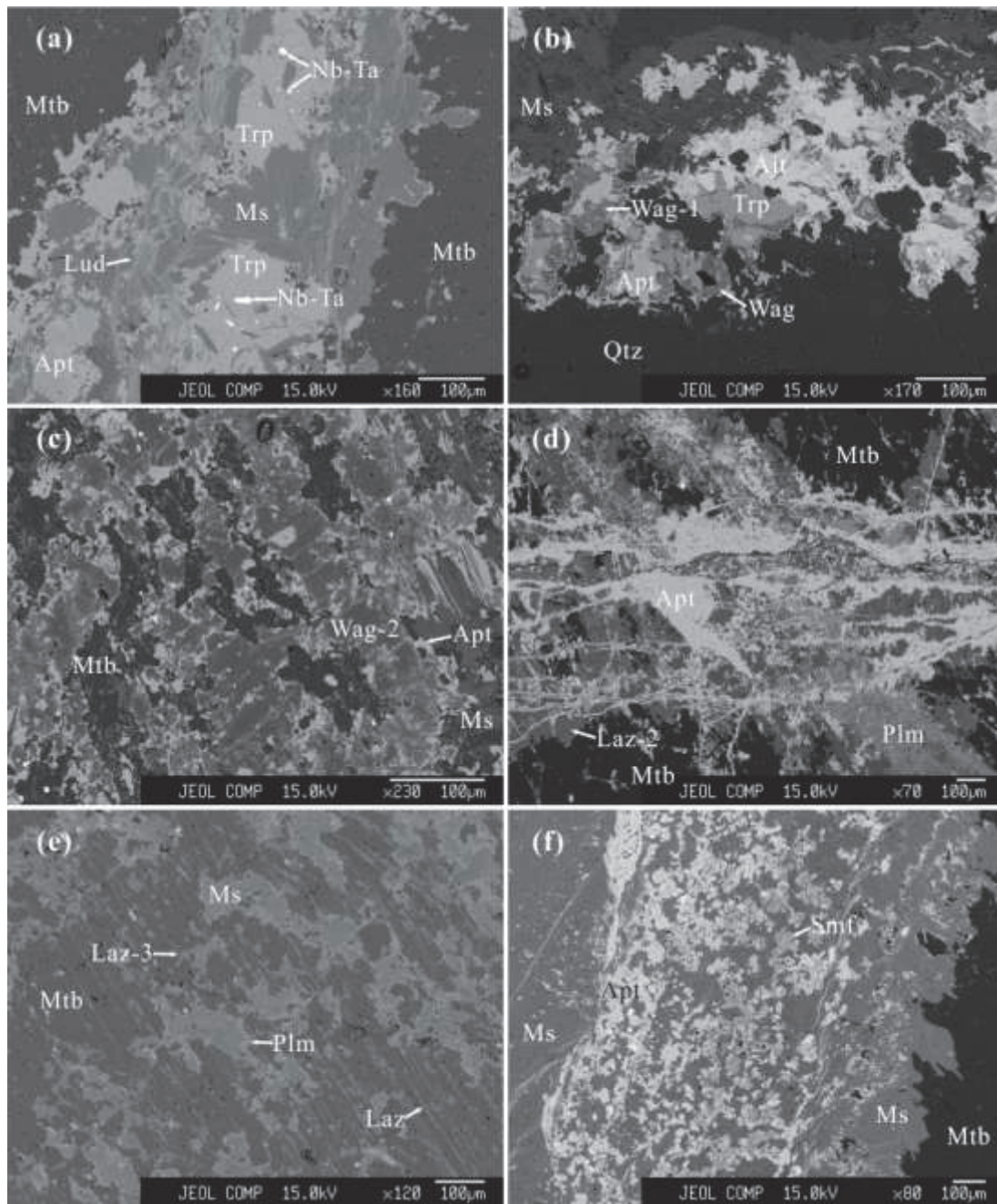


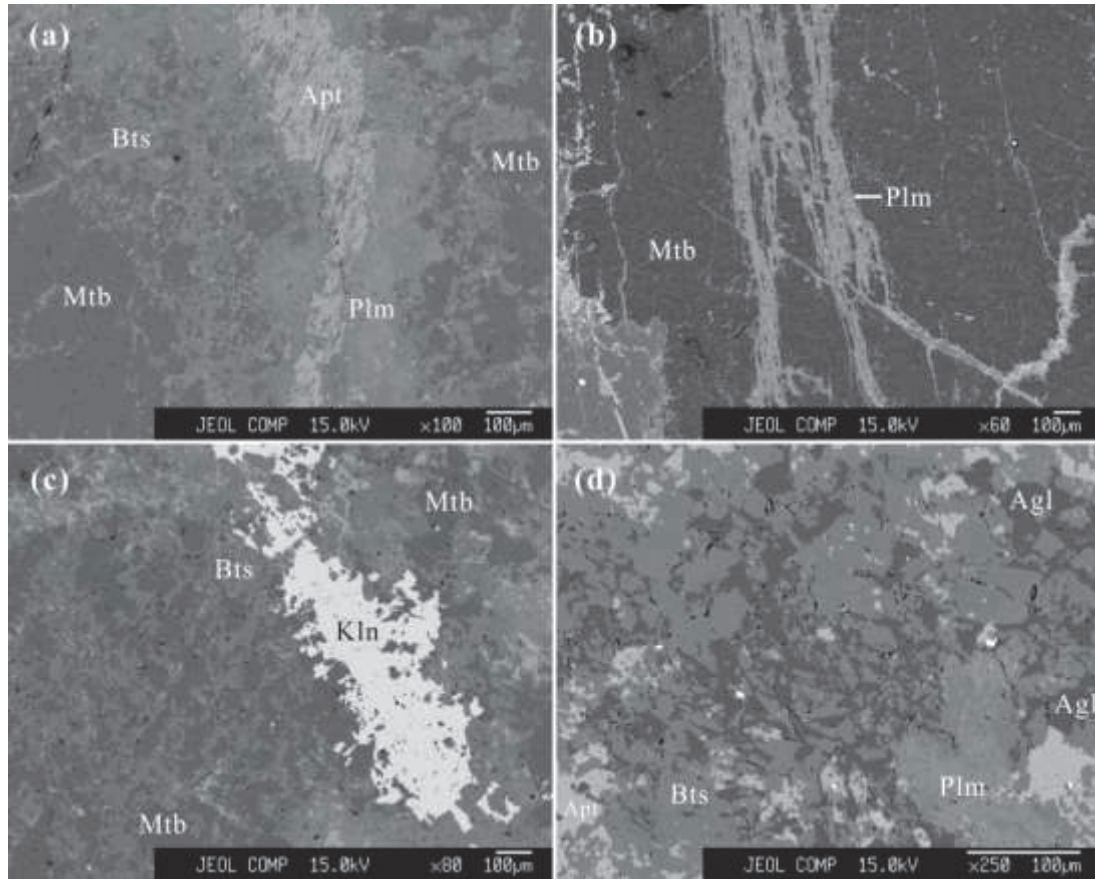


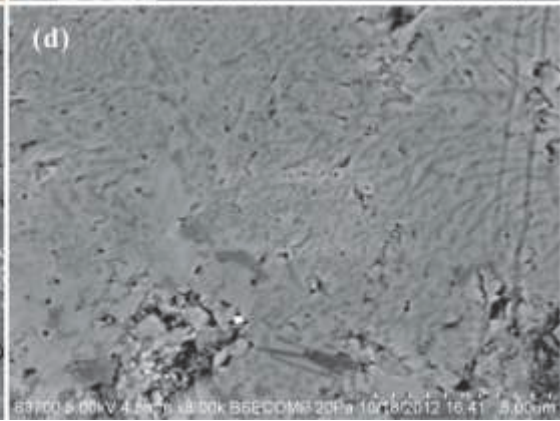
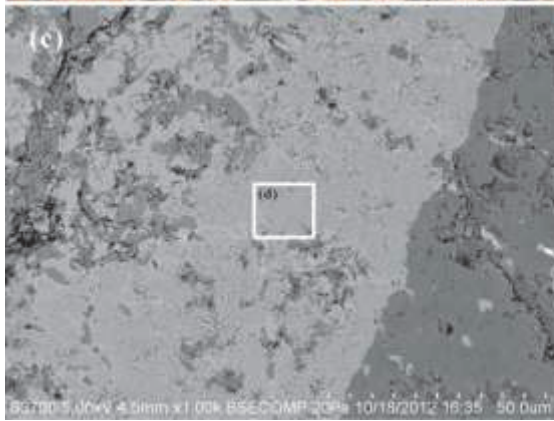
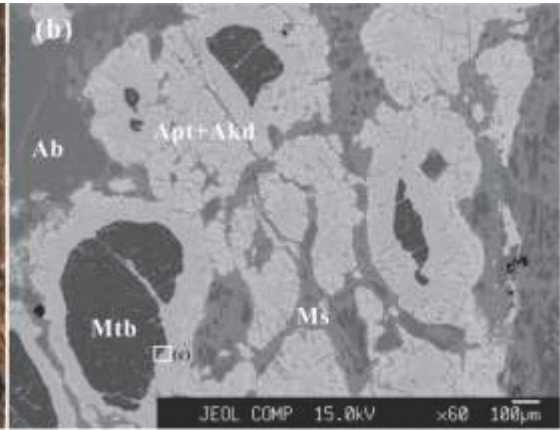
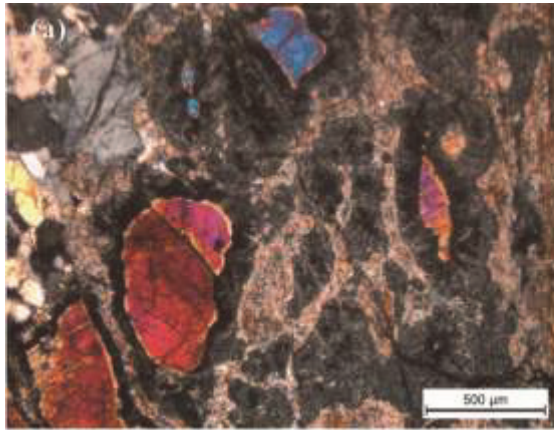


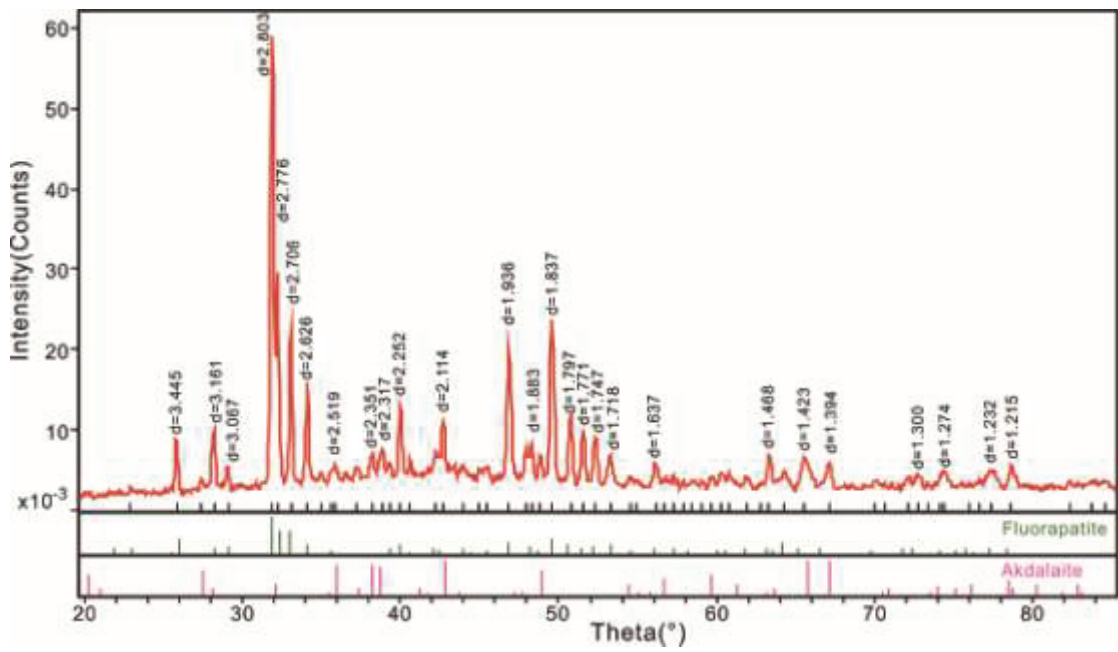












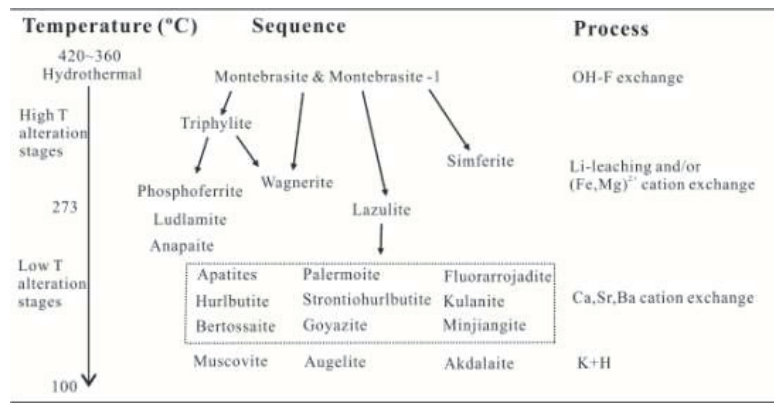


Table 1

List of the phosphate minerals and their schematic sequence in the Nanping No. 31 pegmatite.

Mineral name	Ideal Formula	Zone					Magmatic → hydrothermal
		I	II	III	IV	V	
Monozite*	(Ce,La,Nd,Th)PO ₄	x					
Xenotime*	YbPO ₄	x					
Apatites	Ca ₅ (PO ₄) ₃ (F,OH)	o	o	o	o	o	
Montebrasite	LiAl(PO ₄)(OH,F)	x		o	v		
Triphylite	LiFe(PO ₄)			#	#		
Lazulite	MgAl ₂ (PO ₄) ₂ (OH) ₂			x	o		
Augelite	Al ₂ (PO ₄)(OH) ₃			#	#		
Palermoite	SrLi ₂ Al ₄ (PO ₄) ₄ (OH) ₄	x		o	v	x	
Bertossaite	CaLi ₂ Al ₄ (PO ₄) ₄ (OH) ₄			o	v		
Goyazite	SrAl ₃ (PO ₄) ₂ (OH) ₅ ·(H ₂ O)		x	#	#		
Crandallite	CaAl ₃ (PO ₄) ₂ (OH) ₅ ·(H ₂ O)				x		
Kulanite	BaFe ²⁺ ₂ Al ₂ (PO ₄) ₃ (OH) ₃			x	#		
Hydroxylherderite	CaBe(PO ₄)(OH)	x	x				
Hurlbutite	CaBe ₂ (PO ₄) ₂	x	x		#		
Strontiohurlbutite	SrBe ₂ (PO ₄) ₂	x	x		#		
Minjiangite	BaBe ₂ (PO ₄) ₂				#		
Fluorarrojadite- (BaNa)	BaNa ₂ Ca(Fe ²⁺ ,Mn,Mg) ₁₃ Al(PO ₄) ₁₁ (PO ₃ OH)(F,OH) ₂			#	#		
Wagnerite	Mg ₂ (PO ₄)F			#	#		
Phosphoferrite	Fe ²⁺ ₃ (PO ₄) ₂ ·3(H ₂ O)				x		
Ludlamite	Fe ²⁺ ₃ (PO ₄) ₂ ·4(H ₂ O)			x	x		
Anapaite	Ca ₂ Fe ²⁺ (PO ₄) ₂ ·4(H ₂ O)			x	x		
Beryllonite	NaBePO ₄	x					
Simferite	Li(Mg,Fe ³⁺ ,Mn ³⁺) ₂ (PO ₄) ₂				x		
Eosphorite*	MnAl(PO ₄)(OH) ₂ ·(H ₂ O)			x			
Childrenite*	Fe ²⁺ Al(PO ₄)(OH) ₂ ·(H ₂ O)				x		
Vivianite*	Fe ²⁺ ₃ (PO ₄) ₂ ·8(H ₂ O)				x		
Ferrisichlerite*	Li(Fe ³⁺ ,Mn ²⁺)PO ₄				x		
Vayrynenite*	MnBe(PO ₄)(OH,F)				x		
Sicklerite*	Li(Mn ²⁺ ,Fe ³⁺)PO ₄				x		
Autunite*	Ca(UO ₂) ₂ (PO ₄) ₂ ·2-6(H ₂ O)				x		

Note: v = abundant, o = common, # = rare, x = very rare

*: not found in this study and referred to Yang et al. 1994

Table 2

The chemical composition of montebrasite in the Nanping No. 31 pegmatite.

	Primary montebrasite			Secondary montebrasite
	Zone I	Zone III	Zone IV	Zone IV
n	3	9	14	24
P ₂ O ₅ wt. %	48.79(0.36)	48.92(0.54)	48.68(0.46)	48.82(0.55)
SiO ₂	0.03(0.03)	0.03(0.02)	0.03(0.03)	0.05(0.09)
Al ₂ O ₃	34.24(0.29)	34.14(0.37)	34.37(0.39)	34.27(0.48)
FeO	0.02(0.02)	0.04(0.05)	0.04(0.05)	0.09(0.10)
MnO	0.02(0.02)	0.01(0.02)	0.01(0.02)	0.01(0.02)
MgO	0.00(0.00)	0.01(0.01)	0.00(0.00)	0.07(0.14)
TiO ₂	0.17(0.05)	0.07(0.06)	0.13(0.10)	0.04(0.04)
CaO	0.01(0.01)	0.02(0.02)	0.03(0.02)	0.04(0.07)
SrO	0.00(0.01)	0.01(0.01)	0.01(0.02)	0.01(0.01)
BaO	0.08(0.08)	0.05(0.06)	0.05(0.08)	0.05(0.07)
Na ₂ O	0.02(0.01)	0.01 (0.01)	0.01 (0.01)	0.01(0.02)
K ₂ O	0.01(0.01)	0.01(0.01)	0.01(0.02)	0.01(0.01)
F	1.59(1.33)	1.57(0.63)	1.27(0.78)	0.10(0.18)
Li ₂ O	10.26(0.08)	10.30(0.11)	10.24(0.09)	10.28(0.12)
H ₂ O	6.04(0.68)	6.07(0.32)	6.20(0.42)	6.83(0.14)
O=2F	-0.67(0.56)	-0.66(0.26)	-0.53(0.33)	-0.04(0.08)
Total	100.62(0.71)	100.59(0.71)	100.54(0.83)	100.63(1.07)
Cations per formula unit: P + Si = 1				
P <i>apfu</i>	0.999	0.999	0.999	0.999
Si	0.001	0.001	0.001	0.001
Al	0.976	0.971	0.982	0.976
Fe	0.000	0.001	0.001	0.002
Mn	0.000	0.000	0.000	0.000
Mg	0.000	0.000	0.000	0.003
Ti	0.000	0.001	0.001	0.001
Ca	0.003	0.001	0.002	0.001
Sr	0.000	0.000	0.000	0.000
Ba	0.001	0.000	0.000	0.000
Na	0.001	0.000	0.000	0.000
K	0.000	0.000	0.000	0.000
Li	0.998	0.999	0.999	0.999
F	0.122	0.119	0.097	0.008
OH	0.878	0.881	0.903	0.992

FeO as the total Fe; n: the number of analyses; standard deviation in parentheses.

Table 3

The chemical composition of apatite group minerals in the Nanping No. 31 pegmatite.

n	Primary apatites				Secondary apatites			
	zone I	zone II	zone III	zone IV				
	11	7	3	16				
P ₂ O ₅	41.69	41.56	41.61	41.66	41.05	39.30	41.00	41.38
SiO ₂	0.01	-	0.01	0.02	0.86	0.01	0.06	-
CaO	53.05	54.18	53.90	53.82	54.62	52.38	52.17	50.77
FeO	0.26	0.26	0.25	0.31	0.23	0.23	0.34	-
MnO	2.74	2.20	2.46	1.65	0.47	0.47	0.97	0.56
MgO	0.01	-	0.01	0.01	0.02	-	0.02	0.01
TiO ₂	0.00	0.00	-	0.01	-	0.03	-	0.01
SrO	0.12	0.10	0.26	0.24	0.59	4.62	2.72	5.93
BaO	0.05	0.06	0.05	0.02	0.05	-	-	-
Al ₂ O ₃	0.02	0.01	0.01	0.04	0.72	0.01	0.03	0.01
Na ₂ O	0.07	0.01	0.01	0.01	-	0.03	-	0.02
K ₂ O	0.03	-	0.01	0.02	0.10	0.01	0.15	-
F	3.83	3.97	3.66	3.66	1.49	2.56	3.04	3.71
H ₂ O	0.09	0.06	0.05	0.18	1.19	0.50	0.33	0.00
O=2F	-1.61	-1.67	-1.54	-1.54	-0.63	-1.08	-1.28	-1.56
Total	100.35	100.75	100.76	100.10	100.76	99.06	99.56	100.84
Cations per formula unit: P + Si = 3								
P	2.999	3.000	2.999	2.998	2.928	3.000	2.994	3.000
Si	0.001		0.001	0.002	0.072	0.000	0.006	
Ca	4.769	4.886	4.854	4.840	4.868	4.995	4.761	4.599
Fe	0.018	0.019	0.018	0.022	0.016	0.017	0.025	
Mn	0.197	0.159	0.177	0.119	0.034	0.036	0.071	0.040
Mg	0.001	0.000	0.001	0.001	0.002		0.002	0.001
Ti	0.004	0.001		0.001		0.002		0.001
Sr	0.006	0.005	0.013	0.012	0.029	0.242	0.136	0.294
Ba	0.002	0.002	0.002	0.001	0.002			
Al	0.002	0.001	0.001	0.004	0.072	0.001	0.003	0.001
Na	0.011	0.002	0.001	0.002		0.005		0.003
K	0.003	0.000	0.001	0.002	0.012	0.001	0.019	
F	1.030	1.071	0.985	0.984	0.397	0.730	0.828	1.005
OH			0.048	0.179	0.603	0.270	0.172	

FeO as the total Fe; -: below the detection limits.

Table 4

The chemical composition of phosphate minerals in the Nanping No. 31 pegmatite.

	Laz-1	Laz-2	Laz-3	Plm	Bts	Bts	Kul	Wag-1	Wag-2	Goy
n	2	22	5	Rep	Rep	Rep	3	10	8	15
P ₂ O ₅	42.89	44.38	44.54	43.13	44.46	44.65	34.41	39.24	41.16	30.92
SiO ₂	0.02	0.14	0.05	0.36	0.45	0.35	1.08	0.09	0.04	0.22
CaO	0.02	0.03	0.02	0.66	8.65	4.79	1.75	0.16	0.13	1.66
FeO	11.09	4.06	3.97	0.23	0.77	0.57	9.48	11.24	5.61	0.08
MnO	0.07	0.04	0.02	0.16	0.14	0.19	5.72	4.20	0.79	0.04
MgO	6.98	10.94	11.05	0.02	0.17	0.21	0.85	37.74	45.29	0.02
TiO ₂	0.11	0.06	0.04	0.03	0.07	0.14	0.85	0.05	0.04	0.11
SrO	-	0.01	-	13.01	0.45	6.43	0.12	0.01	0.01	16.92
BaO	0.02	0.08	0.11	0.09		0.29	24.54	0.02	0.04	1.44
Al ₂ O ₃	31.58	33.35	33.39	31.00	32.04	31.27	16.26	0.05	0.04	33.50
Na ₂ O	-	0.01	-	0.02	0.03	0.06	0.11	0.01	-	0.04
K ₂ O	-	0.05	0.01	0.00	0.00	0.00	0.22	0.01	0.01	0.04
F	-	-	-	0.25	1.38	1.73	0.01	11.08	7.77	0.85
H ₂ O	6.05	6.28	6.28	6.01	5.61	5.44	5.02	-0.29	1.72	14.93
Li ₂ O				4.57	4.72	4.71				
O=2F				-0.10	-0.58	-0.73	0.00	-4.65	-3.26	-0.36
Total	98.83	99.44	99.48	99.43	98.37	100.09	100.42	98.95	99.39	100.41
P	1.999	1.992	1.997	3.961	3.953	3.964	2.903	0.997	0.999	1.983
Si	0.001	0.008	0.003	0.039	0.047	0.036	0.097	0.003	0.001	0.017
Ca	0.001	0.002	0.001	0.076	0.961	0.531	0.167	0.005	0.004	0.133
Fe	0.511	0.180	0.176	0.021	0.068	0.050	0.821	0.282	0.135	0.005
Mn	0.003	0.002	0.001	0.014	0.012	0.016	0.504	0.107	0.019	0.002
Mg	0.573	0.865	0.873	0.004	0.027	0.033	0.129	1.690	1.936	0.002
Ti	0.004	0.002	0.001	0.002	0.006	0.011	0.063	0.001	0.001	0.007
Sr		0.000		0.818	0.028	0.391	0.006	0.000	0.000	0.744
Ba	0.000	0.002	0.002	0.004		0.012	0.958	0.000	0.000	0.043
Al	2.049	2.084	2.085	3.963	3.966	3.864	1.916	0.002	0.001	2.992
Na		0.001		0.005	0.006	0.011	0.021	0.001		0.005
K		0.004	0.001	0.000	0.000	0.000	0.030	0.000	0.000	0.005
Li				1.995	1.994	1.988				
F				0.084	0.460	0.575	0.003	1.053	0.704	0.204
OH	2.000	2.000	2.000	3.916	3.540	3.425	2.997	0.000	0.296	6.796

FeO as the total Fe; -: below the detection limits. Structural formulas were calculated on the basis of 2, 4, 3, 1, 2 and 2 (P+Si) atoms for lazulite, palermoite-bertossaite, kulanite, wagnerite and goyazite, respectively. Abbr.: Laz - lazulite, Plm - palermoite, Bts - bertossaite, Kul - kulanite, Wag - wagnerite, Goy - goyazite.

