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1	Into the high to ultrahigh temperature melting of Earth's crust: Investigation of
2	melt and fluid inclusions within Mg–rich metapelitic granulites from the Mather
3	Peninsula, East Antarctica
4	
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31 ABSTRACT

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Precise constraints on the compositions of melts generated by anatexis under ultrahigh 33 34 temperature (UHT) conditions are critical for understanding processes of partial melting and differentiation of the Earth's crust. Here we reveal geochemical and physical signatures of anatectic 35 melts preserved as nanogranitoids (i.e. crystalized melt inclusions) within sapphirine-bearing UHT 36 metapelitic granulites from the Mather Peninsula, East Antarctica. Their coexistence with high-Al 37 orthopyroxene as inclusions in garnets strongly suggests that the investigated melts were at least 38 partially UHT in origin. The nanogranitoids are enriched in SiO_2 (69.9–75.6 wt.%), strongly 39 peraluminous (ASI values = 1.2-1.6) and potassic to ultrapotassic (Na₂O + K₂O = 7.1-9.5 wt.%, 40 K/Na = 2.2-9.3). When compared to the granulitic restite, the melts are enriched in Li, Cs, Rb, Ta, 41 Sm, Nd, Zr, U and Pb, and depleted in Ce, Th, Ba, Sr and Nb. Their geochemical characteristics are 42 consistent with biotite-dehydration melting in the absence of plagioclase. Our calculation results 43 indicate that these hot crustal melts have low densities of 2.47 ± 0.07 g/cm³, low viscosities of 44 $10^{4.9\pm1.2}$ Pa·s and high heat production values of ~2.8 μ W/m³. Therefore, such melts are mobile and 45 susceptible to be extracted from the source, and consequently their flow and removal from the deep 46 crust may greatly affect the chemical and thermal structure of the continental crust. 47

Secondary C–O–H fluid inclusions within garnet and orthopyroxene have also been detected. 48 These inclusions contain magnesite, pyrophyllite, corundum, with or without residual CO₂. The 49 minerals within the fluid inclusions are interpreted as stepdaughter minerals, which were produced 50 by the reaction of the fluid with their host. The metamorphic timing of the investigated rocks is still 51 a matter of debate. Zircon U-Pb dating results obtained in this study suggest that the metapelitic 52 granulites may have undergone two separated thermal events at ~ 1000 and ~ 530 Ma, respectively. 53 The presence of fluid inclusions indicates that fluid infiltration and Pan–African reworking may 54 have played an important role in obscuring chronological information of the early thermal scenario 55 in poly-metamorphic terranes. 56

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58 Key Words:

59 *UHT* metamorphism; Melt inclusions; C–O–H fluid; Mather Peninsula; East Antarctica

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61 INTRODUCTION

Ultrahigh temperature (UHT) metamorphism is the most thermally extreme type of crustal 62 metamorphism, with peak temperatures exceeding 900 °C at moderate pressures (7-13 kbar) 63 (Harley, 1998a, 2008), with dT/dP values of 66–200 °C/kbar (Harley, 2021). UHT metamorphism is 64 now recognized as a relatively common rather than anomalous process of the Earth's crust due to 1) 65 its wide age distribution ranging from Archean to Quaternary (e.g. Brown, 2007; Harley, 2021; 66 Pownall et al., 2014; Zhang et al., 2022); and 2) its extensive exposures around the world (>70 67 localities; Harley, 2021; Lei & Xu, 2018) related to various geodynamic processes. For example, 68 UHT metamorphism may occur in tectonic backgrounds associated with 1) lithospheric extension 69 and underplating of mantle-derived magmas as a result of slab rollback (e.g. Pownall et al., 2014), 70 mantle plume activity (e.g. Liu et al., 2020; Tong et al., 2014a); 2) oceanic ridge subduction (e.g. 71 Santosh et al., 2012), and inversion and thickening of hot back-arc (e.g. Brown, 2007); 3) long-72 lived orogenic self-heating by radioactive elements (e.g. Clark et al., 2011; Zhang et al., 2022); or 4) 73 a combined contribution from several driving forces (Huang et al., 2019, 2021). 74

High temperature (HT) to UHT metamorphism of crustal rocks inevitably produces anatectic 75 melts at depth, and their loss yields a refractory lower crust, whereas ascent and emplacement of 76 granitic melts at shallower levels create a more silicic upper crust (e.g. Brown, 2007, 2013; Sawyer 77 et al., 2011). Furthermore, it may result in decrease of the rock strength and variations in bulk 78 compositions of the granulitic rocks (e.g. Cavalcante et al., 2016; Diener & Fagereng, 2014; 79 Rosenberg & Handy, 2005). However, interpretations of melting mechanisms of crustal rocks 80 remain a long-standing challenge, which have been traditionally tracked with the aid of 1) 81 geochemistry of granites and leucosomes (e.g. Gao et al., 2017; Wolf et al., 2019; Zeng et al., 2005), 82 2) phase equilibria modeling at specific P-T conditions (e.g. Huang et al., 2021; Wei, 2016; White 83 et al., 2004, 2014), and 3) melting experiments on natural rocks under variable $P-T-X_{H2O}$ 84 conditions (refer to Gao et al., 2016 and references therein). Furthermore, a recent progress is to 85 conduct in-situ observations and analyses on nanogranitoid inclusions (partially to totally 86 crystallized melt inclusions preserved in peritectic phases; named after Cesare et al., 2015) and their 87 glassy counterparts (e.g. Acosta-Vigil et al., 2010, 2012; Bartoli et al., 2013a, b, 2014, 2016, 2019; 88 Bartoli & Carvalho, 2021; Borghini et al., 2018, 2023; Carvalho et al., 2019, 2023a; Cesare et al., 89 2009, 2015; Ferrero et al., 2012, 2016, 2018, 2021a, b; Ferrero & Angel, 2018; Ferri et al., 2020; 90

Gao et al., 2014; Gianola et al., 2021; Higashino & Kawakami, 2022; Tacchetto et al., 2021). Thus 91 far, melt inclusions have been only recognized in a few UHT metapelitic terranes, including 1) the 92 Ivrea Zone (NW Italy) (Carvalho et al., 2019), 2) the Kerala Khondalite Belt (India) (Cesare et al., 93 2009; Ferrero et al., 2012), 3) the Gruf Complex (European Central Alps) (Gianola et al., 2021), 4) 94 the Lützow-Holm Complex (East Antarctica) (Carvalho et al., 2023a; Hiroi et al., 2019; Kawakami 95 & Motoyoshi, 2004; Suzuki & Kawakami, 2019), 5) the Sør Rondane Mountains (East Antarctica) 96 (Higashino & Kawakami, 2022), and 6) the Chinese Altai orogen (NW China) (Liu et al., 2020). It 97 is worth noting that the occurrence of nanogranitoids in UHT rocks does not imply automatically 98 that they reflect the UHT crustal melts because melts can be trapped during the HT prograde 99 metamorphism (refer to Gianola et al., 2021). A better understanding of potential effects of UHT 100 metamorphism on the differentiation and dynamic evolution of the crust demands that the hottest 101 crustal melts are chemically characterized, in particular for their major and trace elements. 102

The Mather metapelitic granulites from the Rauer Islands (East Antarctica) are a world-103 renowned poly-metamorphic UHT lithology, with peak P-T conditions of 8.5-12 kbar and 950-104 1050 °C (Chen et al., 2023; Clark et al., 2019; Harley, 1998b, 2016; Harley et al., 2009; Harley & 105 Fitzsimons, 1991; Kelsey et al., 2007; Tong & Wilson, 2006). In this contribution, we conduct an 106 integrated investigation of microstructural features, major and trace element contents of 107 nanogranitoid inclusions (NIs) preserved within garnet porphyroblasts. Data collected in this work 108 are fundamental to the understanding of melting regimes of crustal rocks under UHT conditions, 109 and their contribution to crustal differentiation and orogenic evolution. Secondary fluid inclusions 110 (FIs) in peritectic phases are also characterized to reveal the nature of post-peak infiltrated fluid, 111 and to evaluate its potential role in resetting the U–Pb clock of zircons. 112

113

114 GEOLOGICAL SETTING

The Antarctic continent is separated by the Transantarctic Mountains into two distinct geological units, namely East Antarctica and West Antarctica. East Antarctica is one of the largest cratons around the world, documenting more than 3.5 billion years of evolutional history (Harley et al., 2013). The Prydz Belt was recognized as a tectonic mobile belt in East Antarctica that has been extensively affected by the Pan–African (~530 Ma) thermal event (e.g. Grew et al., 2012; Harley et al., 1998; Harley & Kelly, 2007; Hensen & Zhou, 1997; Kelsey et al., 2003, 2007; Liu et al., 2009,

2021; Tong et al., 2014b, 2019; Wang et al., 2007, 2008, 2022). The tectonic nature of the Prydz 121 Belt remains debated and there are two main schools of thought; one considers that it represents a 122 Pan-African suture zone associated with the final assembly of East Gondwana (e.g. Fitzsimons, 123 2003; Hensen & Zhou, 1997; Kelsey et al., 2007; Liu et al., 2021; Wang et al., 2022; Zhao et al., 124 1995), and the other views it as a Pan-African intraplate orogen related to intracontinental 125 reworking (e.g. Phillips et al., 2007; Ren et al., 2016; Tong et al., 2014b, 2019; Wilson et al., 2007). 126 The Rauer Islands are located on the eastern margin of the Prydz Belt, with the Brattstrand 127 Bluff and Larsemann Hills to the southwest, and the Vestfold Hills to the northeast. The Rauer 128 Islands were thought to be texturally correlated with the Rengali–Eastern Chats Province, forming 129 parts of the Indo-Antarctica supercontinent (e.g. Sawant et al., 2017). Archean and Proterozoic 130 crustal components have been recognized from the Rauer Islands (Kinny et al., 1993). The 131 Proterozoic domain comprises metamorphosed 1420-1000 Ma mafic-intermediate-felsic intrusive 132 rocks (Kinny et al., 1993; Liu et al., 2021). The early Neoproterozoic (i.e. Grenville-aged; 1000-133 900 Ma) and late Neoproterozoic/Cambrian (i.e. Pan-African-aged; 590-500 Ma) ages have been 134 identified from metamorphic rocks in this region (e.g. Kelsey et al., 2007; Kinny et al., 1993; Liu et 135 al., 2021). The Archean domain predominantly consists of tonalitic to granitic orthogneisses with 136 three age clusters of 3470-3270, 2840-2800 and ~2550 Ma (Harley et al., 1998; Hokada et al., 137 2016; Kinny et al., 1993; Liu et al., 2021). This domain also contains Fe- or Mg-rich layered mafic 138 complexes that were subjected to a granulite facies metamorphism in the Cambrian (Chen et al., 139 2023; Harley et al., 1998; Harley & Kelly, 2007). Although Pan–African ages have been widely 140 recognized in the Archean domain (e.g. Clark et al., 2019; Hokada et al., 2016; Kelsey et al., 2003, 141 2007; Liu et al., 2021; Wang et al., 2007), it is still unclear whether it was also involved in the 142 Grenville-aged tectono-thermal event. 143

The Rauer Islands consist of two paragneiss successions, namely the Filla and the Mather Paragneisses (Fig. 1a; Harley & Fitzsimons, 1991). These two paragneiss successions are mainly distributed on the Proterozoic and Archean crustal domains, respectively. Deposition ages of the Filla Paragneiss have been constrained to be Mesoproterozoic (Liu et al., 2021; Sims et al., 1994). This unit was considered to have experienced a prominent metamorphic event at 1030–970 Ma and a high–grade reworking at 575–510 Ma (e.g. Kelsey et al, 2007; Kinny et al., 1993; Liu et al., 2021). The Mather Paragneiss typically comprises magnesian and aluminous garnet–orthopyroxene–

sillimanite-bearing metapelite, orthopyroxene-sillimanite metaquarzite, magnesian garnet-151 orthopyroxene metapelite and garnet-bearing mafic granulite (Fig. 1b; Harley et al., 1995; Harley, 152 1998b). This suite usually occurs as thin and laterally discontinuous horizons from the Mather 153 Peninsula to the Short Point in the eastern Rauer Islands (Hokada et al., 2016). Mather Paragneiss 154 preserves strong evidence for UHT metamorphism, with exposures of sapphirine-bearing 155 metapelitic granulites on the Mather Peninsula (e.g. Harley, 1998b; Tong & Wilson, 2006) and the 156 Torckler Island (Harley et al., 2009), with peak P-T conditions of 8.5-12 kbar and 1000-1050 °C 157 (e.g. Harley, 1998b, 2016; Tong & Wilson, 2006). The sapphirine-bearing UHT gneisses preserve 158 inherited zircon ages of 2800–2400 Ma (e.g. Hokada et al., 2016; Wang et al., 2007), and are hosted 159 by \sim 3267 Ma tonalitic orthogneisses (Hokada et al., 2016). However, timing of the UHT 160 metamorphism is still a matter of debate. Some researchers argued that the UHT metamorphism 161 occurred at ~ 1000 Ma and the subsequent decompression and final cooling happened at ~ 530 Ma 162 (e.g. Tong & Wilson, 2006; Wang et al., 2007) while others considered that the UHT 163 metamorphism occurred in the Cambrian (e.g. Clark et al., 2019; Kelsey et al., 2003, 2007) or prior 164 to 590–580 Ma (Hokada et al., 2016; Harley et al., 2009). 165

166

167 **PETROGRAPHY**

The granulite sample RG16-80 (68°50'50" S and 77°54'43" E) was collected from the Mather 168 Peninsula in the Rauer Islands, East Antarctica (Fig. 1b). Compositions and mineral assemblages 169 vary considerably at outcrop scale, ranging from quartz-bearing to SiO₂-undersaturated rocks (refer 170 to Harley, 1998b for more details). The studied rocks occur as discrete rafts or boudins interlayering 171 with Grt–Opx–Sil–Spr granulites, Grt–rich quartzites, Grt–Sil metapelites, two–pyroxene granulites, 172 Opx–Sil–(Per)–bearing and Spr–Opx–Kfs–bearing leucogneisses (Fig. 2). Mineral abbreviations are 173 from Warr. (2021). The studied Mg-rich metapelites constitute 5-8% of the outcrop, and are 174 roughly parallel to foliation S6 (Tong & Wilson, 2006), corresponding to the composite high-strain 175 fabric described by Dirks & Wilson (1995). The samples are coarse-grained, with an assemblage of 176 orthopyroxene (35–45%), biotite (25–30%), garnet (~15%), cordierite (7–10%), minor sapphirine, 177 plagioclase, K-feldspar, quartz (<5%), and accessory spinel, ilmenite, apatite, zircon and monazite 178 (Fig. 3). 179

Garnet porphyroblasts are subhedral to anhedral, with sizes varying from 0.1 to 30 mm. They

usually preserve corroded biotite and quartz inclusions in cores. Most orthopyroxene grains are 181 present as porphyroblasts (up to 1.2 cm in diameter), and in rare cases, they may contain platy or 182 rounded biotite inclusions (Figs. 3a, b). The inclusions within garnet cores and orthopyroxene, such 183 as biotite and quartz, are considered as pre-peak (M0) minerals. The cores of orthopyroxene 184 porphyroblasts are considered to have been formed at M0 stage because they contain relatively low 185 Al_2O_3 contents (<6.5 wt.%) compared with the rim (>8.0 wt.%; Opx₁) (see details on the section of 186 mineral chemistry), and high Al₂O₃ contents in orthopyroxene usually suggest that they were 187 formed under UHT conditions (e.g. Harley, 1998a). 188

Sometimes, orthopyroxene with high Al₂O₃ contents (up to 8.85–9.04 wt.%) may occur as inclusions within garnet mantle (Fig. 3a), and both minerals are considered to form at, or close to, peak conditions. Collectively, garnet mantles, orthopyroxene rims, high–Al orthopyroxene inclusions within garnets and biotite porphyroblasts (Fig. 3b) are interpreted to be formed at the metamorphic peak (M1) stage. K–feldspar occurs in very local domains, and is usually 0.5–4.0 mm in size (Fig. 3c). However, it is difficult to determine whether K–feldspar was formed at M0 or M1 stage based on petrographic observations.

The presence of biotite and quartz inclusions within garnets, coupled with the formation of garnet and orthopyroxene porphyroblasts, implies the progress of the following Bt-dehydration melting reaction:

199 Bt $(+ Kfs) + Qz \rightarrow Grt + Opx + melt.$

Garnet porphyroblasts are commonly replaced at their margins by the intergrowth of orthopyroxene (Opx_2) + cordierite ± biotite ± plagioclase ± spinel (Figs. 3d, e), which are assigned to the post-peak (M2) assemblage. Formation of the intergrowth is considered to result from the following decompression-related reaction:

204

Grt \rightarrow Opx₂ + Crd \pm Spl (e.g. Harley, 1998b; Santosh et al., 2012).

Intergrowths of sapphirine + cordierite are found in the matrix (Fig. 3f). Occasionally,
 vermicular low-Al orthopyroxene and plagioclase are preserved in the rim of high-Al
 orthopyroxene porphyroblasts (Fig. 3g). These microstructures are also interpreted to form at the
 post-peak (M2) stage.

209 Some medium– to fine–grained sapphirine strips are associated with oriented biotite, which 210 sometimes, may surround orthopyroxene + cordierite intergrowths (Fig. 3h). In strongly deformed domains, elongated garnet, cordierite, biotite, plagioclase, rare orthopyroxene and spinel grains
show a similar orientation (Fig. 3i), which may have been formed at the final cooling (M3) stage.
The appearance of interstitial plagioclase films suggests the presence of former melts (Fig. 3i) (e.g.

214 Holness et al., 2011).

The presence of retrograde biotite and plagioclase is in agreement with progress of the following reaction:

217 Grt (Opx) + melt \rightarrow Bt + Pl \pm Qz.

In summary, the Spr-bearing UHT granulites (sample RG16-80) from the Mather Peninsula 218 preserve four-stage metamorphic assemblages: 1) M0 stage is represented by the Bt-Qz-bearing 219 inclusions and the cores of garnet and orthopyroxene porphyroblasts; 2) M1 stage is marked by the 220 growth of garnet–orthopyroxene₁–biotite in the presence of melt, with or without the presence of 221 K-feldspar; 3) M2 stage is featured by the development of orthopyroxene₂ + cordierite \pm biotitet \pm 222 plagioclase \pm spinel and sapphirine + cordierite intergrowths surrounding the peak garnet or 223 orthopyroxene; and 4) M3 stage is characterized by the growth of cordierite-biotite-plagioclase-224 bearing assemblages during the crystallization of melts. 225

226

227 ANALYTICAL METHODS

228 Back–scattered electron imaging, X–ray mapping and EPMA analysis

Analyses of major elements in minerals and back-scattered electron (BSE) imaging were 229 conducted using JEOL JXA-8230 electron probe microanalyzer (EPMA) at the Department of 230 Geology, Northwest University, Xi'an, China. Additional X-ray mapping of NIs was done using 231 JEOL JXA-8230 EPMA at the Wuhan Sample Solution Analytical Technology Co., Ltd., Wuhan, 232 China. Analytical conditions were: an accelerating voltage of 15 kV, and a beam current of 2×10^{-8} 233 A, a beam diameter of 2 µm. Counting times were 20 s on peaks, with 10 s on backgrounds. Data 234 correction was processed by using a ZAF method. Fe³⁺ was calculated by charge balance. The 235 analysis results are shown in Table 1. X-ray mapping was performed using an accelerating voltage 236 of 20 kV, a beam current of 5×10^{-8} A, a beam diameter of 0.3 µm and a dwell time of 50 ms. 237

238

239 Trace elements in garnet

240 Trace element analyses of garnets were conducted with LA-ICP-MS at the Wuhan Sample Solution

Analytical Technology Co., Ltd., Wuhan, China. Detailed operating conditions of the laser ablation 241 system, the ICP–MS instrument and the data reduction were as same as in Zong et al. (2017). Laser 242 sampling was performed using a GeolasPro laser ablation system that consists of a COMPexPro 243 102 ArF excimer laser and a MicroLas optical system. An Agilent 7700e ICP-MS instrument was 244 used to acquire ion-signal intensities. A "wire" signal smoothing device was included in this laser 245 ablation system (Hu et al., 2015). Spot size and frequency of the laser were set to be 44 μ m and 5 246 Hz, respectively. Trace element compositions of garnets were calibrated based on various reference 247 materials (BHVO-2G, BCR-2G and BIR-1G) without using an internal standard (Liu et al., 2008). 248 An Excel-based software ICPMSDataCal was used to perform quantitative calibration for trace 249 250 element analyses (Liu et al., 2008).

251

252 **Raman measurements**

Micro-Raman measurements were performed on representative thick sections at the School of Earth 253 Science and Resources, Chang'an University, using a HORIBA LabRam HR Raman 254 microspectrometer. A frequency-doubled Nd:YAG laser with 532 nm excitation wavelength was 255 used. A petrographic microscope Olympus ($50 \times$ objective) was used to focus the laser on the 256 selected inclusions. Raman spectra in the ranges 100–4000 cm⁻¹ were collected with an integration 257 time of 10 s, and three accumulations. Data were processed using LabSpec v5.58.25 and the phases 258 inside the inclusions were identified according to the data from the literature (Frezzotti et al., 2012; 259 Ferrero et al., 2016, 2021b; Carvalho et al., 2020; Gianola et al., 2021). 260

261

262 **Re-homogenization** of nanogranitoids

Re-homogenization experiments of NIs were conducted following the procedure described in 263 Bartoli et al. (2013a), using a piston cylinder apparatus at the CAS Key Laboratory of Crust–Mantle 264 Materials and Environments, University of Science and Technology of China and the QUICKpress 265 non-endloaded piston cylinder available at the Laboratory of Experimental Petrology and 266 Geochemistry, Dipartimento di Geoscienze, Università di Padova. During each run, separated 267 garnet grains were charged into Au capsules with external diameters of 3-5 mm and length of 5-10 268 mm. They were alternated by powdered silicon dioxide. The experiments were performed under the 269 following conditions: 1000 °C and 6 hours, 950 °C and 4 hours, 950 °C and 2 hours, and 950 °C and 270

1 hour. A fixed pressure of 12 kbar was adopted during the experiments.

272

273 Major elements in nanogranitoids

Major element compositions of the re-homogenized NIs were determined using a JEOL JXA 8530F 274 plus EPMA at the CAS Key Laboratory of Crust-Mantle Materials and Environments, University of 275 Science and Technology of China, Hefei, China. During EMPA analysis of hydrous granitic glasses, 276 migration and loss of alkali from the inclusions may happen in the case that small beam sizes and 277 high beam currents are adopted (e.g. Cesare et al., 2015). To minimize the alkali loss, K₂O and 278 Na₂O were determined at first. An anhydrous glass B (Morgan & London., 2005) was analyzed to 279 obtain correction coefficients, which were used to correct the data of the unknown glasses. 280 Analytical conditions were set as: a 15 kV accelerating voltage, a 5 nA beam current, variable beam 281 diameters of $2-5 \mu m$ and counting times of 10 s on peak and 5 s on backgrounds. 282

283

284 Trace elements in nanogranitoids

Trace elements in NIs were analyzed at the State Key Laboratory for Mineral Deposit Research, 285 Nanjing University, using a Coherent GeolasHD system equipped with a 193 nm excimer laser 286 coupled to a NexION 350 ICP mass spectrometer (Pan et al., 2019). The samples were loaded and 287 analyzed in an in-house rhomb-shaped plexiglass cell of relatively small internal volume to enable 288 fast washout of aerosol. The He-based aerosol was admixed with nebulizer gas via a T-junction 289 prior to injection into the plasma. The ICP-MS was tuned to achieve maximum sensitivity and low 290 oxide production rate by lowering the ThO/Th ratios to <0.5 % before analyses. The analyzed 291 inclusions are 2-25 µm beneath the surface to ensure they were representative of the original melts. 292 The NIs close to the surface $(2-10 \ \mu m \text{ in depth})$ were preferentially selected for analyses. In the rare 293 294 case of deep-seated inclusions (15–25 µm in depth), the host garnet next to the inclusion was also analyzed and the ablation signal from the same depth interval as the inclusion was integrated, which 295 will minimize the down-hole fractionation of elements during ablation (Zajacz and Halter, 2007). 296 Repetition rate of the laser was increased to 10 Hz, on-sample energy densities of 5 J/cm² and spot 297 size of 24 or 32 µm based on the size of NIs. Generally, measurement of each melt inclusion 298 299 consists of 20-30 s instrumental background followed by 50-80 s ablations depending on the depth of melt inclusion. The glass NIST SRM 610 was employed as external standard. For the analyses of 300

the NIST SRM 610, the spot size and energy density were set as 32 μ m and 5 J/cm², respectively. 301 The acquired time-resolved signals were processed using the software SILLS (Guillong et al., 302 2008). Due to the small size of NIs, part of the host garnet has also been ablated along with target 303 inclusions. Therefore, deconvolution of the resulting mixed garnet-glass signal was applied. 304 following the procedures of Halter et al. (2002). The Al₂O₃ of the re-homogenized inclusions 305 derived from EPMA was adopted as reference standards for inclusions and garnet hosts, and MgO 306 was used as the matrix-only tracer. Uncertainties and calculated limits of detection increase with 307 308 the concentrations of elements in the host, thus, some elements which are strongly compatible in the host garnets (e.g. Y, Gd, V, Zn, Sc and heavy rare earth elements-HREEs) are normally not 309 detectable in the inclusions. Some inclusions show anomalously high concentrations of Zr, Th, U, 310 Nb and Sr, suggesting the occurrence of accidental trapped phases (e.g. zircon, monazite, rutile and 311 apatite) (Bartoli et al., 2019). For these inclusions with extremely high contents of Zr, Th, U, Nb 312 and Sr, only Rb, Cs, Ba and Li are reported in Table 4 and discussed in the text. 313

314

315 U–Pb dating of zircon

Conventional magnetic and heavy liquid techniques followed by hand-picking under a binocular 316 microscope were used for separation of zircons from the Mather UHT metapelitic granulites. 317 Morphology and internal structure of the zircons were documented with transmitted, reflected light 318 microphotographs and cathodoluminescene (CL) images. Zircon U-Pb dating was carried out using 319 LA-ICP-MS at the Wuhan Sample Solution Analytical Technology Co., Ltd., Wuhan, China. 320 Detailed operating conditions for the laser ablation system and the ICP-MS instrument and data 321 reduction were described in Zong et al. (2017). Laser sampling was performed using a GeolasPro 322 laser ablation system that consists of a COMPexPro 102 ArF excimer laser (wavelength of 193 nm 323 and maximum energy of 200 mJ) and a MicroLas optical system. Each analysis incorporated a 324 background acquisition of approximately 20–30 s followed by 50 s of data acquisition from the 325 sample. Spot size and frequency of the laser for analyses were set to 24 mm and 10 Hz, respectively. 326 Zircon 91500 and glass NIST610 were used respectively as external standards for U-Pb dating and 327 trace element calibrations. In order to monitor the external uncertainties of zircon dating, a zircon 328 standard GJ-1 was alternately analyzed together with the other unknown zircons. Fourteen 329 measurements of the zircon GJ-1 yielded a weighted mean ${}^{206}Pb/{}^{238}U$ age of 602.1 ± 4.4 Ma 330

(MSWD = 0.22), which is roughly in agreement with the recommended 206 Pb/ 238 U age value of 600.4 ± 0.6 Ma (Jackson et al., 2004). An Excel–based software ICPMSDataCal was used to perform off–line selection and integration of background and analyzed signals, time–drift correction and quantitative calibration for trace element analyses and U–Pb dating (Liu et al., 2010). Programs SQUID 1.0 and ISOPLOT (V.3.0; Ludwig, 1999) were used for data processing.

336

337 **RESULTS**

338 Mineral Chemistry

Garnet porphyroblasts are chemically homogeneous from core to mantle (Alm₃₄₋₃₈Prp₅₉₋₆₂Grs₃₋₄Sps₀₋₁). Small garnet crystals and garnet porphyroblast rims have slightly lower pyrope contents, with compositions of Alm₄₀₋₄₆Prp₅₅₋₅₆Grs₄₋₃Sps₀₋₁ (Figs. 4a, b). The garnet relicts from the strongly deformed domain display the lowest magnesium contents (Alm₄₈₋₅₀Prp₄₆₋₄₄Grs₄₋₅Sps₀₋₁). NI– bearing and NI–absent domains display similar REE patterns in the chondrite–normalized diagram, with enrichments of HREEs and slightly negative Eu anomalies (Fig. 4c).

Orthopyroxene porphyroblasts preserve evident Al–zoning patterns, with y_{Al} (= Al^{IV} = Al/2) 345 values increasing from 0.14 (Al₂O₃ = \sim 6.49 wt.%) in the core to 0.19 (Al₂O₃ = \sim 8.79 wt.%) in the 346 rim (Fig. 4d). The highest Al_2O_3 content of orthopyroxene detected in the sample is ~9.16 wt.%, 347 with corresponding y_{A1} value of 0.19. However, orthopyroxene porphyroblasts show relatively 348 restricted $X_{Mg} (X_{Mg} = Mg/(Mg + Fe^{2+}))$ variations (0.77–0.79). Orthopyroxene inclusions in garnet 349 have Al₂O₃ contents of 8.85–9.04 wt.% ($y_{Al} = \sim 0.19$) and X_{Mg} values of 0.78–0.80. Orthopyroxene 350 from the orthopyroxene-cordierite symplectites have y_{Al} of 0.14–0.16 and X_{Mg} of 0.75–0.80. 351 Vermicular low-Al orthopyroxene in the rims of some orthopyroxene porphyroblasts displays Al₂O₃ 352 contents of ~6.88 wt.% ($y_{Al} = ~0.15$) and X_{Mg} values of ~0.77. Orthopyroxene grains associated 353 with the retrograde biotite display Al₂O₃ contents of 7.67–7.69 ($y_{Al} = 0.16-0.17$) and X_{Mg} of 0.74– 354 0.76. 355

Biotite porphyroblasts show relatively homogeneous compositions, with TiO₂ contents of 4.29– 4.61 wt.% and X_{Mg} of 0.79–0.81. The low TiO₂ contents for peak biotite can be partially ascribed to crystal–chemical constraints, which hamper the temperature control on Ti content of biotite (Cesare et al., 2008). Biotite inclusions within garnet have relatively higher X_{Mg} values (0.84–0.87) than those within orthopyroxene (~0.79). However, they show similar TiO₂ content variations ranging Sapphirine associated with retrograde biotite has slightly higher X_{Mg} values (0.84–0.86) than that from sapphirine–cordierite intergrowths (0.80–0.82), both of which show indistinguishable Fe^{3+}/Fe^{T} ratios varying from 0.71 to 0.81. Cordierite grains from different positions show restricted X_{Mg} variations from 0.90 to 0.92. Spinel has X_{Mg} of 0.43–0.48 and ZnO contents of 0.64–1.72 wt.%. Plagioclase from the orthopyroxene–cordierite intergrowths is rich in anorthite, with a typical X_{An} (= Ca/(Ca + Na + K)) of ~0.81. However, plagioclase associated with the retrograde biotite has lower X_{An} values of 0.51–0.58.

371

372 Microstructures of melt and fluid inclusions

Nanogranitoids are randomly distributed within garnet crystals, but absent in the outer rims (Figs. 373 3a, 4a, 5a). They are also absent in the small garnet crystals from the strongly deformed domains 374 (Fig. 3i). These inclusions show spherical or polygonal shape, and are $5-15 \mu m$ in size (Figs. 5b-f, 375 6). They usually display negative crystal shapes, and may contain nano-scale porosities (Fig. 5e), 376 implying the possible presence of fluid which was dissolved in the former melts and subsequently 377 exsolved upon crystallization (Bartoli et al., 2013b). Coupled Raman analyses and compositional 378 maps suggest that they contain daughter phases of quartz, kokchetavite (KAlSi₃O₈), minor 379 kumdykolite (NaAlSi₃O₈) and biotite (Figs. 5g, 6), indicating the former presence of silicate melts. 380 Moreover, an unknown phase with Raman peaks at 292, 429 and 476 cm⁻¹ has also been detected 381 (Fig. 5g). Such phase has been previously recognized within garnet-hosted melt inclusions from the 382 Gruf Complex (Gianola et al., 2021) and the Central Maine Terrane (Ferrero et al., 2021b). This 383 phase was described as phase "430" by Ferrero et al. (2021b). Apatite, rutile, zircon and monazite 384 may be present as accidentally trapped phases, as indicated by the relatively large size of trapped 385 phases compared with the size of the melt inclusions, and also by their low solubility in granitic 386 melts (e.g. Bartoli et al., 2019). Daughter biotite crystals in NIs have very low TiO₂ contents of 387 0.03–0.50, and is characterized by high X_{Mg} values ranging from 0.92 to 0.95, whereas F and Cl 388 contents in biotite are negligible, lower than 0.07 and 0.03 wt.%, respectively. Kokchetavite has a 389

typical composition of $Or_{94}Ab_6$ and kumdykolite displays compositions of $Or_5An_8Ab_{87}$. For detailed mineral compositions, please refer to Table 2.

Fluid inclusions are also found within garnet (Figs. 3a, 7a, b) and orthopyroxene (Fig. 7c). 392 These fluid inclusions display granular or vermicular shapes, with sizes of 5–40 μ m (Fig. 7). They 393 are usually present along healed cracks, suggesting that they are secondary in origin. Some early 394 trapped fluid has reacted with their hosts to produce stepdaughter phases (i.e. their precipitation 395 results from fluid-host interaction; Carvalho et al., 2020) of pyrophyllite, magnesite, corundum and 396 quartz, with or without residual CO_2 (Figs. 7d, e). The measured CO_2 Fermi diad splits range from 397 102.8 to 105.1 cm⁻¹, corresponding to densities of 0.2–1.0 g/cm³ calculated with the densimeter of 398 Wang et al. (2011). Secondary pure CO₂ inclusions may also occur along healed garnet cracks (Fig. 399 7f), which have a density of ~ 0.8 g/cm³. Although not detected in this study, the existence of trace 400 amounts of free H₂O cannot be precluded because H₂O-rich hand may be present as a thin film 401 (<0.1 µm) wetting the walls of the FIs at ambient conditions (e.g. Berkesi et al., 2009; Lamadrid et 402 al., 2014). Other fluid species such as N₂ and CH₄ have not been detected in this study. 403

404

405 Microstructures of re-homogenized nanogranitoids

Experiments at 950 °C with run durations of 2–4 hours yielded some homogenized melt inclusions. 406 However, those at 950 °C with run duration of 1 hour produced the largest proportion of perfectly 407 re-homogenized inclusions (Fig. 8). Some inclusions show clear evidence for overheating, such as 408 the occurrences of 1) irregular boundaries between inclusions and garnet hosts (Fig. 8a); 2) 409 secondary minerals (i.e. orthopyroxene) formed during experiments or upon quenching (Figs. 8a-d); 410 3) new growth of garnet on the inclusion walls (Fig. 8c); and 4) offshoots around melt inclusions 411 formed as a consequence of decrepitations (Figs. 8a, b). The compositions of these inclusions 412 showing interaction with the host have been disregarded. Shrinkage bubbles have also been 413 observed within the overheated inclusions (Figs. 8b, c), suggesting that the fluid may have 414 undergone incomplete dissolution in the melts during the experimental run or, alternatively, it 415 diffused during cooling to form shrinkage bubbles (Lowenstern, 1995; Bartoli et al., 2013a). Some 416 inclusions were not re-homogenized properly during the experiments, as indicated by the 417 coexistence of melts with rounded residual quartz (Fig. 8d). Some melt inclusions still contain 418 accessory minerals such as apatite, rutile, zircon and monazite (Figs. 8a, d-f). This strongly 419

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422 Major element compositions of re-homogenized nanogranitoids

Re-homogenized inclusions are strongly peraluminous (ASI = $[Al_2O_3/(CaO+Na_2O+K_2O)] =$ 1.24–1.61) (Fig. 9) and rhyolitic in composition, with high SiO₂ contents of 69.89–75.61 wt.%, high Na₂O+K₂O contents of 7.08–9.53 wt.% (K/Na = 2.21–9.31) and extremely low CaO (< 0.7 wt.%) and FeO+MgO contents (Table 3). Difference of EMPA totals from 100% suggest that the H₂O contents in the analyzed MIs are 1.49 to 4.24 wt.%, with an average value of 2.79 wt.%.

In the CIPW–normative Qz–Ab–Or diagram (Fig. 10a), the high–SiO₂, potassic to ultrapotassic 428 melts plot far away from haplogranitic minimum melts, and define an elongated trend paralleling to 429 the Qz-Or sideline, which are consistent with the melt compositions found in other UHT 430 occurrences reported by Cesare et al. (2009), Gianola et al. (2021) and Ferrero et al. (2012). Similar 431 compositions have also been obtained by melting experiments of pelitic protoliths at 5-7 kbar and 432 900-950 °C (Droop et al., 2003; Patiño Douce & Johnston, 1991) (Fig. 10a). In the Ab-An-Or 433 diagram, the NIs show variable Or/Ab ratios and low An contents. They plot in the granite field, and 434 are close to the boundary between granite and Qz-monzonite (Fig. 10b). 435

436

437 Trace element compositions of nanogranitoids

In the average upper continental crust (UCC) normalized diagram (Rudnick & Gao, 2014), NIs 438 from the Mather Peninsula are enriched in Rb, Sm, Ta and depleted in Ba, Pb, Th, Ce and Sr (Fig. 439 11). Compared with the granulitic restite (i.e. granulites that have experienced partial melting and 440 substantial melt loss), NIs are enriched in Li (5.3–76.9 ppm), Cs (0.9–35.5 ppm), Rb (126.7–1325.0 441 ppm), Ta (1.4-3.2 ppm), Sm (5.5-27.2 ppm), Zr (84.2-276.5 ppm), Nd (7.2-22.8 ppm), and 442 depleted in Ce (2.4–13.4 ppm), Th (0.4–3.8 ppm), Ba (56.3–565.6 ppm), Sr (5.8–27.6 ppm) and Nb 443 (4.0–15.7 ppm) (Table 4; Fig. 11). The melts have relatively higher U (0.6–9.3 ppm) and Pb 444 contents (3.9-12.8 ppm), and display higher Rb/Sr (7.69-52.08) and Sm/Nd (0.52-1.14) ratios 445 relative to the granulitic restite (Rb/Sr = ~ 8.56 ; Sm/Nd = ~ 0.25). They also contain variable 446 amounts of Sc (9.7-42.0 ppm), V (29.4-146.6 ppm) and Zn (83.3-130.7 ppm), which are discarded 447 in this study because these concentrations might be affected by contamination from the garnet hosts 448 (Bartoli et al., 2019). 449

450 Zircon morphology, U–Pb dates and trace elements

Zircon grains from the Mather UHT metapelitic granulites are usually rounded, and have grain sizes 451 of 80–250 µm. They typically display complex core-mantle-rim structures and occasionally, they 452 may be homogeneous in the CL images (Fig. 12a). The cores are generally 30-120 µm in diameter, 453 and show oscillatory zoning or homogeneous patterns. Most zircon cores are enclosed by dark-grey 454 mantles, which usually show blurred zoning or are unzoned. Sometimes, zircon cores and mantles 455 are enveloped by pale-grey to bright rims (Fig. 12a). A total of 46 analyses of zircon were 456 conducted on 34 grains. Zircon cores yield inherited ${}^{207}Pb/{}^{206}Pb$ dates of 2623–2088 Ma (N = 13; 457 Th/U = 0.28–1.81). Zircon mantles document 206 Pb/ 238 U dates ranging from 1050 to 607 Ma (N = 458 27; Th/U = 0.01–0.09) while the rims record 206 Pb/ 238 U dates varying from 569 to 472 Ma (N = 6; 459 Th/U = 0.11-0.34) (Table 5). 460

Orthopyroxene, biotite, plagioclase, apatite, chlorite, quartz and monazite inclusions were 461 observed within zircons (Fig. 12c). Some trivial inclusions are extensively present along the healed 462 fractures in zircons (Figs. 12b, d). They cannot be analyzed due to their small sizes. However, their 463 appearance resembles FIs reported in literature (Fig. 6d in Li et al., 2018). Thus, these inclusions 464 are suspected to also be FIs. Accordingly, the large variations of U-Pb dates for zircon mantles and 465 rims (Fig. 12e) and Pb loss could result from extensive fluid infiltration and the Pan-African HT-466 *UHT* overprinting. Zircon cores are characterized by the presence of low–Mg biotite ($X_{Mg} = \sim 0.71$) 467 and low-Ca plagioclase ($X_{n} = -0.35$) inclusions (Table 1), which were not found in the matrix, 468 suggesting that these cores are detrital in origin. It is worth noting that an orthopyroxene grain 469 enclosed in a detrital zircon (i.e. it shows a near concordant date of ~ 2381 Ma) shows high Al₂O₃ 470 contents of ~6.79 wt.% (Fig. 12c), indicative of a possible high-grade metamorphism in the 471 Paleoproterozoic. 472

473

474 PHASE EQUILIBRIA MODELING

Phase equilibria modeling for the Mather *UHT* metapelitic granulite sample (RG16–80) was conducted with Perple_X (v. 6.9.1; Connolly, 2005) and the ds622 thermodynamic data set (Holland & Powell, 2011). Pseudosections were calculated in the 11–component MnO–Na₂O–CaO– K_2O –FeO–MgO–Al₂O₃–SiO₂–H₂O–TiO₂–O₂ (MnNCKFMASHTO) system. This system is considered to provide a realistic approximation to the composition of the samples within which melt

and Fe³⁺ can be assessed. The solution model for sapphirine is from Wheller & Powell (2014) and 480 a-x models for other phases are same to those adopted by White et al. (2014). Whole-rock 481 composition of a piece of relatively homogeneous rock ($5 \times 5 \times 3$ cm³), obtained from XRF analysis, 482 was used as the bulk rock composition for the modeling. The formation of fine-grained 483 intergrowths is usually controlled by local effective bulk compositions. In this study, we assume 484 that whole-rock composition obtained from XRF can roughly represent the local effective bulk 485 composition of the symplectite formation. The H₂O contents were adjusted to ensure that the final 486 assemblage is stable just above the solidus (Korhonen et al., 2012). A preliminary $T-X(Fe_2O_3)$ 487 diagram was calculated to evaluate the effect of Fe³⁺contents on phase assemblage (Supplementary 488 file). Based on the predicted peak assemblages across the range of Fe³⁺ contents, a Fe³⁺/Fe^T value of 489 ~ 0.16 is finally adopted. 490

491

492 Peak and post-peak evolution of the Mather *UHT* metapelitic granulites

A pseudosection for the rocks was constructed within a P-T window of 3–15 kbar and 750–1200 °C. 493 Due to lower diffusion rates of Al^{3+} compared to Fe^{2+} and Mg^{2+} , aluminum contents in 494 orthopyroxene are considered reliable to document the peak temperatures of UHT metamorphism. 495 As shown in Figure 13a, the pseudosection is contoured by $(X_{Mg}(Opx) = Mg/(Fe^{2+} + Mg))$ and 496 $y_{Al}(Opx)$ in order to retrieve peak P-T conditions. The interpreted peak assemblage of garnet + 497 orthopyroxene + biotite + melt is stable over a wide P-T range (>7.5 kbar and >860 °C). The 498 measured highest y_{Al} (0.18–0.19) and X_{Mg} (0.775–0.780) in orthopyroxene intersect at 12–13 kbar 499 and 1070–1130 °C, which is interpreted as the peak P-T conditions. 500

The peak minerals usually show breakdown to intergrowths of orthopyroxene₂ + cordierite \pm 501 biotite ± plagioclase ± spinel or sapphirine + cordierite at the post-peak stage. As shown in the 502 enlarged P-T diagram (P = 5.0-9.0 kbar and T = 800-1000 °C) (Fig. 13b), these assemblages 503 occupy a narrow field, constraining the P-T conditions to 6.5–7.0 kbar and 910–950 °C (outlined by 504 purple bold lines). Plagioclase from the orthopyroxene-cordierite-bearing intergrowths has a 505 typical X_{An} of 0.80–0.81, consistent with the prediction by the modeling. The inferred final 506 assemblage is featured by later growth of sapphirine, biotite and plagioclase, which is predicted to 507 be stable at a narrow pressure range of 6.5–6.8 kbar, with temperatures varying from 825 to 900 °C. 508 This field is bounded by the fluid-absent solidus, rutile-in, garnet- and ilmenite-out curves. The 509

measured X_{Mg} in biotite (~0.81) and X_{An} in plagioclase (0.51–0.58) are consistent with the P–T field, further constraining the cooling path through *P–T* conditions of 6.5–7.0 kbar and 830–870 °C (outlined by black bold lines).

513

514 Tracking the pre-peak melting process with melt-reintegration

Phase equilibria modeling suggests that the solidus is located between 800 and 870 °C (at 3–7 kbar) 515 and the rock produces <15 vol.% melt at the peak stage, which is much lower than the expected 516 values for metapelites. In addition, biotite is predicted to be stable at >1100 °C (Fig. 13a). All these 517 observations are in agreement with the strongly residual character of the bulk rock composition 518 $(SiO_2 = 44 \text{ wt.\%})$. Thus, this composition obtained from XRF analysis is only appropriate for 519 modeling the peak and post-peak metamorphic evolutions (White et al., 2004; Liu et al., 2020). 520 Here we explore the potential effects of melt loss on the topology during pre-peak evolution by 521 reintegrating a certain amount of melts to the residual composition (Bartoli, 2017, 2019). The 522 simplified chemical systems with respect to natural occurrences (Bartoli & Carvalho, 2021) and the 523 limitations of solution models (White et al., 2011) make it impossible to precisely predict the 524 primary melt compositions of partially melted rocks. As a consequence, NIs are used as proxy of the 525 pristine melts that was present in the system (e.g. Bartoli, 2019). This single-step melt-526 reintegration may be only suitable to reconstruct the probable bulk rock compositions (BRC) at or 527 near UHT conditions as the rock may have undergone several episodes of melt loss (e.g. Bartoli, 528 2017; White et al., 2014) and the compositions of NIs cannot represent the melt compositions 529 generated at low temperatures. 530

The presence of trapped rutile grains within NIs are confirmed by Raman analyses (Fig. 5g), 531 but not observed in the matrix or predicted by the phase equilibria modeling (Fig. 13). As shown in 532 Figure 14a, rutile is expected to appear at >950 °C if >20 mol.% melts are reintegrated back into the 533 rock. This is consistent with their presence as trapped phase within the NIs (Fig. 5g), and is strongly 534 supportive of the entrapment of rutile-bearing melt inclusions at UHT conditions. With 25 mol.% 535 melts reintegrated back into the BRC, the modeling predicts that most melts (~40 mol.%) were 536 produced in the temperature range of 850-950 °C. The melt reintegration also results in a reduction 537 of the stability field of biotite, which is predicted to disappear at >950 °C (Fig. 14a). The melt 538 composition calculated at 12.5 kbar and 950 °C shows slightly higher Na₂O and K₂O contents 539

(Table 3). This divergence between NIs and the calculated melts was also documented in other case
studies, and was ascribed to the lack of equilibrium with the residuum or diffusion in the melts, and
alternatively, the imperfections of forward phase equilibria modeling (Bartoli & Carvalho, 2021).

The P-T pseudosection reconstructed with a reintegration of 25 mol.% melts into the BRC is 543 shown in Figure 14b. The most evident changes are expansions of the stability fields of K-feldspar 544 and rutile, which were predicted to be stable during prograde melting. Despite the fact that 545 kyanite/sillimanite is absent in the investigated rocks, kyanite pseudomorphs (Tong & Wilson, 2006) 546 547 and kyanite relics replaced by sapphirine + quartz (Harley, 2021) have been identified in rocks from the same region, suggesting that the rocks may have been in the kyanite stability field. The quartz-, 548 kyanite-in and plagioclase-out curves suggest that the rocks underwent a nearly isobaric heating 549 from M0 to M1 stage (shown by dashed lines in Figure 13a). Accordingly, the Mather UHT 550 granulites may have experienced a clockwise P-T path with peak conditions of 12–13 kbar and 551 1070–1130 °C, followed by decompression and cooling to 6.5–7.0 kbar and 910–950 °C, and finally 552 isobaric cooling (IBC) to solidus conditions (P = 6.5-7.0 kbar; T = 830-870 °C). 553

554

555 **DISCUSSIONS**

556 *P–T* conditions of melt entrapment and anatectic regime

In East Antarctica, NIs in high–grade metamorphic rocks have been reported from the Lützow– Holm Complex (Spr–bearing *UHT* metapelitic granulites; Carvalho et al., 2023a), the Dronning Maud Land (ultramafic granulites; Ferrero et al., 2018), the Northern Victoria Land (metagreywackes; Ferri et al., 2020), and the Sør Rondane Mountains (East Antarctica) (Sil–Grt–Bt gneiss; Higashino & Kawakami, 2022). The present study expands the existing database to include the melts produced by *UHT* metamorphism of the Mg–rich and Sil–absent metapelitic rocks from the Rauer Islands.

The presence of kokchetavite and kumdykolite is confirmed by Raman analyses (Fig. 5g). Kokchetavite has been found within multiphase inclusions in high to ultrahigh pressure (*HP–UHP*) rocks, which was considered to crystalize from K–rich melts and may exist as a metastable phase (e.g. Borghini et al., 2018, 2023; Ferrero et al., 2016; Gao & Liu, 2018; Hwang et al., 2004, 2013). Kumdykolite was not only present as inclusions within *HP–UHP* rocks (e.g. Borghini et al., 2018, 2023; Ferrero et al., 2016; Gao & Liu, 2018; Hwang et al., 2009; Kotková et al., 2014), but also

found in chondrites (Németh et al., 2013). However, Ferrero et al. (2016) concluded that high 570 pressure is not required for the formation of kumdykolite and kokchetavite, and their presence is 571 thus not an indicator of UHP conditions. More recently, NIs from granulites have also been reported 572 to preserve kokchetavite and/or kumdykolite as daughter phases (Carvalho et al., 2019; Ferrero & 573 Angel, 2018; Gianola et al., 2021). Therefore, both kokchetavite and kumdykolite may have 574 crystalized from HT to UHT melts, and cannot be used to infer the P-T entrapment conditions. 575 According to Ferrero et al. (2016), the presence of feldspar polymorphs indicates that the melt 576 577 compositions (in terms of major elements and volatiles) are mostly preserved.

Melting during heating is predicted by phase equilibria modeling (Figs. 13, 14). Rutile is 578 present within NIs as an accidentally trapped phase (Fig. 5g). However, it is not observed in the 579 matrix of the rocks. This suggests that rutile existed in the pre-peak stage and became metastable as 580 a consequence of melt extraction and loss. After an integration of $\sim 25\%$ melts into the granulitic 581 restite, phase equilibria modeling predicts that rutile appears when temperature exceeds 950 °C at 582 12.5 kbar (Fig. 14), highly in agreement with the remelting experimental constraints given by NIs at 583 12.5 kbar. Furthermore, some NIs occur near to high-Al orthopyroxene inclusions (Al₂O₃ = 584 8.85-9.04 wt.%) within garnets (Fig. 3a). These are strong evidence that at least some NIs 585 investigated in this study were trapped at UHT conditions. Although the rocks studied here have 586 undergone peak temperatures of 1070-1130 °C, the melts may be produced at or near UHT 587 conditions (850–950 °C). 588

The H₂O contents of NIs show variable values ranging from 1.49 to 4.24 wt.% (Table 3), 589 suggestive of heterogeneous distributions of water (Bartoli et al., 2014) or existences of C-bearing 590 phases in some NIs (e.g. Carvalho et al., 2019; Ferri et al., 2020; Gianola et al., 2021). Estimation 591 of H₂O by difference of EMPA totals from 100% usually overestimates the H₂O contents of NIs (e.g. 592 Cesare et al., 2015), and specific reasons for this inconsistency still need further investigations. As 593 shown in Figures 9 and 10, the highly silicic, peraluminous and potassic to ultrapotassic NIs have 594 also been found in peritectic garnets from the Kerala Khondalite Belt (Cesare et al., 2009; Ferrero et 595 al., 2012), the Gruf Complex (Gianola et al., 2021) and the Lützow-Holm Complex (Carvalho et al., 596 2023a). Moreover, K-rich melts have also been experimentally derived from pelitic starting 597 materials under UHT conditions (e.g. Droop et al., 2003; Patiño-Douce & Johnston, 1991). Ti and 598 Fe³⁺ are expected to stabilize biotite to expand the Bt-dehydration melting to occur at higher 599

temperatures (e.g. White et al., 2014), which likely accounts for the enrichments of K_2O in HT to 600 UHT melts (Gianola et al., 2021). However, it is also evident that melt compositions of HT-UHT 601 granulites from the Ivrea Zone plot away from the K-apex of the triangle (Fig. 10), with low K₂O 602 contents of ~2.0 wt.% (Carvalho et al., 2019). The authors argued that the low-K melts may be 603 produced at P-T conditions approaching the Bt-out curves (Carvalho et al., 2019). Alternatively, 604 participation of plagioclase in the prograde melting reactions may also be responsible for the 605 relatively higher CaO/NaO concentrations in that study. The potassic nature of melts, coupled with 606 607 phase equilibria modeling (Fig. 14), supports that the melts were produced by Bt-dehydration melting of pelites under HT to UHT conditions (850–950 °C). 608

Our data show that the melts are enriched in Li, Cs, Rb, Ta, Sm, Nd, Zr, U and Pb and depleted 609 in Ce, Th, Ba, Sr and Nb relative to the granulitic restite (Table 4; Fig. 11). Dissolution of zircon or 610 monazite during high temperature melting (Kelsey & Powell, 2011) may contribute to the U, Pb and 611 Zr enrichments in anatectic melts. NIs in this study are depleted in Ce (<12.8 ppm) and Th (<3.8 612 ppm), which may be ascribed to the shielding of some monazites by porphyroblastic phases (Huang 613 et al., 2021) or disequilibrium melting (Watt & Harley, 1993). It suggests that the melts are likely 614 undersaturated with respect to monazite. Besides, the melts show mild depletion of Ba compared 615 with the restite. K-feldspar is the most important repository of Ba in the rocks and it was still stable 616 during melting, which might reserve appreciable amounts of Ba (Zeng et al., 2005). 617

The investigated melts contain higher Rb contents than the melts produced through 618 Ms-dehydration melting (Acosta-Vigil et al., 2010, 2012; Bartoli et al., 2019), consistent with 619 melts yielded by Bt-dehydration melting because biotite is the major reservoir of Rb (e.g. 620 Acosta-Vigil et al., 2010; Wolf et al., 2019). They also show higher Rb contents relative to the 621 matrix glasses from El Hoyazo (Fig. 11), which were considered to be yielded at the onset of 622 biotite-dehydration melting (Acosta-Vigil et al., 2010). Moreover, Sr contents are usually low in 623 micas but abundant in plagioclase (Weinberg & Hasalová, 2015), and plagioclase was not involved 624 in the incongruent melting in this study. Thus, higher Rb/Sr ratios of the investigated NIs can be 625 attributed to Bt-dehydration melting in the absence of plagioclase. On the other hand, although 626 Zeng et al. (2005) previously pointed out that that Ms/Bt-dehydration melting and dissolution of 627 apatite will produce melts with high Sm/Nd values, the mechanisms that caused elevated Sm/Nd 628 ratios in the melts are still vague. Assuming no isotopic fractionations of Nd and Sr, enrichments of 629

630 Sm and Rb in anatectic melts are expected to result in higher 143 Nd/ 144 Nd and 87 Sr/ 86 Sr values in 631 melts than the granulitic restite after enough long time.

632

633 Physical properties of the hottest crustal melts

The chemical data of NIs allow to peer into physical properties of the melts and therefore, may provide insights into rheology and mobility of the hottest granitic melts and their potential effects on orogenic evolution. Assuming conditions of P = 12 kbar and T = 950 °C, the average compositions of re–homogenized melt inclusions are considered to have densities of 2.47 ± 0.07 g/cm³, calculated with program DensityX (Iacovino & Till, 2019). These values are slightly higher than previously estimated densities for hydrous granitic melts (~2.20 g/cm³; Clemens & Vielzeuf, 1987), but lower than the average density for granites (~2.70 g/cm³; Artemieva et al., 2017).

Viscosity of melts may be affected by several potential factors, such as fluid contents (Holtz et 641 al., 2001; Scaillet et al., 1996) and melt compositions (Giordano et al., 2008). At 950 °C, 642 applications of the equations of Giordano et al. (2008) and Scaillet et al. (1996) yield melt 643 viscosities of $10^{4.1}-10^{6.1}$ Pa·s and $10^{4.0}-10^{5.2}$ for the investigated melts, respectively. When 644 considering the average melt compositions, these methods produce results of $10^{4.9\pm1.2}$ and $10^{4.6\pm0.8}$ 645 Pa·s. As the estimated H₂O contents for the melts based on difference of EMPA totals from 100% 646 usually slightly higher than their real values due to the possible presence of CO_2 (e.g. Cesare et al., 647 2015), the calculated viscosities may represent the minimum values. Viscosities for the investigated 648 melts are roughly in agreement with the calculation results for melt inclusions within UHT 649 granulites from the Gruf Complex $(10^{4.1}-10^{4.8} \text{ Pa}\cdot\text{s}; \text{Gianola et al., 2021})$. These values also overlap 650 the estimates of granitic melts that were experimentally derived at 5–8 kbar and 700–950 °C ($10^{4.3}$ – 651 10^{5.4} Pa-s) (Hess & Dingwell., 1996; Scaillet et al., 1996). However, melts formed at low 652 temperature (~700 °C) showed higher viscosities (~10^{6.0} Pa·s) (Bartoli et al., 2013b), and therefore 653 less mobile and result in slower rates of melt segregation and ascent (Bartoli et al., 2013b; Brown, 654 2013; Holtz et al., 2001). Clemens et al. (2020) concluded that melting at high temperatures 655 (mostly >850 °C) under fluid-deficient or -absent conditions are usually highly mobile, and 656 therefore, the low viscosity melts may facilitate melt extraction, and consequently crustal 657 differentiation and redistribution of heat-producing elements (such as K, Th and U) (Gianola et al., 658 2021). 659

The melts are enriched in heat producing elements such as K and U. Despite that the melts are depleted in Th, the heat production rate of Th (0.072 μ W/m³) is much lower than that of U (0.265 μ W/m³) (Bea, 2012). The capacity for heat production of the hot melts was determined with the following equation (Rybach, 1988):

664 $H = \rho \times (0.0952 \times C_U + 0.0348 \times C_K + 0.0256 \times C_{Th})$

665 Where H is heat production in μ W/m³, ρ is density in g/cm³, and C_U, C_K, and C_{Th} are 666 concentrations of radioactive isotopes of U, K, and Th in ppm, wt.% and ppm, respectively.

As calculated above, the density of melts is estimated as 2.47 ± 0.07 g/cm³. Calculated heat production value for the average melt composition is ~2.3 µW/m³. However, the obtained K, U and Th contents of melts cannot represent their real concentrations during the thermal event because of long–standing decay of the parental elements. Accordingly, heat production of average melt compositions was back-calculated to ~530 Ma based on appropriate decay equations for ²³⁸U, ²³⁵U, ²³²Th and ⁴⁰K. The resultant average value is ~2.8 µW/m³, which is higher compared with bulk heat production in granitic rocks of all ages (~2.0 µW/m³; Artemieva et al., 2017).

Granitic melts investigated in this study are particularly rich in heat-producing elements (such as K and U), and show densities of 2.47 ± 0.07 g/cm³ and viscosities of $10^{4.9\pm1.2}$ Pa•s. Such melts are mobile and prone to be extracted from the source, and as a consequence, might be responsible for the formation of large granitic intrusions. Stagnation and flow of such melts in the deep crust provides a potential source at the base of the crust to heat the orogenic hinterlands through radioactive decay of heat producing elements (Clark et al., 2011; Gianola et al., 2021), and may have a great influence on the thermal structure of the continental crust.

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Secondary fluid infiltration and its implications for the timing of UHT metamorphism: open questions

Fluid infiltration is a well-documented phenomenon in the Rauer Islands. For example, Harley (1998b) has suggested the growth of cordierites in Grt–Sil–Qz gneiss was linked with the ingress of CO₂-rich fluid ($X_{CO2} > 0.8$) based on secondary ion mass spectrometry analyses of H₂O and CO₂ in the secondary cordierite. On the contrary, coarser cordierites adjacent to leucosomes show lower X_{CO2} values of 0.05. Such features indicate that the fluid regime in this region has changed markedly as a consequence of ambient C–O–H fluid ingress. As zircon U–Pb systematics of the

layered mafic complex and intrusive tonalites from the nearby Scherbinina Island show clear 690 resetting at ~520 Ma, the fluid ingress was considered to have occurred at 530-510 Ma (Harley et 691 al., 1998; Harley & Kelly, 2007). On the other hand, Liu et al. (2021) reported that hornblende and 692 clinopyroxene in mafic granulites from the Filla Island were partially replaced by actinolite, which 693 was also ascribed by the authors to strong fluid activities during the Cambrian reworking. In this 694 study, we report on the occurrence of multiphase fluid inclusions (i.e. magnesite, pyrophyllite, 695 corundum and CO_2) within orthopyroxene and garnet (Fig. 7), which are considered to be secondary 696 697 in origin. Such stepdaughter phases were previously recognized in garnet-hosted primary FIs (e.g. Carvalho et al., 2020, 2023b; Gianola et al., 2021; Tacchetto et al., 2019). Here, we confirm that the 698 secondary C-O-H fluid along the healed fractures of garnet and orthopyroxene at high 699 temperatures could not survive neither, and has reacted with its hosts to produce CO₂ and 700 701 OH-bearing phases during cooling (possibly at 400-600 °C; Carvalho et al., 2020). Pure CO₂ inclusions within garnets are also present, which might represent the fluid that infiltrated into the 702 rocks at a very late stage. 703

In regard of the origin of the fluid, several potential sources can be considered. Harley et al. 704 (1998) speculated that the infiltration of CO₂-rich fluid might have occurred during the 705 decompression as a consequence of emplacements of granitic pegmatites in the Cambrian (Kinny et 706 al., 1993). Additionally, coeval graphite-bearing sediments (e.g. Harley & Fitzsimons, 1991) and 707 forsterite marbles (e.g. Buick et al., 1994; Harley, 1998b) have been reported in the Rauer Islands, 708 709 and their decarbonation may also be external sources for the fluid. Collectively, an external source for the C–O–H fluid is favored in this study. Fluid infiltration not only lead to possible transfer of 710 fluid-related elements (e.g. Liu et al., 2018), but also caused partial to complete resetting of U-Th-711 Pb systems in zircon and monazite (e.g. Das et al., 2021; Harley et al., 1998; Harley & Kelly, 2007; 712 Taylor et al., 2014). 713

As addressed on the section of geological setting, the timing of the *UHT* metamorphism is still a matter of debate. Constraints on its timing are critical to understanding the tectono--thermal evolution of the Rauer Islands, and even the whole Prydz Belt. Some works proposed that the Prydz Belt represents an early Neoproterozoic (~1000 Ma) metamorphic complex that was reworked by a Pan-African (~530 Ma) thermal event (e.g. Liu et al., 2009, 2014; Tong et al., 2014b, 2019; Wang et al., 2008), and the *UHT* metamorphism in the Rauer Islands occurred at ~1000 Ma (Tong &

Wilson, 2006; Wang et al., 2007). This point of view has not been accepted by other researchers, 720 who argued that this episode of UHT metamorphism may have taken place in the late 721 Neoproterozoic to Cambrian (e.g. Clark et al., 2019; Kelsey et al., 2003, 2007) or prior to ~590 Ma 722 (Hokada et al., 2016; Harley et al., 2009). Our dating results for zircon mantles show variable dates 723 ranging from 1050 to 607 Ma (Fig. 12e; Table 5). Previous U-Pb dating on Spr-bearing UHT 724 granulites from the same locality also showed similar age spreads from 1000 Ma to 500 Ma 725 (Hokada et al., 2016; Wang et al., 2007). Such scattered ages were previously interpreted as mixed 726 727 ages, detrital ages, or ascribed to variable effects of the Pan–African thermal event on pre–existing zircons (Hokada et al., 2016; Spreitzer et al., 2021; Wang et al., 2022). Although we cannot 728 preclude the possibility of melt-mediated-coupled dissolution-precipitation of zircons in the 729 Cambrian (Wang et al., 2022), findings of secondary FIs in peritectic phases (such as garnet, 730 731 orthopyroxene) and zircons in this study seem to support that the broad range of isotopic dates may be partially caused by extensive C–O–H fluid infiltration at high temperature conditions (>600 °C). 732 Till now, all the U-Th-Pb dating on monazite pointed to metamorphic ages of 580-510 Ma (Clark 733 et al., 2018; Hokada et al., 2016; Kelsey et al., 2003, 2007; Liu et al., 2021). Here, we prefer the 734 interpretation that U-Th-Pb clock of monazites has been completely reset as they were more 735 vulnerable to fluid ingress compared with zircons (e.g. Das et al., 2021; Taylor et al., 2014). 736

Additionally, garnet inclusions within zircon mantles from mafic granulites in the Archean 737 crustal domain indicate that the post-peak decompression and cooling probably occurred at 540-738 471 Ma (our own unpublished data), roughly consistent with the dates documented by zircon rims 739 in this study (569–472 Ma). If it is the case, the peak metamorphism took place at least prior to 740 ~570 Ma. Hokada et al. (2016) reported the microstructure and CL image of an 741 orthopyroxene-hosted zircon inclusion from the same outcrop, and it documents ages of 2502 Ma 742 in the core and 1254–1148 Ma in the rim. This suggests that this type of zircons has not been 743 affected by the Pan–African thermal event likely due to the shielding effect of orthopyroxene. Five 744 dark-CL mantles show ²⁰⁶Pb/²³⁸U dates of 1050-940 Ma (Table 5), which however, are slightly 745 discordant (Fig. 12f). Up to now, the most concordant ages for the Spr-bearing UHT granulites 746 from the Mather Peninsula were obtained by Wang et al. (2007), which yielded a weighted mean 747 207 Pb/ 206 Pb age of 995 ± 15 Ma. Nonetheless, their dating results also showed some discordance 748 from 1000 Ma toward 530 Ma (Wang et al., 2007). The present study highlights that C-O-H fluid 749

infiltration and alternatively, melt-mediated-coupled dissolution-precipitation during the Pan-750 African reworking, have contributed to the partial resetting of U-Th-Pb systems in zircons. 751 Collectively, the P-T framework (Fig. 15) might be linked with two separated thermal events that 752 happened at ~1000 Ma and ~530 Ma, respectively. We propose that the UHT metamorphism and 753 relevant melting most likely occurred prior to ~570 Ma (Harley et al., 2009; Hokada et al., 2016), 754 while the post-peak decompression and the final cooling happened at 570-470 Ma (Fig. 15). 755 However, it is still far from precisely determining the timing of UHT metamorphism in this region, 756 757 which requires further investigations.

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759 CONCLUSIONS

Phase equilibria modeling suggests that the Spr-bearing *UHT* granulites from the Mather Peninsula (East Antarctica) have undergone a clockwise P-T path, with peak conditions of 12–13 kbar and 1070–1130 °C. Melt and fluid inclusions are characterized in this study. Coupled with chronological dating results, the main conclusions drawn are described below:

1) Abundant NIs are preserved within garnet, which are usually 5–15 μ m in diameter and display typically negative crystal shapes. NIs have crystallized into polycrystalline aggregates, comprising daughter phases of quartz, biotite, kokchetavite, kumdykolite and an unknown phase (phase "430"). Petrographic observations and mineral chemistry suggest that the melts were at least partially yielded at *UHT* conditions. The NIs were perfectly re–homogenized with a piston cylinder apparatus at ~12 kbar and ~950 °C, and show SiO₂–rich, strongly peraluminous and potassic to ultrapotassic composition.

2) Compared with the granulitic restite, the NIs are enriched in Li, Cs, Rb, Ta, Sm, Nd, Zr, U 771 and Pb, and depleted in Ce, Th, Ba, Sr and Nb. Melting mechanisms and dissolution of accessory 772 773 minerals played a key role in the partitioning of elements between anatectic melts and their granulitic counterparts. High Rb/Sr and Sm/Nd ratios for the melts, coupled with phase equilibria 774 modeling, suggest that the melts were formed by fluid-absent Bt-dehydration melting. These UHT 775 crustal melts have densities of 2.47 \pm 0.07 g/cm³, viscosities of 10^{4.9\pm1.2} Pa·s and high heat 776 production values of ~2.8 μ W/m³. As a consequence, these melts are expected to be mobile and 777 may have played an important role in redistribution of heat-producing elements (such as K and U). 778

3) Secondary FIs are distributed along healed fractures of the peritectic phases (such as garnet

and orthopyroxene), typically comprising magnesite, corundum, pyrophyllite and residual CO_2 780 $(\rho_{CO2} = 0.2 - 1.0 \text{ g/cm}^3)$. They are interpreted as stepdaughter phases due to interaction between 781 C-O-H fluid and their hosts during post-peak metamorphic evolutions. Pure CO₂ inclusions with a 782 density of ~ 0.8 are also present, representing the fluid that infiltrated the rock at a very late stage. 783 Ingress of the C–O–H fluid under high temperatures probably resulted in mobility and loss of Pb in 784 zircons, and was likely responsible for the large date span of zircon mantles from 1050 to 607 Ma. 785 Another possible reason is melt-mediated-coupled dissolution-precipitation during the Pan-786 787 African reworking. Therefore, the retrieved P-T path might be linked with two separated thermal events happened at ~ 1000 and ~ 530 Ma, respectively. Although metamorphic timing for the UHT 788 metamorphism is still debated, we conclude that the peak metamorphism occurred prior to ~ 570 789 Ma. 790

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- 1243

1244 TABLE CAPTIONS

Table 1 Representative mineral compositions of the Mather *UHT* granulites. The major element

1247	compositions are	shown in	wt.%.

	G	G	G	G	0	0	0	Op	0	0	Sp	S	С	В	В	В	В	Р	Pl	S	Pl	Bt	Op	
	rt	rt	rt	rt	р	р	р	X _{ex}	р	р	r _{wi}	pr	rd	ti	t _p	ti	t-	l_i	3	р	-in	-in	x-i	
	-с	-	-r	3	x _{in}	х	х	solv	xi	х	th	wit	int	n	orp	nt	3	nt		1	Zr	Zr	n	
		m			Grt	-	-r	ed	nte	3	Crd	h	er	Gr	hy	er		er			n	n	Zr	
						с			r			Bt		t								× .	n	
S	4	4	4	3	5	5	4	51	5	5	13	13	4	3	3	3	3	4	5	0	59	38	50.	
i	0.	0.	0.	9.	1.	2.	9	.0	0.	0	.2	.4	9.	9.	6.	6	7	7	5.	\.``	.8	.5	28	
0	9	5	6	9	1	3		7	2		7	1	2	7	6				1	0	7	1		
2	4	1	3	4	4	4	1		0	6			0	1	7	6	4	0	9	4				
							1			8						9	8	3						
Т	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	3.	4.	4)4	0	0.	0	0.	3.	0.0	
i	0	0	0	0	0	0		11	0		05	10	0	4	0				0		01	54	5	
0	3	6	0	0	4	7	0		9	0			3	7	8	3	1	0	0	0				
2							8			6						2	2	0		4				
А	2	2	2	2	8.	6.	9	6.	7.	7	62	61	3	1	1	1	1	3	2	6	24	16	6.7	
l_2	3.	3.	3.	2.	8	4		88	3		.2	.3	4.	5.	6.	7	6	3	9.	1	.9	.9	9	
0	2	0	1	3	5	9	1		7	6	2	4	0	4	9				0		5	6		
3	3	1	1	8			6			7			3	9	3	1	8	7	4	1				
																3	8	3		1				
С	0.	0.	0.	0.	0.	0.	0	0	0.	Ø	0.	0.	0.	0.	0.	0	0	0	0.	1	0.	0.	0.1	
\mathbf{r}_2	1	1	0	0	1	0	• _	07	`		32	49	0	0	1				0		06	08	1	
0	2	0	9	1	0	3	0	$\backslash acket$	2	0			6	9	3	2	1	0	0	2				
3							3	\sum		7						5	0	0		8				
F	1	1	1	2	1	1	1	15	1	1	7.	8.	2.	5.	8.	8	8	0	0.	2	0.	11	17.	
e	7.	8.	9.	3.	2.	3.	4	.2	5.	5	79	51	6	4	2				2	1	03	.5	62	
Т	2	1	7	8	0	0	•	1	2	•			1	4	5	9	6	2	5			3		
0	6	7	3	4	4	6	9		4	4						2	8	4		8				
				Y			3			9										5				
Μ	0.	0.	9.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0		0.	0.0	
n	1	1	2	3	0	0		07	0	•	04	06	0	0	0	•	•		0			06	8	
0	8	6	6	8	4	3	1		8	1			3	0	2	0	0	0	1	0				
							0			1						0	1	2		8				
M	1	1	1	1	2	2	2	26	2	2	16	16	1	2	1	1	1	0	0.	1	0.	15	24.	
g	6.	5.	5.	1.	7.	7.	5	.2	7.	5	.2	.7	2.	1.	8.	7	8		1	2	00	.5	69	
0	8	9	0	8	3	6		3	0	•	6	0	6	9	2	•	•	0	3	•		8		
	7	6	0	1	1	1	5		0	1			0	0	4	7	1	0		7				
							8			7						9	6			1				
С	1.	1.	1.	1.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	1	1	0	7.	0.	0.1	

	a	4	1	6	7	1	0		12	0		00	02	0	0	0			6	0.		15	00	0	
	0	7	8	3	8	0	9	1		9	0			3	2	2	0	0		4	0				
								1			7						0	0	4	0	0				
																			6						
	Ν	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	2	5.	0	7.	0.	0.0	
	а	0	0	0	0	0	0		03	0		02	00	0	4	4				3		24	12	1	
	2	1	3	2	2	5	0	0		2	0			4	3	4	3	4	1	3	0				
	0							0			4						3	1	4		8			V.	/
	Κ	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	9.	9.	9	9	0	0.	0	0.	8.	0.0	
	2	0	0	0	0	0	0		00	0		00	00	0	4	9				5	•	16	56	0	
	0	0	0	0	1	0	0	0		0	0			1	6	6	5	8	0	3	0				
								1			3						2	5	3	X	2				
	Ζ	Ι		_		_	-	-	_	_	_	_	Ι	-	-	_		-	_	-	1	_	_	_	
	n																								
	0																	K			3				
																	< T		Y		7				
	F	—	_	_	_	—	_	_	—	_	—	—	_	_	0.	0.	0	Ø	—	_	_	—	—	—	
															3	0									
															7	9	2	2							
																	9	2							
	С	—	_	_	_	—	_	_	—	_	—	—		Ê)	0.	0.	0	0	—	_	_	—	—	—	
	1											\checkmark			1	1									
												\sim	Y		0	2	1	1							
												, ,					1	2							
	Т	1	9	1	1	9	9	9	99	1	9	-99	10	9	9	9	9	9	9	1	9	99	94	99.	
	0	0	9.	0	0	9.	9.	9	.74	0	9	.9	0.	8.	6.	4.	5	6	9	0	8	.5	.9	72	
	ta	0.	1	0.	0.	6	7	•	9	0.		7	63	6	4	9				0.		2	5		
	1	1	8	4	1	7	2	0	$\langle \rangle$	2	3			3	9	4	3	0	6	8	5				
		1		7	7			9	\mathcal{A}	1	9						4	3	6	7	8				
	0	1	1	1	1	6	6	6	6	6	6	20	20	1	1	1	1	1	8	8	4	8	11	6	
		2	2	2	2		Y							8	1	1	1	1							
	S	2.	2.	2.	3.	1	1.	1	1.	1.	1	1.	1.	4.	2.	2.	2	2	2	2.	0	2.	2.	1.8	
	i	9	9	9	0	8	8	•	83	7		57	58	9	8	6	•			4		67	80	26	
		8	9	8	Ŏ	1	6	7	4	9	8	8	7	2	0	8	6	7	1	6	0	9	7		
		4	5	7	9	5	4	7		1	3			1	2	1	7	1	6	8	0				
		$ \land $						3			3						6	0	5		1				
	T	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
$\left(\right)$	i	0	0	0	0	0	0		00	0		00	00	0	1	2				0		00	19	01	
	\mathcal{I}	0	0	0	0	0	0	0	3	0	0	4	9	0	8	2	2	2	0	0	0	0	4		
		2	3	0	0	1	2	0		2	0			2	4	4	3	2	0	0	0				
								2			2						7	4	0		1				
	Α	1.	2.	2.	1.	0.	0.	0	0.	0.	0	8.	8.	4.	1.	1.	1	1	1	1.	1	1.	1.	0.2	
	1	9	0	0	9	3	2		29	3		72	56	0	2	4				5		31	45	91	
		9	0	0	8	7	7	3	1	1	3	3	1	1	8	5	4	4	8	3	9	6	8		
		9	0	0	8	/	/	3	I	1	3	3	I	1	8	5	4	4	8	3	9	6	8		

		6	5	3	8	0	2	9		0	2			2	9	9	7	3	3	1	3				
								0			7						2	9	1		6				
	С	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
	r	0	0	0	0	0	0		00	0		03	04	0	0	0				0		00	00	03	
	_	0	0	0	0	0	0	0	2	0	0	0	6	0	0	0	0	0	0	0	0	2	4		
		7	6	5	1	3	1	0	2	3	0	Ū	U	5	5	8	1	0	0	0	2	2	-		
		/	0	5	1	5	1	1		5	2			5	5	0	5	6	0	0	27				
	Б	0	0	0	0	0	0	1	0	0		0	0	0	0	0	5	0	0	0	/	0	0		Y
	Г	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
	e 3⊥	0	0	0	0	0	0	•	03	0	•	08	20	0	0	0	•	•	•	0	•	00	00	51	
	51	2	0	2	0	0	0	0	5	9	0	1	1	4	0	0	0	0	0	0	0		0		
		8	0	0	0	0	0	6		1	0			4	0	0	0	0	0	9	3				
								0			5						0	0	9	\checkmark	7				
	F	1.	1.	1.	1.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.4	
	e	0	1	1	5	3	3		42	3	•	68	64	1	3	5	•			0		00	70	84	
	2+	2	2	9	0	5	8	3	2	6	4	8	2	7	2	0	5	5	0	0	4	0	3		
		4	3	3	2	7	9	9		4	6			5	1	4	4	2	0	0	5				
								1			4						4	5	0		6				
	Μ	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
	n	0	0	0	0	0	0		00	0		00	00	0	0	0				0		00	00	02	
		1	1	1	2	0	0	0	2	0	0	4	6	0	0	0	0	0	0	0	0	1	4		
		1	0	6	4	1	1	0		2	0			3	0	1	0	0	0	0	0				
								3			3		$\langle \cdot \rangle$	Y			0	1	1		2				
	М	1.	1.	1.	1.	1.	1.	1	1.	1.	1	2	2.	1.	2.	1.	1	1	0	0.	0	0.	1.	1.3	
	σ	8	7	6	3	4	4	_	40	4		88	94	8	3	9	_			0		00	69	37	
	D	3	5	4	2	4	6	3	4	3		2	6	7	0	8	. 9	9	0	0	5	0	2	57	
		2	8	4	6	4	5	7		6	5	-	Ū	, 8	3	7	3	5	0	9	0	Ŭ	2		
		2	0	-	0	-	5	6			7			0	5	,	1	7	0)	0				
	C	0	0	0	0	0	0	0			/	0	0	0	0	0	4	/	0	0)	0	0	0.0	
	C	U.	0.	1.	0.	0.	0.	0		0.	0	0.	0.	0.	0.	0.	U	0	0	0.	0	0.	0.	0.0	
	а	1	0		1		0		-00	0	•	00	00	0	0	0	•	•	•	4	•	34 2	00	04	
		l z	9	2	4	0	0	0	5	0	0	0	3	0	0	0	0	0	8	9	0	3	0		
		5	3	8	4	4	3	0		3	0			3	2	2	0	0	1	8	0				
					\square			4	_		3						0	0	2		0				
	Ν	0.	0.	0.	0.	$\Box 0.$	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
	а	0	0	0	0	0	0	•	00	0	•	00	00	0	0	0	•	•	•	4	•	62	01	00	
		0	0		0	0	0	0	2	0	0	5	0	0	5	6	0	0	1	6	0	8	7		
		1	4	3	3	3	0	0		1	0			8	8	3	4	5	9	2	0				
	\checkmark		/					0			3						7	8	1		4				
\bigcap	K	0.	0.	0.	0.	0.	0.	0	0.	0.	0	0.	0.	0.	0.	0.	0	0	0	0.	0	0.	0.	0.0	
	\mathcal{I}	0	0	0	0	0	0		00	0		00	00	0	8	9				0		00	79	00	
		0	0	0	0	0	0	0	0	0	0	0	0	0	5	2	8	9	0	3	0	9	6		
		0	0	0	1	0	0	0		0	0			1	2	9	8	0	0	0	0				
								0			1						5	9	2		1				
	Ζ	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	0	_	_	_	
	n																								
	L		1	1	I	l	1	I		I	1		l	I			1	1							



	Bt	Kok	Kum
SiO ₂	39.22	75.17	66.34
TiO ₂	0.50	0.00	0.03
Al ₂ O ₃	19.22	11.89	19.5
Cr ₂ O ₃	0.18	0.00	0.01
Fe ^T O	2.88	0.75	0.83
MnO	0.02	0.03	0
MgO	23.88	0.04	1.67
CaO	0.06	0.02	1.46
Na ₂ O	0.73	0.44	8.38
K ₂ O	9.29	10.85	0.71
F	0.00	-	× -
Cl	0.02	-	_
Total	95.99	99.21	98.93
0	11	8	8
Si	2.729	3.345	2.934
Ti	0.026	0.000	0.001
Al	1.576	0.624	1.017
Cr	0.010	0.000	0.000
Fe ³⁺	0.025	0.028	0.031
Fe ²⁺	0.142	0.000	0.000
MnO	0.001	0.001	0.000
Mg	2.477	0.003	0.110
Ca	0.004	0.001	0.069
Na	0.098	0.038	0.719
K	0.824	0.616	0.040
F	0.000	_	-
Cl	0.002	_	_
Total	7.914	4.656	4.921
X _{Mg}	0.95		
X _{An}		0.94	0.08
	1	1	

Table 2 Compositions of daughter phases within NIs. The major element compositions are shown in

1257 wt.%.

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	ave	σ	PE	
																	Μ	
SiO ₂	73.	70.	72.	69.	70.	69.	73.	75.	72.	74.	73.	71.	73.	70.	72.	1.	69.	
	60	36	93	94	99	89	16	61	78	94	14	33	86	48	36	80	33	
TiO ₂	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	0	5	4	7	3	1	2	0	6	6	0	4	2	3	3	02	0	
Al_2O_3	14.	14.	15.	15.	14.	14.	12.	13.	13.	13.	15.	14.	13.	14.	14.	0.	14.	
	28	21	21	28	30	49	70	47	33	05	06	21	89	52	14	76	19	
Cr_2O_3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	0	4	0	0	0	2	0	1	0	0	0	3	0	2	1	01	0	
Fe ^T O	1.1	1.1	1.2	1.1	1.2	1.4	1.0	0.9	1.0	0.9	1.3	1.4	1.1	1.0	1.1	0.	0.4	
	0	2	2	1	2	3	3	8	0	9	3	4	1	9	5	15	8	
MnO	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	0	2	2	4	0	0	0	0	1	2	0	Ø	2	4	1	02	0	
MgO	0.5	1.2	0.7	1.3	0.8	0.9	0.3	0.4	0.4	0.4	0.6	1.8	0.5	0.5	0.7	0.	0.2	
	4	3	7	4	3	4	9	1	1		2	7	8	4	8	42	9	
CaO	0.3	0.4	0.4	0.4	0.5	0.3	0.3	0.3	0.5	0.4	0.6	0.5	0.3	0.3	0.4	0.	0.6	
	5	5	3	6	6	2	6	8	8	1	3	1	9	0	4	10	3	
Na ₂ O	1.2	0.9	0.9	1.0	1.3	1.3	0.9	0.9	1.0	1.3	1.6	0.7	0.6	0.6	1.0	0.	1.7	
	6	4	9	6	0	9	9	9. '	5	1	3	4	4	1	6	28	0	
K ₂ O	6.5	8.4	6.5	8.4	6.4	7.5	7.3	6.6	7.1	6.6	5.4	7.4	7.2	8.6	7.1	0.	8.5	
	3	3	0	7	7	2	3	0	5	8	6	0	0	2	7	86	3	
Cl	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	5	5	6	5	6	4	6	6	6	5	6	4	8	2	5	01	0	
P_2O_5	0.0	0.0	0.0	0.0	0,0>	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	0	2	0	1	0	0	0	0	0	0	3	0	3	0	1	01	0	
Total	97.	96.	98.	97.	95.	96.	96.	98.	96.	97.	97.	97.	97.	96.	97.	0.	96.	
	71	91	18	84	76	05	04	51	44	92	95	61	80	29	21	89	29	
H ₂ O by	2.2	3.0	1.8	2.1	4.2	3.9	3.9	1.4	3.5	2.0	2.0	2.3	2.2	3.7	2.7	0.	4.8	
diff	9	9	2	6	4	5	6	9	6	8	5	9	0	1	9	89	5	
ASI	1.4	1.2	1.6	1.3	1.4	1.3	1.2	1.4	1.2	1.2	1.5	1.4	1.4	1.3	1.4	0.	1.0	
	6	4	1	0	1	2	4	2	6	9	5	0	5	3	0	11	8	
AI	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.	0.0	
	3	2	4	3	3	2	2	3	2	2	4	3	3	2	3	01	1	
Na ₂ O+K	7.7	9.3	7.4	9.5	7.7	8.9	8.3	7.5	8.2	7.9	7.0	8.1	7.8	9.2	8.2	0.	10.	
₂ O	9	7	9	3	7	1	2	9	1	9	8	4	4	3	3	72	23	
K/Na	3.4	5.9	4.3	5.2	3.2	3.5	4.8	4.3	4.4	3.3	2.2	6.5	7.4	9.3	4.4	1.	3.3	
	0	0	0	5	9	7	5	8	6	4	1	9	4	1	3	89	1	
Qz norm	39.	29.	39.	28.	36.	31.	38.	43.	37.	40.	40.	34.	40.	32.	36.	4.	25.	
	73	93	93	14	84	02	69	01	99	32	33	53	99	80	75	48	86	
Crn	4.6	2.8	5.8	3.6	4.3	3.6	2.5	4.0	2.8	2.9	5.5	4.1	4.5	3.7	3.9	0.	1.0	ĺ

1264 **Table 3** Major element concentrations of re-homogenized NIs. The compositions are shown in

1265 wt.%.

	norm	1	6	5	5	4	4	7	7	9	8	2	8	0	8	6	94	7	
	Or norm	39.	51.	39.	51.	39.	46.	45.	39.	43.	40.	32.	44.	43.	52.	43.	5.	52.	
		50	44	15	17	97	31	12	64	86	31	93	83	53	96	60	40	98	
	Ab	10.	8.2	8.5	9.1	11.	12.	8.7	8.5	9.2	11.	14.	6.4	5.5	5.3	9.2	2.	15.	
	norm	95	1	8	9	47	22	6	2	6	36	07	1	2	6	7	45	12	
	An	1.7	2.1	2.1	2.2	2.9	1.6	1.8	1.9	3.0	2.0	3.0	2.5	1.8	1.5	2.1	0.	3.2	
	norm	6	9	9	3	0	3	9	1	1	8	1	5	1	4	9	48	8	
	Density	2.4	2.4	2.5	2.5	2.4	2.4	2.4	2.5	2.4	2.4	2.5	2.5	2.4	2.4	2.4	0.	2.3	
		8	7	1	1	2	3	1	1	3	8	0	0	9	4	7	03	0	
	Viscosity	10 ⁴	10 ⁴	10 ⁵	10 ⁴	10 ⁴	10^{4}	10 ⁴	10 ⁵	10 ⁴	104	10	10 ³						
	(S96)	.8	.5	.0	.9	.0	.1	.1	.2	.3	.9	.9	.8	.9	.2	.6	0.4	.8	
	Viscosity	10 ⁵	10 ⁴	10 ⁵	10 ⁵	10 ⁴	10 ⁴	10 ⁴	10 ⁶	10 ⁴	10 ⁵	10 ⁵	10 ⁴	10 ⁵	< 10 ⁴	10^{4}	10	10^{3}	
	(G08)	.3	.5	.6	.0	.1	.1	.4	.1	.6	.5	.4	.9	.4	, A	.9	0.6	.7	
	ASI value	es are o	calcula	nted or	n anhv	drous	basis	S96	Scai	llet et	al (19	96)· (liorda	no et a				
	(2008).						045151	270	,			,,,,							
	PEM, melt compositions calculated by phase equilibria modeling at 950 °C with 25 mol% melts reintegrated back into bulk rock compositions.																		
	PEM, melt compositions calculated by phase equilibria modeling at 950 °C with 25 mol% melts reintegrated back into bulk rock compositions.																		
1200	back into bulk rock compositions.																		
1266	back into bulk rock compositions.																		
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1278	Δ																		
1279	Y																		
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1200																			
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1202																			
1283																			

1284	Table 4 Trace element	concentrations of	f NIs an	nd the granu	ilitic restite	from the	Mather	Peninsula.
			•••					

1285 The trace element compositions are shown in ppm.

	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	a	σ	W	
	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	v		R	
	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	e		C	h.,
	0	0	0	1	1	1	1	2	2	2	2	3	4	0	0	2	1	4	4	4	4	1				
	5	7	9	0	2	6	8	1	4	5	7	8	8	3	8	6	3	4	6	7	2	1				
														*	*	*	*	*	*	*	*	*			Y	
R	2	7	3	7	7	6	7	2	3	2	1	1	6	4	7	1	1	3	4	1	4	6	5	3	1	
b	6	5	8	0	1	7	3	8	8	1	5	2	0	1	9	1	3	8	8	0	0	9)7	1	6	
	8	5	7	8	1	6	2	2	5	2	6	6	5	1.	8.	3	2	0.	9.	7	2.	1.	8	2	2	
	•	•	•	•		•	•	•	•	•	•	•	•	1	8	1.	5.	3	2	(7.)	9	•	•		
	8	9	1	9	7	6	7	3	7	2	6	7	0			7	0			4			1	9	4	
С	0	5	2	9	2	1	4	3	6	1	1	b	2	2.	3.	b.	1	3.	9.	3	3.	2.	7	8	1	
S	•	•	•	•		4	•	•	•	•	•	•	6	3	3	d.	5.	4	0	5.	5	9	•		•	
	9	6	6	5	8	· _	6	2	9	8	8	d	•			l.	3	Y		5			8	8	2	
						7						l.	1													
D	1	2	1	1	2	5	1	1	0	4	2	•	2	1	6		1	4	5	4	2	2	2	4	7	
В	1 0	2			2	5 6		1	9	4	3	2			3	2	4	4	5 6	4	3	2	5	4	 6	
a	0	0		4	0	0	0	1	1	4	с С	0	4		~	5 6	っ っ	0	0 5	0	2	7	07	3 7	0	
	0	3	/	3	9	3	2	4	• 1	0	Ζ		0	9.	0. 3	0	2. 8	2. 3	5. 6	3. 7	5. 7	J. 1	/	/	9	
	· 2		6	3	7	5	3	7	1	7		•	$\dot{\mathbf{x}}$	[′	5	4. 7	0	5	0	/	/	1	7	· 2	5	
T	1	b	1	<u>з</u>	3	h	<u>з</u>	, h	2	h	1	Б	$\frac{2}{2}$	8	7	$\frac{7}{2}$	3	h	1	h	3	h	γ Δ	$\frac{2}{4}$	1	
i	8	U	3	6	1	U	$\frac{1}{2}$	U	$\frac{z}{x}$		6		1	6.	6	$\begin{bmatrix} 2\\ 0 \end{bmatrix}$	8	d.	7	d.	3	d.	$\frac{1}{2}$	7	5	
•		d				d	_	d				d		Ŭ	9	0.	8	1.	8	1.	9	1.	_	,		
	8	.1	5	5	5	.1	7	Λ.	8	.1	0	.1	5			8			-		-		5	0	6	
)																	
Р	7	1	6	b	1	5	b	8	3	9	1	8	1	n.	n.	n.	n.	n.	n.	n.	n.	n.	8	2	2	
b		0			0						0		2	С	С	С	С	С	С	С	С	С				
	2		3	đ	. \	8	d	9	9	9	•	6											5	4	1	
		4		.1	Ì		.1				1		8													
				X			•																			
Т	0	0	ን '	0	b	b	b	0	b	3	2	3	1	n.	n.	n.	n.	n.	n.	n.	n.	n.	1	1	4	
h		·	•	•	•	•	•	•	•	•	•	•	•	С	С	С	С	С	С	С	С	С	•	•		
\sim	7	4	d	8	d	d	d	6	d	8	8	0	7										7	2	5	
			.l		.l	.l	.l		.l																	
		_	•	<u> </u>	•	•			•				0									-			-	
U	4	5	2	4	6	1	0	2	1	8	6	6	9	n.	n.	n.	n.	n.	n.	n.	n.	n.	4	2	0	
	•			•	•	•	•	•	•	•		•	•	С	С	С	С	С	С	С	С	С	•	•	•	
NT	8	2	5	9	4	6	0	1	8	/	5	8	5										/	/	4	
IN 1-	9	8	5	6		4	/	8	4	1	8		8	n.	n.	n.	n.	n.	n.	n.	n.	n.	8	5		
D	•	•	•	•	С	•	•	•	•	4	•	1	•	С	С	С	С	С	С	С	С	С	•	•	С	

	r	1	r	r	r	1	-	-	-	r	1	r	1	r	r	r	1	-	-	-			-	-		
	3	7	6	2	7	2	8	4	0	3	4		0										6	4	Д	
Т	3	3	1	b	b	b	2	1	2	3	b	1	2	n.	n.	n.	n.	n.	n.	n.	n.	n.	2	0	1	
a	•		•							•		•		С	С	С	С	С	С	С	С	С			•	
	2	2	4	d	d	d	0	8	0	2	d	8	7										4	7	0	2
				l.	l.	l.					1.															
Т	9	1	7	9	9	· 3	4	4	2	7	6	7	3	n.	n.	n.	n.	n.	n.	n.	n.	n.	6	2	n	
i	9	0	9	6	0	3	1	7	7	3	5	0	1	С	С	С	С	С	С	С	С	С	6	6		
	7	5	3	3	4	4	9	0	4	4	4	5	4										2	5	С	
	5	2		•	. 2	· 2	• 1	5	•		5		5								C).'	•		
	5	6	0	1	5	2	1	5	1	0	5	5	5							$\langle \checkmark$			9	1		
С	7	1	7	6	5	4	b	3	2	1	5	1	8	n.	n.	n.	n.	n.	n.	n.	n.	n.	7	3	1	
e		0								3	•	2	•	С	С	С	С	с	¢	С	С	С			6	
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		0					.1			4		0													0	
S	1	1	1	1	2	8	1	1	7	2	1	1	2	n.	n.	n.	n.	n.	n.	n.	n.	n.	1	5	1	
r	6	7	1	7	4		4	0		7	9	7	0	e	C	c	С	С	С	С	С	С	6		9	
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	2			d		3	d	d	d	$\langle \cdot \rangle$	7		•											0	1	
		7	0	.1	8		.1	.1	A	8		7	2										9			
S	5	1	7	· 2	1	h	· h	h	·	1	1	1	h	n	n	n	n	n	n	n	n	n	1	6	1	
m		2		7	1					9	1	5		п. С	п. С	п. С	<i>с</i>	<i>с</i>	п. С	<i>с</i>	п. С	п. С	3			
	5		4		•	d	d	d	d				d											4	8	
		6		2	8	.1	.1	.1	.1	0	1	3	.1										7			
7	1	1	1	1	$ \rightarrow $	/. 8	2	1	1	2	2	1	0	12	12	11	11	n	11	n	11	11	1	Δ	8	
r	9	8	1	7	9	4	$\begin{bmatrix} 2\\ 0 \end{bmatrix}$	6	2	5	7	8	6	п. С	п. С	п. С	п. С	п. С	п. С	п. С	п. С	п. С	7	7	4	
	4	5	2	3	7		6	3	1	6	6	2											7			
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ч	9	1	9 1	9 12	$\frac{0}{2}$	h	3	8 1] h	0	5	8	1	10	10	70	10	70	70	70	70	70	8	2	n	
f	0	ש א	0	0		0	ש	0	0	$\frac{1}{2}$) .	5	$\begin{vmatrix} 1 \\ 2 \end{vmatrix}$	п. С	п. С	c	r	п. С	п. С	п. С	п. С	п. С	0		2	
-		7	d	d	6	d	0	d	d		9	4		C	C	C	Ũ	C	C	Ū	e	e	6	4	2	
	7		.1	.1		.1		.1	.1	5			7													
Б	1.	2	•	•	1.	•	1.	•		1.	1.	1.	1.										2	0	0	
E U	D	2	D	D	D	D	D	D	5	D	D	D	D	n. C	n. C	п. С	n.	n.	п. С	n. C	n. C	n. C	2	0	U	
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																										I
G	6	b	1	b	b	1	b	2	b	b	b	2	2	n	n	n	n	n	n	n	n	n	1	6	2	I
d	Ŭ		1	U		7		1			U	3	5	с.	с.	<i>C</i>	<i>c</i>	с.	с.	с.	с.	<i>с</i>	7	Ŭ	2	I
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		1.	9	l.	l.	1	l.	9	l.	l.	l.	9	1										7		5	
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V	b	b	5	b	b	2	1	1	5	3	4	7	1	n.	n.	n.	п.	n.	<i>n</i> .	n.	n.	<i>n</i> .	6	3	2	i i
			2			9	8	0	5	5	6	9	4	С	С	С	С	С	С	С	С	C	2	8	7	I
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n		3				1							3	e	C	c	С	С	С	С	С	С	1	0	1	i i
	d		d	d	d	9	d	d	d	d	d	d	0	$\langle \rangle$						-			1		4	i i
	1		1	1	1	ĺ	1	1	1	1	1	1	X										1	. 3		i i
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S	b	b	I	b	b	b	b	4	b	b	9		b	n.	n.	n.	n.	n.	n.	n.	n.	n.	1	1	1	I
с	•	•	2	•	•	•	•	2	•	$\langle \cdot \rangle$	\cdot	3	•	С	С	С	С	С	С	С	С	С	9	3	9	I
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		0	2					1	1	2	2	0	L F										5	7	0	1
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S	0	0	0		0					1	1	0											0	0	0	1
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/	7	7	7		5					1	1	8											8	2	2	I

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	N	6	6	4		2					3	4	2													5	
	d U	6	1		6				3		2	2	2	5										5	3	0	
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	h	2	4 7		9				4		1	2	1	1												9	
	b.d	l.l. b	elov	<i>w</i> de	etect	ion	limi	it; <i>n</i> .	c n	ot co	onsi	dere	ed; V	VRO	C , w	hole	-rocl	k coi	npo	sitio	n; *	Nan	ogra	anite	oid		7
	inc	lusi	ons	with	n ac	cide	ntal	ly tr	app	ed p	bhas	es s	uch	as z	zirco	n, m	onaz	zite,	apat	ite a	nd r	utile	e.	2		, 7	
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	Th	U	Th/	²⁰⁶ Pb/ ²³⁸	1σ	²⁰⁷ Pb/ ²⁰⁶	1σ	²⁰⁷ Pb/ ²³⁵	1σ	²⁰⁶ Pb/ ²³⁸	1	²⁰⁷ Pb/ ²⁰⁶	1σ	²⁰⁷ Pb/ ²³⁵	1	
Spot			U	U		Pb		U		U	σ	Pb		U	σ	
	(pp	(pp								(Ma)		(Ma)		(Ma)		
	m)	m)														
Rim																
1	49	145	0.34	0.076	###	0.060	0.00	0.633	###	472	8	611	15	498	2	
					#		4		#				7		5	
2	45	167	0.27	0.080	###	0.057	0.00	0.621	###	497	8	480	12	490	2	
					#		3		#				0		0	
3	40	143	0.28	0.081	###	0.059	0.00	0.656	###	500	9	565	13	512	2	
					#		4		#						5	
4	98	437	0.22	0.081	###	0.062	0.00	0.699	###	502	7	676	96	538	1	
					#		3		#						8	
5	47	420	0.11	0.091	###	0.065	0.00	0.842	###	564	#	770	96	620	2	
					#		3		#		#				4	
6	57	271	0.21	0.092	###	0.061	0.00	0.774	### /	569	#	657	10	582	2	
					#		3		#	$\mathbf{\mathbf{Y}}$	#		0		0	
Mant																
le																
7	8	877	0.01	0.099	###	0.062	0.00	0.848	###	607	5	661	66	624	1	
					#		2		#						3	
8	23	481	0.05	0.101	###	0.062	0.00	0.920	###	623	#	687	95	662	3	
					#		3		#		#				2	
9	18	####	0.02	0.120	###	0.068	0.00	1.136	###	730	8	865	65	771	1	
					#		2		#						8	
10	32	659	0.05	0.120	###	0.067	0.00	1.115	###	730	8	843	70	761	1	
					#	<i>y</i>	2		#						8	
11	18	811	0.02	0.121	###	0.066	0.00	1.110	###	735	8	1200	71	758	1	
					#		2		#						8	
12	22	####	0.02	0.122	###	0.066	0.00	1.123	###	743	7	817	59	765	1	
				/	#		2		#						5	
13	48	524	0.09	0.123	###	0.070	0.00	1.210	###	750	#	937	67	805	2	
	$\left(\right)$	$\langle \rangle$	<i></i>		#		2		#		#				1	
14	42	521	0.08	0.124	###	0.065	0.00	1.115	###	754	#	759	70	760	2	
					#		2		#		#				1	
15	31	####	0.03	0.130	###	0.072	0.00	1.301	###	788	9	983	55	846	1	
					#		2		#						6	
16	21	955	0.02	0.130	###	0.070	0.00	1.273	###	788	9	943	54	834	1	
					#		2		#						6	
17	42	848	0.05	0.131	###	0.072	0.00	1.310	 ###	793	9	981	56	850	1	
- /		010	0.00	0.101	#	0.072	2	1.510	#	.,,,	Ĺ	,,,,			7	
18	15	821	0.02	0.133	" ###	0.067	0.00	1 245	" ###	806	#	835	61	821	2	
10	15	021	0.02	0.155	$\pi\pi\pi$	0.007	0.00	1.273	aππ	000	π'	055	01	021	4	I

Table 5 LA–ICP–MS analysis results for zircons from the Mather UHT granulites. 1309

1		1	1		I		T	1	1			1	1		1		
						#		2		#		#				0	
	19	12	675	0.02	0.133	###	0.068	0.00	1.257	###	807	9	861	72	827	1	
						#		2		#						9	
	20	43	666	0.06	0.133	###	0.073	0.00	1.376	###	808	#	1013	70	879	2	
						#		3		#		#				7	
	21	14	####	0.01	0.136	###	0.071	0.00	1.345	###	822	8	955	49	865	1	
						#		2		#						5	
	22	22	825	0.03	0.138	###	0.076	0.00	1.448	###	836	7	1087	55	909	1	
						#		2		#					\sim	7	
	23	10	548	0.02	0.139	###	0.073	0.00	1.421	###	837	#	1020	54	898	2	
						#		2		#		#)	0	
	24	21	765	0.03	0.139	###	0.072	0.00	1.384	###	838	#	977	61	882	1	
						#		2		#		#				7	
	25	24	####	0.02	0.145	###	0.069	0.00	1.391	###	873	#	889	57	885	1	
						#		2		#		#				8	
	26	47	758	0.06	0.148	###	0.073	0.00	1.552	###	889	#	1013	67	951	3	
						#		2		#		#				0	
	27	10	####	0.01	0.151	###	0.069	0.00	1.456	###	908	9	898	56	912	1	
						#		2		#	7					5	
	28	20	####	0.01	0.152	###	0.076	0.00	1.654	###	915	#	1098	57	991	2	
						#		2	\sim	#		#				6	
	29	23	944	0.02	0.157	###	0.078	0.00	1.708	###	940	#	1139	57	1012	2	
						#		2		#		#				1	
	30	26	972	0.03	0.160	###	0.081	0.00	1.820	###	954	#	1232	56	1053	2	
						#		2		#		#				0	
	31	27	805	0.03	0.161	### <	0.078	0.00	1.783	###	964	#	1158	55	1039	2	
						#		2		#		#				2	
	32	19	707	0.03	0.167	###	0.078	0.00	1.924	###	994	#	1154	64	1090	4	
						#		3		#		#				0	
	33	21	####	0.02	0.177	###	0.084	0.00	2.069	###	1050	7	1292	40	1139	1	
						#		2		#						4	
	Core			\sum	Y .												
	34	###	529	0.28	0.330	###	0.129	0.00	6.343	###	1840	#	2088	56	2024	5	
				7		#		4		#		#				3	
	35	###	219	0.79	0.393	###	0.150	0.00	8.258	###	2135	#	2340	53	2260	3	
						#		5		#		#				8	
	36	###	351	0.69	0.394	###	0.151	0.00	8.407	###	2140	#	2361	48	2276	3	
						#		4		#		#				3	
	37	58	289	0.20	0.367	###	0.152	0.00	7.803	###	2017	#	2374	41	2209	2	
						#		4		#		#				2	
	38	###	144	0.70	0.382	###	0.153	0.00	8.080	###	2085	#	2381	44	2240	2	
						#		4		#		#				4	
	39	###	167	0.83	0.453	###	0.154	0.00	9.704	###	2410	#	2387	57	2407	3	

					#		5		#		#				1	
40	###	378	0.57	0.371	###	0.154	0.00	7.970	###	2035	#	2391	39	2228	2	
					#		4		#		#				1	
41	80	140	0.57	0.441	###	0.155	0.00	9.582	###	2355	#	2406	54	2395	3	
					#		5		#		#				0	
42	###	441	1.81	0.391	###	0.160	0.00	8.661	###	2126	#	2452	46	2303	2	
					#		4		#		#				5	
43	###	337	0.69	0.446	###	0.160	0.00	9.951	###	2376	#	2461	41	2430	2	
					#		4		#		#			\sim	3	
44	50	49	1.01	0.436	###	0.161	0.00	9.656	###	2331	#	2465	58	2403	3	
					#		6		#		#)	2	
45	###	234	0.59	0.448	###	0.168	0.00	10.468	###	2384	#	2540	38	2477	2	
					#		4		#		#	$\boldsymbol{\wedge}$			4	
46	###	227	0.52	0.458	###	0.177	0.00	11.225	###	2429	#	2623	39	2542	2	
					#		4		#		#				3	

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1329 FIGURE CAPTIONS

1330

Figure 1 (a) Simplified geological map of the Rauer Islands, East Antarctica (after Harley &
Fitzsimons, 1991; Tong & Wilson, 2006; Tong et al., 2021). The black dashed line represents the
boundary between Mesoproterozoic and Archean crustal components. (b) Location of the Rauer
Islands in the reconstructed Gondwana supercontinent at ~500 Ma (modified after Fitzsimons, 2000;
Wilson et al., 2007; Hokada et al., 2016). (c) Lithological distribution and sample locality on the
Mather Peninsula (after Harley, 1998b; Hokada et al., 2016).

1337

Figure 2 (a) A Field photograph showing the occurrence of *UHT* granulites from the Mather Peninsula, East Antarctica. (b) and (c) Enlarged photographs of white boxes in (a) displaying the presence of different lithological components. The sampling sites are shown by orange stars. The hammer for scale is 25 cm in length. (d) A representative hand specimen of the studied rocks. The coin for scale is about 2.4 cm in diameter.

1343

Figure 3 Plane polarized light photomicrographs and BSE images of the Mather UHT granulite 1344 sample RG16-80. (a) Co-existence of garnet and orthopyroxene porphyroblasts. High-Al 1345 orthopyroxene inclusion within garnet is outlined by green dashed lines in the figure, which 1346 coexists with primary NIs (outlined by red dashed lines). Also shown in this figure are secondary 1347 FIs, highlighted with blue dashed lines. (b) Coexistence of biotite and orthopyroxene 1348 porphyroblasts, with a biotite inclusion within orthopyroxene. (c) K-feldspar grain rimmed by the 1349 intergrowth of orthopyroxene + cordierite + plagioclase. (d) Garnet and orthopyroxene 1350 porphyroblasts separated by fine-grained intergrowths of orthopyroxene + cordierite. (e) A BSE 1351 1352 image showing the intergrowth of orthopyroxene + cordierite + plagioclase around garnets. (f) Intergrowths of sapphirine + cordierite in the matrix. (g) BSE image showing vermicular low-Al 1353 orthopyroxene in the rim of a high-Al orthopyroxene porphyroblast. (h) Biotite + plagioclase 1354 symplectites surrounding orthopyroxene. (i) Elongated garnet, cordierite, biotite, plagioclase in 1355 strongly deformed domains. Interstitial plagioclase is outlined by dashed black lines while quartz 1356 inclusions within garnets are highlighted with white dashed lines. 1357

Figure 4 (a) Garnet grain showing the analysis positions. (b) Compositional zoning profile across
garnet from core to rim. (c) Chondrite–normalized REE patterns of garnets for the NI–bearing and
NI–absent domains. (d) Compositional zoning profiles across an orthopyroxene porphyroblast.

Figure 5 Plane polarized light photomicrographs and BSE images of NIs within garnet porphyroblasts and Raman spectrum of these inclusions. (a) Garnet crystal mounted on an epoxy resin disc containing abundant NIs. (b) and (c) Plane polarized photomicrographs showing NIs within peritectic garnets. (d)–(f) BSE images showing NIs with daughter phases of quartz, biotite and kokchetavite. Some cavities are also observed and might be related to the exsolution of fluid during the retrograde crystallization of melt. (g) Mixed Raman spectrum of NIs and their garnet hosts.

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Figure 6 A BSE image of a nanogranitoid inclusion and corresponding X–ray maps of Si, Al, Mg,K and Na.

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Figure 7 (a)–(c) Secondary FIs within garnet and orthopyroxene porphyroblasts. Pink and green arrows respectively point to nanogranitoid and fluid inclusions, where NIs are randomly distributed while FIs are mostly present along the cracks. (d–f) Mixed spectra of FIs and their hosts.

1377

Figure 8 BSE images of experimentally re-homogenized NIs under 1000 °C and 12 kbar with a run duration of 6 hours (a), 950 °C and 12 kbar with run durations of 4 hours (b, d), 2 hours (c, e), and 1 hour (f). Perfectly re-homogenized inclusions with well-developed negative shapes are shown in Figures e and f. Other inclusions show clear evidence for overheating (a–d).

- **Figure 9** Plot of aluminum saturation index vs. alkalinity index $[AI = molar Al_2O_3 (Na_2O + K_2O)$ (from Frost & Frost, 2008). Dotted line separates the compositional fields of I– and S–type granites (from Chappell, 1999).
- 1386 KKB, Kerala Khondalite Belt (India) (Cesare et al., 2009; Ferrero et al., 2012), with peak P-T1387 conditions of 900°C and 6–8 kbar (Cenki et al., 2002; Shabeer et al., 2005); Grf Type–A and –C,
- 1388 type–A or –C granulites from the Gruf Complex (European Central Alps) (Gianola et al., 2021),

with peak P-T conditions of 920–940 °C and 8.5–9.5 kbar (Galli et al., 2012); Iv, metapelitic migmatites from the Ivrea Zone (NW Italy) (Carvalho et al., 2019), with P-T conditions of 650–730 °C and 3.5–6.5 kbar for the UA, and 900–950 °C and 11–12 kbar for the G (Redler et al., 2013; Ewing et al., 2013). UA, upper amphibolite facies zone; Tr, transition zone; G, granulite facies zone.

1394

Figure 10 (a) CIPW Qz–Ab–Or diagram showing normative compositions of analyzed NIs (normalized to 100% on anhydrous basis). Pink dashed lines show the eutectic points and cotectic lines for the subaluminous haplogranite system at $a_{H2O} = 1$ and different pressures (Luth, et al., 1964). References to experimental glasses: PDJ91 (shown in green), 950 °C and 7 kbar (Patiño Douce & Johnston 1991); D03 (shown in blue), 900 °C and 5 kbar (Droop et al., 2003). (b) An–Or– Ab diagram showing the normative compositions of analyzed NIs (after O'Connor, 1965).

- 1401 For the symbols, please refer to Figure 9.
- 1402

Figure 11 Trace element compositions of NIs and granulitic restite, normalized to the average upper continental crust (UCC) composition (data from Rudnick & Gao 2014). Trace elements of matrix glasses within metapelitic enclaves from El Hoyazo (SE Spain) (samples HO–50A, –33A and –54; Acosta–Vigil et al., 2010) are also shown for comparison. Note that peak P-T conditions for the metapelitic enclaves from El Hoyazo are 850 ± 50 °C and 5–7 kbar, slightly lower than the investigated rocks in this study.

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1414

Figure 12 (a) CL images of representative zircons mounted on epoxy resins. Green circles represent analyzed spots and the nearby blue numbers represent their isotopic dates. (b)–(d) Photographs showing inclusions within zircons. (e) Concordia diagram showing the results of U–Pb zircon dating. (f) Enlarged concordia diagram of the box in Figure e.

Figure 13 (a) *P*–*T* pseudosection calculated at $X(Fe_2O_3) = Fe^{3+}/Fe^T = 0.35$, with a *P*–*T* range of 3–15 kbar and 750–1200 °C. The fluid–absent solidus is shown in red dashed curves and the interpreted peak *P*–*T* conditions are shown in pale yellow. The grey curves in Figure a represent the inferred *P*–*T* path. (b) Enlarged *P*–*T* pseudosection with a *P*–*T* range of 5–9 kbar and 800–1000 °C. 1419 The constrained P-T fields for the M2 and M3 stage are outlined by purple and black bold lines, 1420 respectively.

Figure 14 (a) $T-X_{melt}$ pseudosection showing the pre-peak evolution of the investigated rocks. Average re-homogenized nanogranitoid compositions are adopted as the melt compositions. The pressure is fixed at ~12.5 kbar. (b) Reconstructed P-T pseudosection after 25% melt reintegration into the bulk rock composition.

1427Figure 15 Suggested P-T path of UHT granulites from the Mather Peninsula, East Antarctica. Also1428shown are previously retrieved P-T paths for the UHT granulites from the Rauer Islands (modified1429after Tong et al, 2021).

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1	4	3	9

1449 Fig. 1.



1452 **Fig. 2.**



1467 **Fig. 3.**







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Fig. 6. 1508



1528 Fig. 7.
















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