

Evidence for Low-pressure Crustal Anatexis During the Northeast Atlantic Break-up

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38 **Key Points:**

- 39 • A dacitic unit was recovered in early Eocene sediments on the Vøring margin during
40 International Ocean Discovery Program Expedition 396.
- 41 • The dacite was formed by upper crustal anatexis at 54.6 ± 1.1 Ma, postdating the
42 Paleocene-Eocene Thermal Maximum.
- 43 • The dacite is evidence for a pre-breakup phase associated with significant continental
44 lithospheric extension.

45 Abstract

46 While basaltic volcanism is dominate during rifting and continental breakup, felsic magmatism
47 may also comprise important components of some rift margins. During International Ocean
48 Discovery Program (IODP) Expedition 396 on the continental margin of Norway, a graphite-
49 garnet-cordierite bearing dacitic, pyroclastic unit was recovered within early Eocene sediments
50 on Mimir High (Site U1570), a marginal high on the Vøring transform margin. Here, we present
51 a comprehensive textural, mineralogical, and petrological study of the dacite in order to assess its
52 melting origin and emplacement. The major mineral phases (garnet, cordierite, quartz,
53 plagioclase, alkali feldspar) are hosted in a fresh rhyolitic, highly vesicular, glassy matrix, locally
54 mingled with sediments. The xenocrystic major element chemistry of garnet and cordierite, the
55 presence of zircon inclusions with inherited cores, and thermobarometric calculations all support
56 a crustal metapelite origin. While most magma-rich margin models favor crustal anatexis in the
57 lower crust, thermobarometric calculations performed here show that the dacite was produced at
58 upper-crustal depths (< 5 kbar) and high temperature (750–800 °C) with up to 3 wt% water
59 content. In situ U-Pb analyses on zircon inclusions give a magmatic age of 54.6 ± 1.1 Ma,
60 revealing the emplacement of the dacite post-dates the Paleocene-Eocene Thermal Maximum
61 (PETM). Our results suggest that the opening of the North Atlantic was associated with a phase
62 of low-pressure, high-temperature crustal melting at the onset of the main phase of magmatism.

63

64 Plain Language Summary

65 Fifty-six million years ago, the continents were beginning the final phase of their journey to their
66 current configuration. This included the rifting and formation of the Northeast Atlantic Ocean,
67 known in particular for the excess amounts of magmatic activity that accompanied continental
68 breakup. The International Ocean Discovery Program organized Expedition 396 to collect
69 volcanic and sedimentary rocks deposited during this time off the coast of present-day Norway to
70 investigate the cause of the excess magmatism and its potential implications for the global
71 climate. While sampling sediments on the expedition, an unexpected volcanic unit, a garnet-
72 cordierite, glassy dacite, was recovered. To determine the origin and emplacement of this unit we
73 combined multiple methods (petrography, stratigraphy, thermodynamic calculations,
74 geochronology, in site compositional analyses) and showed that the unit was produced by

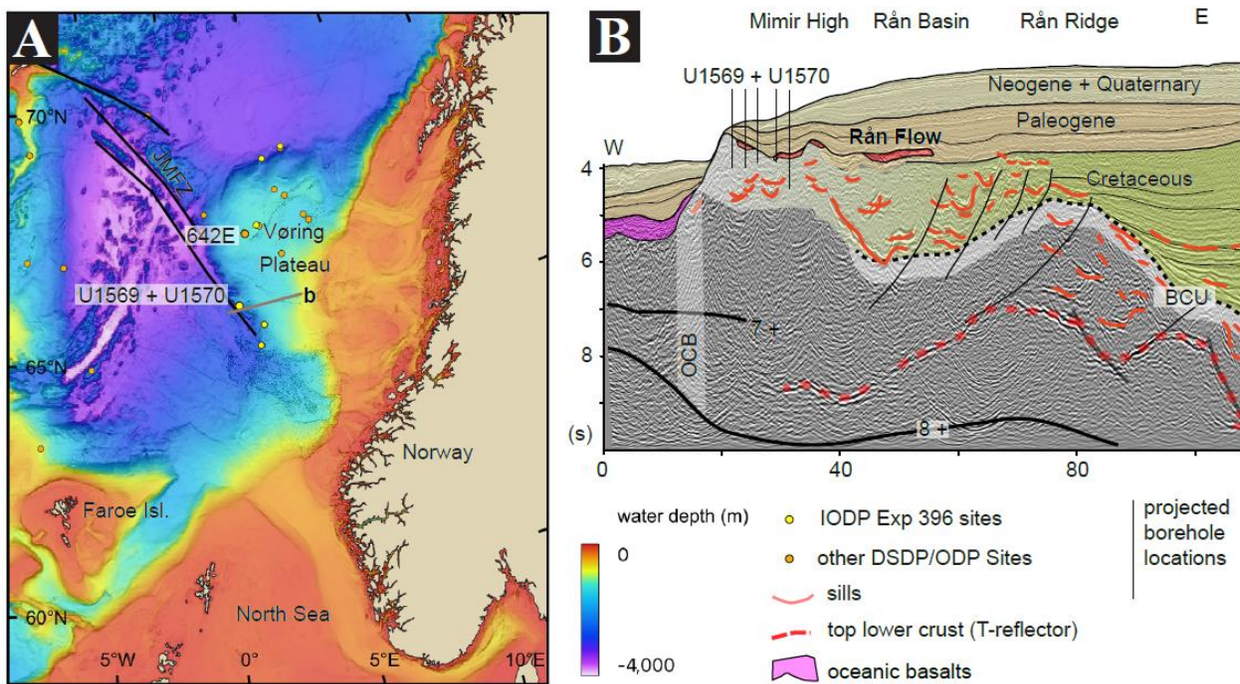
75 melting of the upper continental crust at depth during the rifting process and likely later
76 emplaced as a pyroclastic flow in shallow water. Our results demonstrate that the rifting process
77 in the Northeast Atlantic included a long and intense period of continental lithospheric thinning.
78 This research provides evidence needed to reconstruct the evolution of the North Atlantic Ocean
79 and associated magma production.

80 **1 Introduction**

81 The North Atlantic Igneous Province (NAIP) is an estimated $6\text{--}10\times 10^6$ km³ of magma
82 that was emplaced during the Paleogene (64–35 Ma; Storey et al., 2007). Much of this volume
83 was emplaced 56 to 54 million years ago (Ma), which coincides with the opening of the North
84 Atlantic, which began around 56 Ma (Saunders et al., 2007; Horni et al., 2017; Westerhold et al.
85 2020; Planke et al., 2023a). In the NE Atlantic, on the mid-Norwegian rift margin, NAIP
86 material is present as successions of voluminous extrusive basalt flows, including seaward
87 dipping reflectors (SDRs), magmatic intrusions within the continental crust and associated
88 sedimentary basins, and high-velocity bodies underlying the continent-ocean boundary (COB) at
89 the base of the continental crust which, at least in the distal margin, are interpreted as a magmatic
90 underplate (Fig. 1; Berndt et al., 2001; Mjelde et al., 2005; Planke et al. 2005; Planke et al.,
91 2023a; Svensen et al., 2004). Passive rift margins whose volume of magmatic production and
92 volcanic activity cannot be explained by decompression melting of the sub-lithosphere at
93 ambient mantle temperature, like the mid-Norwegian margin, are classified as excess magmatic
94 margins (Lu & Huisman, 2001), or volcanic rifted margins.

95 Excess magmatism at volcanic margins was first attributed to active rifting (as opposed to
96 passive rifting in non-volcanic margins; Celal Sengör & Burke, 1978). In a purely active model,
97 a hot mantle (related, for example, to the presence of a mantle plume) thermally erodes the
98 bottom of the lithosphere (Fleitout et al., 1896) and results in two stages of rifting: (1) flood
99 basalt stage, coeval with very small crustal extension, and (2) a break-up stage associated with
100 the volcanic margin formation (Courtilot et al., 1999). During active rifting, regional uplift
101 predates and accompanies the volcanic margin development (e.g. Clift et al., 1998; White &
102 Lovell, 1997). However, the purely active upwelling model is limited by evidence that many
103 volcanic rifted margins are associated with regions that have been previously subjected to far-
104 field extension (White & McKensie, 1989). This relationship suggests that prior extension may

105 facilitate volcanic activity and that deep mantle plumes may not be required to explain all cases
 106 of excess magmatism (e.g. Hill, 1991; White & McKensie, 1989). However, it is important to
 107 highlight that volcanic margins can also develop in areas that were not subjected to significant
 108 lithospheric stretching leading up to the flood basalt stage (e.g. in Afar; Courtillot et al. 1999).
 109 While accurate timing of the initiation and end of continental rifting, of the peak of magmatic
 110 activity, and of the main continental break-up phase is crucial to refine models of margin
 111 development, the causality and timing between rifting and excess magmatism is still poorly
 112 constrained.



113 **Figure 1.** (a) Bathymetry map showing relative locations for IODP Expedition 396 (yellow
 114 circles), and for other DSDP/ODP sites (orange circles). The brown line shows the location of
 115 the seismic profile in (b). (b) Interpreted seismic section along Mimir High and the Rån Basin.
 116 The 7+ km/s and T-reflection (TR) are interpreted as the top of the lower crustal body (LCB), the
 117 8+ km/s as the seismic Moho, and BCU, this Base Cretaceous Unconformity. Adapted from
 118 Abdelmalak et al. (2017).
 119

120 During the recent International Ocean Discovery Program (IODP) Expedition 396, a
 121 dacitic unit was recovered in lower Eocene sediments deposited on the Mimir High (Fig. 1A;
 122 Planke et al., 2023b). Mimir High is a marginal high adjacent to the Vøring transform margin,
 123 the landward extension of the Jan Mayen Fracture Zone that segments the mid-Norwegian rift

124 margin. Dacite flows in oceanic or transitional regimes outside of volcanic arcs are unusual but
125 not unique. For instance, they have been collected at several locations in the NAIP, including in
126 Well 163/6-1A in the northern Rockall Trough (Morton et al., 1988a), in Well 209/3-1 in the
127 Faroe-Shetland Basin (Kanaris-Sotiriou et al. 1993), in Hole 642E on the Vøring Plateau from
128 Ocean Drilling Program (ODP) Leg 104 (Fig. 1; Eldholm et al., 1987; Viereck et al., 1988), and
129 on the southeast Greenland margin (Larsen et al., 1995). Garnet and cordierite-bearing dacitic
130 flows have also been reported in the Neogene Volcanic Province (NVP) in SE Spain that are
131 associated with the opening of the Alborán Domain in the late Tertiary (Platt et al., 1998; Vissers
132 et al., 1995). Dacitic flows have previously been interpreted as being products of crustal melting
133 at various depths, with the units recovered in the NE Atlantic interpreted as produced in shallow
134 magma chambers (Eldholm et al., 1989) and the units from SE Greenland as derived from
135 magma differentiation and crustal contamination in the lower crust (e.g., Fitton et al., 1995).

136 Dacites previously collected in the NE Atlantic are peraluminous, suggesting they were
137 generated through the melting or assimilation of pelitic sediments or metapelitic rocks. However,
138 previously recovered units have experienced varying degrees of alteration (e.g., Abdelmalak et
139 al., 2016; Morton et al., 1988) and, to our knowledge, no thermobarometric calculations were
140 performed to constrain the origin of these rocks. Here, we use petrographic analyses, major and
141 trace element compositions of mineral phases, in-situ U-Pb zircon dating, and thermodynamic
142 modeling to discuss the petrogenesis and emplacement of the dacite in the context of the
143 emplacement of the NAIP and explore the geodynamic implications for the development of the
144 Vøring margin.

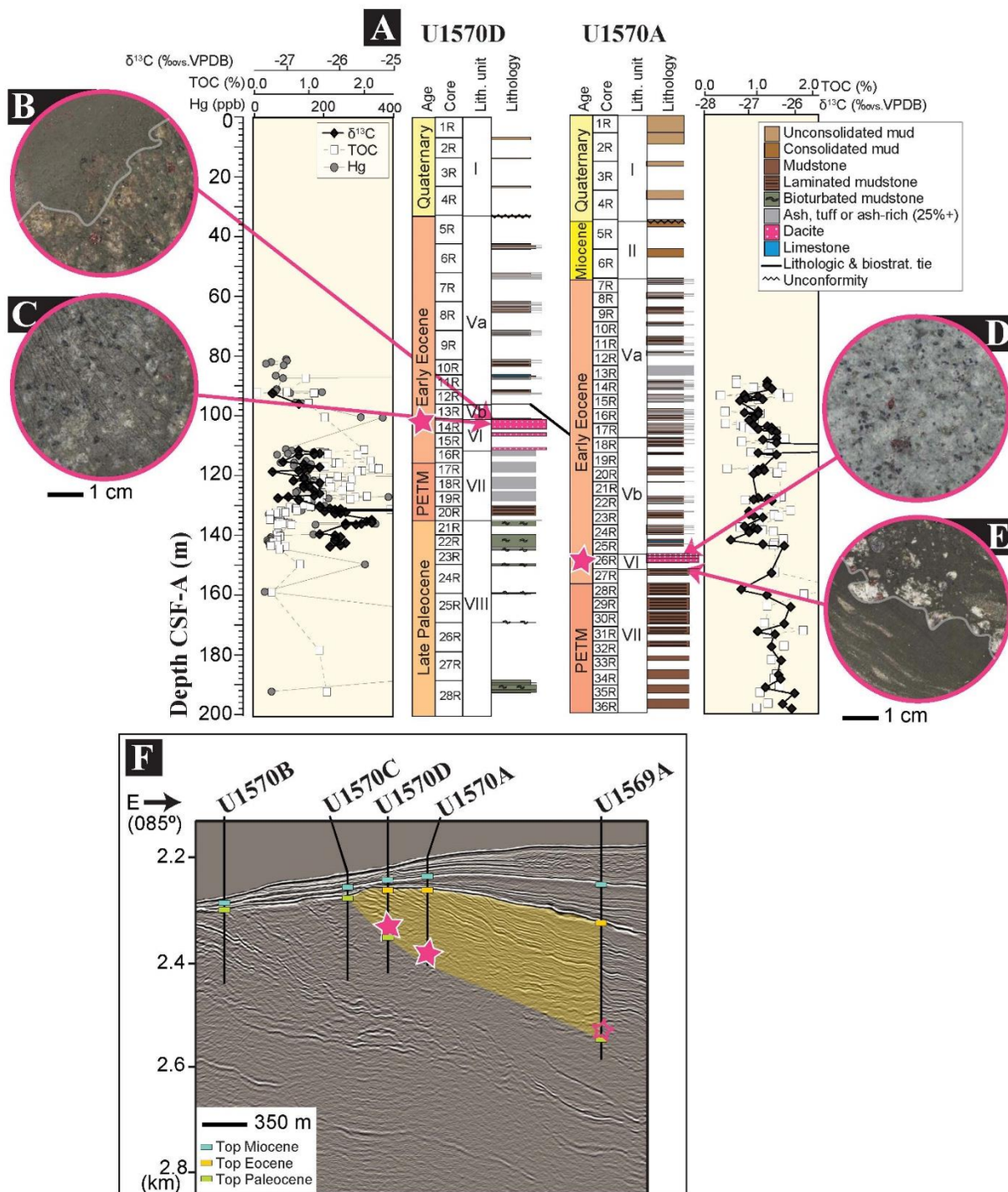
145 1.1 Geologic setting

146 Mimir High (Fig. 1) is a marginal high on the continent side of the continent-ocean
147 boundary of the Vøring transform margin. This margin is characterized by diminished extrusive
148 volcanism relative to the adjacent rifted margin segments of the Vøring and Møre basins (Berndt
149 et al., 2001). The absence of basalt cover makes the Mimir High an exceptional place to study
150 the stratigraphic successions and effects of sill intrusions (Planke et al., 2023). In fact, seismic
151 interpretation suggests that the Mimir High was a depocenter during the Eocene, which makes it
152 an ideal candidate for recovering expanded Paleocene–Eocene sedimentary sections. During
153 Expedition 396, a transect of five boreholes was drilled along the Mimir High with the intention

154 of forming a composite ribbon core through most the Paleocene and Eocene strata (Fig. 2). In
155 two of these boreholes (U1570A and D), the early Eocene sediments hosted a discrete (<10 m
156 thick) glassy, garnet-cordierite-graphite dacitic igneous unit (Planke et al., 2023).

157 The dacite was recovered from between 146.84 and 151.96 meters below surface (mbsf)
158 in Hole U1570A and between 100.75 and 110.91 mbsf in Hole U1570D (Fig. 2). The contact
159 between the base of the dacite and dark gray and grayish-brown ash-rich siltstone was recovered
160 in the cores from both boreholes. This basal contact is 8 and 9 m above the Paleocene–Eocene
161 Thermal Maximum (PETM) interval in holes U1570D and U1570A, respectively (Vickers et al.,
162 2023). Within both holes, the dacitic unit is observed as semi-discrete layers with local peperitic
163 texture and mingling with dark, sand-rich silt with small (2–5 cm) intervals of silt and claystone
164 between them (Fig. 2C). It is overlain by an early Eocene (Ypresian) age dark gray claystone
165 with black ash and intervals of gray limestone. The top contact between the dacite and the early
166 Eocene sediments was also recovered in Hole D (Fig. 2B). While this unit was not encountered
167 in Hole U1569A, millimeter-size garnet grains were recovered in the core catcher of Section
168 U1569A–35R collected at 315.66 mbsf. The location of this section is indicated by an open pink
169 star in Figure 2F.

170



171

172 **Figure 2.** (a) Stratigraphic columns of the boreholes U1570D and U1570A cores modified from
 173 Planke et al. (2023b) and associated TOC, Hg concentrations, and $\delta^{13}\text{C}_{\text{org}}$ measurements in the
 174 sediments. The dacitic unit is represented in pink with white dots. Large circles show various
 175 facies of the dacitic unit at the locations indicated by arrows in the stratigraphic column. (b)

176 Contact between the layered sediments and the dacite (sample 1570D-14R1-11-18, read as
177 borehole U1570D, core 14, section 1 between 11 and 18 cm), (c) peperitic texture (sample
178 1570D-14R2-7-11), (d) a fresh, unmingled section of dacite (sample 1570A-26R2-51-54), and
179 (e) basalt contact between the dacite and early Eocene sediment below (highlighted with a gray
180 line; sample 1570A-27R1-9-13). The 1cm scale bar refers to all images of the core. (f) The
181 interpreted seismic section at sites U1570 and U1569, adapted from Planke et al. (2023),
182 showing the spatial relationship between the boreholes, geological epochs, and the locations of
183 dacite recovery (pink stars); smaller open pink star is the recovery location of individual garnet
184 grains at site U1569.

185 **2 Methods**

186 2.1 Bulk rock geochemical analyses

187 We analyzed two samples for major and trace element bulk rock concentrations; a
188 representative fresh sample with little to no mixing with sediments (U1570A-26R2-51-54 cm;
189 Fig. 2D) and a sample collected for being representative of mingling textures with the sediments
190 (U1570D-14R2-7-11 cm; Fig. 2C). Sample U1570A-26R2-51-54 major element composition
191 was done using Thermo Scientific ARL X-ray fluorescence (XRF) spectrometer at Institute of
192 Geochemistry, Chinese Academy of Sciences. BHVO-2 and BCR-2 were used as secondary
193 standards. Trace element concentrations were acquired using the Agilent-7900 inductively
194 coupled plasma mass spectrometer (ICP-MS) at Institute of Oceanography, Chinese Academy of
195 Sciences, following procedures of Chen et al. (2017). In brief, ~50 milligrams of the sample
196 were dissolved with an acid mix of double distilled, concentrated HCl+HNO₃ and HF in a high-
197 pressure bomb for 15 hours and then re-dissolved with distilled 20% HNO₃ for 2 hours until
198 complete digestion/dissolution. BHVO-2 was used as a secondary standard. Analytical
199 accuracy on secondary standards is better than 5% for all major elements and most trace
200 elements, between 5% and 10% for Li, Cr, Mn, and Cu, and better than 15% for Be, Ti, V, Zn
201 and Ta.

202 Sample U1570D-14R2-7-11 major element composition was analyzed using X-ray
203 fluorescence spectrometry (Rigaku RIX3000) at Niigata University following the analytical
204 method of Takahashi and Shuto (1997), with an optimization for ultramafic rocks. Trace element

205 concentrations were determined using a Yokogawa HP4500 ICP-MS at Niigata University,
206 following the acid digestion method described in Senda et al. (2014). In brief, for each sample,
207 100 mg of material was weighed and placed in a Teflon vial and underwent step-by-step acid
208 treatment with heating until the sample reached complete dissolution. The solution was finally
209 diluted using HNO₃ and an internal standard (¹¹⁵In) and was measured along with secondary
210 standards (BHVO-2, W2a and JB2). Analytical accuracy on secondary standards is better than
211 5% for most elements, between 5 and 10% for Sc, Ga, Sr, Nb, Gd, Dy, Yb, Lu, Hf, and Pb, and
212 12% for Cs.

213 2.2 Sample selection for in situ analysis

214 Fourteen samples of the dacite unit were collected from both Holes U1570A and
215 U1570D. Materials were sampled to include a range of apparent volcanic facies (undisturbed
216 versus higher degrees of sediment mingling). Four thin sections (30 μm thick) were impregnated
217 with blue epoxy to highlight the porosity. The remaining ten samples were cut, polished, and
218 carbon-coated into thick sections (300–1000 μm thick) to minimize potential issues with the
219 fragility of the sample and to facilitate laser ablation analyses.

220 2.3 Scanning electron microscopy and electron microprobe analysis

221 Scanning electron microscope (SEM) images presented in this study were collected using
222 a JEOL JSM-IT300 at the Energy and Geoscience Institute (EGI) at the University of Utah.
223 Major and minor element compositions were acquired using a Cameca SX100 electron
224 microprobe at the University of Utah. A 15 keV beam voltage and 20 nA beam current were used
225 for all analyses. Single spot measurements were used for mineral phases and glass using beam
226 diameters of 5 μm and 10 μm, respectively, and 1–3 spots were analyzed on each large mineral
227 phase. Sodium (Na) and potassium (K) were analyzed first, and a time-dependent intensity (TDI)
228 correction was applied to correct the signal from devolatilization (Nielson & Sigurdsson, 1981;
229 Siivola, 1969). Compositional profiles were also acquired across large (1–4 mm) garnet grains
230 using a 5 μm beam diameter every ~120 μm (individual analysis locations within these profiles
231 were adjusted to avoid cracks and inclusions). On-peak counting times were 10 s for major
232 elements and 20 s for minor elements; half of the on-peak time was used on each of the high and
233 low backgrounds. The following ASTIMEX reference standards were used, Rutile (Ti),

234 Chromite (Cr), Diopside (Ca), Albite (Al, Na, Si), Sanidine (K), Apatite (P), and Sphalerite (S).
235 Additionally, we used synthetic Mn and a Ni olivine provided by George Rossman to calibrate
236 Mn and Ni, and the USNM San Carlos olivine distributed by the Smithsonian Institute (USNM
237 111312/44) to calibrate Fe and Mg. Data were processed using the ‘Probe for EMPA (v. 12.7.3)’
238 software and the PAP procedure was used for data reduction (Pouchou & Pichoir, 1991). BHVO-
239 2g and USNM San Carlos olivine 111312/44 were repeatedly analyzed as secondary standards
240 during each analytical session to monitor drift on major and minor elements, but no additional
241 corrections were applied.

242 2.4 Laser ablation inductively coupled plasma mass spectrometry

243 2.4.1 Trace element analysis of garnet, cordierite, and feldspars

244 Trace element concentrations in garnet, plagioclase, alkali feldspar, and cordierite were
245 collected using laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS).
246 Analyses were performed at the University of Utah on a Teledyne-Photon Machines Analyte
247 Excite Excimer LA system attached to an Agilent 8900 ICP-MS. The laser carrier and the
248 nebulizer gas flow were 1 L He/min (0.4 L/min cup +0.6 L/min cell and 1.1 L Ar/min,
249 respectively). The first half of the analytical session was run using a fluence of 2.19 J/cm², which
250 was later increased to 3.29 J/cm² in an attempt to reduce shattering observed on quartz grains
251 (see below). We used a laser repetition rate of 10 Hz. We used 40 μm spot analyses, with each
252 analysis preceded by a cleaning shot. Acquisition times were about 50 s. To assess compositional
253 variability, 1–5 analyses were performed per grain. NIST610 and NIST612 were measured every
254 six unknown analyses for a (total n = 23) and used as primary standards for elements with
255 concentrations higher and lower than 100 ppm, respectively. The Si contents, measured by
256 electron microprobe, served as an internal standard. Each element signal was carefully monitored
257 for spikes that may indicate the presence of cracks or inclusions; in this case, the analysis was
258 discarded. We tested the accuracy of each measurement with BHVO-2G, which was acquired
259 every 12 unknown analyses for a total of 10 analyses. Measured concentrations for BHVO-2G
260 are compared with the GeoReM preferred values (Jochum et al. 2005); the relative errors on each
261 trace element contents are reported in Table S1.

262 *2.4.2 Trace element analysis of quartz*

263 For quartz grains, we performed an additional analytical session with a fluence of 7.90
264 J/cm², a 5 Hz laser repetition rate, and pre-ablation cleaning spots for 85 µm single spot analyses.
265 These conditions minimized the number of quartz grains shattered by the laser, providing 51
266 quartz analysis from 17 individual grains with good analytical signals. Elements analyzed in
267 quartz are Ti, Li, Na, Al, P, K, Ge, Rb, Sr, Zr, and Ba. NIST610 was used as the primary
268 reference material, measured once every five to eight quartz analyses for a total of 21
269 measurements. The data were processed by normalizing analytical peaks from the mass
270 spectrometer by their ²⁸Si values and converted to ppm using the NIST610 standard values as
271 calibration. NIST612 was used as a secondary standard. Comparison of the mean concentrations
272 from 11 NIST612 analyses with the GeoReM preferred values (Jochum et al. 2011) shows that
273 the relative error on the trace element contents in quartz is 6.7% for Ti and lies between 1.0 and
274 12.1% for the other analyzed elements (Table S1). We also performed six analyses on the
275 reference material San Carlos Olivine NMNH 111312/42 and compared the data with data
276 published in the literature (Lambart et al., 2022). The measured Ti concentration in olivine San
277 Carlos reference material overlaps with the published values inside the two standard deviation
278 (Table S1).

279 *2.4.3 U-Pb zircon analyses*

280 Twenty-five zircon crystals across two samples were located and imaged using
281 backscattered electron and cathodoluminescence (CL) detectors. The BSE and CL images (Fig.
282 S1) guided laser spot placements to minimize any mechanical mixing of chemically and/or
283 isotopically distinct domains and to ensure an analysis of the full range of populations in each
284 sample. Both U-Pb and trace element analyses were performed simultaneously for the two
285 samples by Laser Ablation Split Stream (LASS; Kylander-Clark et al., 2013; Stearns et al., 2020)
286 at the Petrochronology lab of the University of California Santa Barbara. The crystals were
287 ablated using a Photon Machines/Teledyne 193 nm Excimer laser equipped with a two-volume
288 Helex® stage (Eggins et al., 1998) and the ablated material was introduced via He carrier gas to
289 the multi-collector ICP-MS and quadrupole (Q-MS) mass spectrometers to measure uranium,
290 thorium, and lead isotopes and trace elements, respectively. The Nu Plasma multicollector
291 measured ²³⁸U, ²³⁵U, ²³²Th, ²⁰⁸Pb, ²⁰⁷Pb, ²⁰⁶Pb, ²⁰⁴Pb, ²⁰²Hg and the Agilent 7700X quadrupole

292 measured Zr, Th, U, Si, P, Ti, V, Sr, Y, Nb, Mo, La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy, Ho, Er,
293 Tm, Yb, Lu, Hf, Ta, and W.

294 A total of 88 unknown analyses on the U1570 dacite's zircons were collected using a 12
295 μm spot size, a 5Hz laser operating frequency, and pre-ablation cleaning shots. A standard
296 sample bracketing method was used, and a series of primary and secondary reference materials
297 were measured, including GJ (608.5 Ma; Ehlou et al., 2006) and NIST612 as primary standards,
298 and 91500 (Weidenbeck et al., 2004), Aus (Kennedy et al., 2014), and Plešovice (337.13 Ma;
299 Sláma et al., 2008) as secondary standards. Comparison between the mean measured values and
300 the published values for trace elements are reported in Table S1.

301 2.5 MAGEMin thermodynamic calculations

302 The Mineral Assemblage Gibbs Energy Minimization (MAGEMin) thermodynamic
303 package (Riel et al. 2022) was used to calculate stability fields for minerals and their modal
304 proportions using the bulk composition of the dacite sample U1570-26R2-51-54 cm (Fig. 2D, 3)
305 for comparison with the phase assemblage observed in the samples. The MAGEMin program is
306 calibrated with the Holland et al. (2018) thermodynamic dataset and KNCFMASHTOCr
307 calculation system to perform single-point calculations at a given pressure, temperature, and
308 bulk-rock composition to find the most thermodynamically stable phase assemblage.
309 Pseudosections were computed using seven refinement levels and six cores for 0 to 6 kbar (1
310 kbar step size) and for 600 to 1000 °C (100 °C step size). Three pseudosections were calculated
311 to constrain the effect of water, using inputs of 1 wt%, 3 wt%, and 5 wt% H₂O.

312 2.6 Carbon isotopes and mercury concentration in sediments

313 We determined the total organic carbon (TOC) contents and stable carbon isotope ratios
314 ($\delta^{13}\text{C}_{\text{org}}$) of sediments around the dacitic unit by powdering and decalcifying samples using 1 M
315 HCl for 72 hours. A brief (c. 1 h) heating step at 50 °C during the acidification process was
316 included to remove any siderite that is occasionally present in these sediments. The samples were
317 then rinsed using deionized water, oven-dried at 50°C, and re-homogenized. We performed
318 analyses of a reconnaissance data set using a Thermo Fisher Scientific Flash Elemental Analyzer
319 coupled to a Thermo Fisher Scientific DeltaV Isotope Ratio Mass Spectrometer at the CLIPT
320 Lab, University of Oslo. Based on these initial results, we analyzed an expanded sample set for

321 TOC and $\delta^{13}\text{C}_{\text{org}}$ using a Thermo Finnigan DeltaPlus XP Mass Spectrometer at the School of
322 Ocean and Earth Science and Technology (SOEST) at the University of Hawaii at Mānoa.

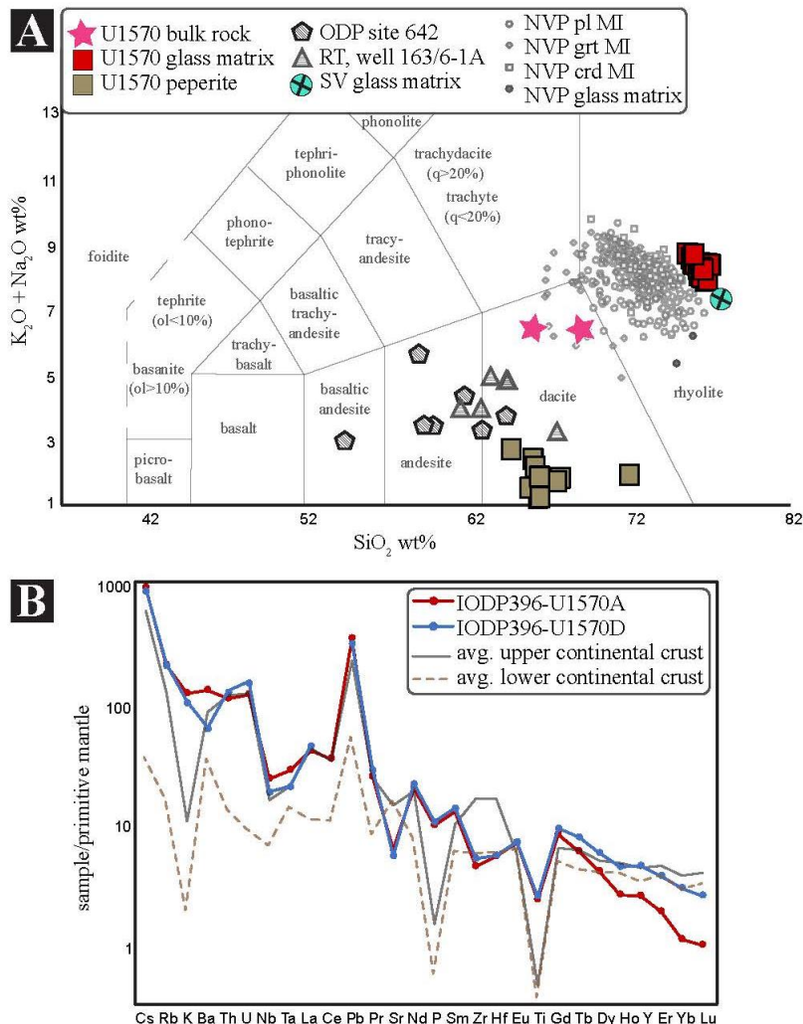
323 We performed mercury analyses at the University of Oxford (UK) using a Lumex
324 915Lab and a Lumex 915+ Portable Mercury Analyzer with a PYRO-915 pyrolyzer attached.
325 Approximately 100 mg of oven-dried (40 °C) and finely powdered sample was heated to >700
326 °C to volatilize the Hg present and Hg concentrations were determined via atomic absorption
327 spectrometry. The instruments were calibrated using the NIST-SRM2587 (paint-contaminated
328 soil) standard (290 ± 9 ppb) and long-term observations of the standards and samples showed
329 that analytical errors for the Hg analyzers were $\pm 6\%$ (Frieling et al., 2023).

330 Total inorganic (TIC) and organic carbon (TOC) concentrations and thermal maturity of
331 organic-carbon on the same sample powders as used for Hg were analyzed at the University of
332 Oxford (UK) with a Vinci RockEval-6 device using standard procedures (Behar et al., 2001). For
333 each sample, ~50 mg of dried powdered material was analyzed, and we assessed reproducibility
334 with an in-house standard of homogenized sediment (~2.8% TOC, ~6% TIC). The standard
335 deviation for TOC, TIC, hydrogen index and Tmax was $\pm 1\%$ of the measured value or better for
336 the in-house standard ($n = 6$). The hydrogen index (HI) and oxygen index (OI) are defined as in
337 Behar et al. (2001) where the mass of released hydrocarbons (“S2”) and CO₂ (“S3”) in mg is
338 multiplied by 100 and divided by TOC to obtain HI and OI, respectively.

339 **3 Results**

340 **3.1 Bulk rock composition**

341 Major and trace element bulk rock compositions are reported in Table S2. Compositions
342 obtained on both samples show similar major element compositions (Fig. 3A). The igneous
343 samples from the two boreholes have a SiO₂ content of 66–70 wt%, an alkali (Na₂O+K₂O)
344 content of 6.3 wt%, and an A/CNK (molar Al₂O₃/(CaO+Na₂O+K₂O)) aluminum saturation value
345 of 1.5–1.6, classifying them as peraluminous dacite. Rare earth element (REE) diagrams from
346 both samples (Fig. S2) also show similar enrichment in light-REE (LREE), with La_N/Sm_N ~3.1,
347 and a moderate negative Eu anomaly ($\text{Eu}/\text{Eu}^* = \text{Eu}_N/(\text{Sm}_N * \text{Gd}_N)^{0.5} \sim 0.65$). The two samples
348 show slightly contrasted heavy-REE (HREE) depletion with a Sm_N/Yb_N ratios of 4.5 and 11.2
349 for the samples from U150D and U1570A, respectively.



350

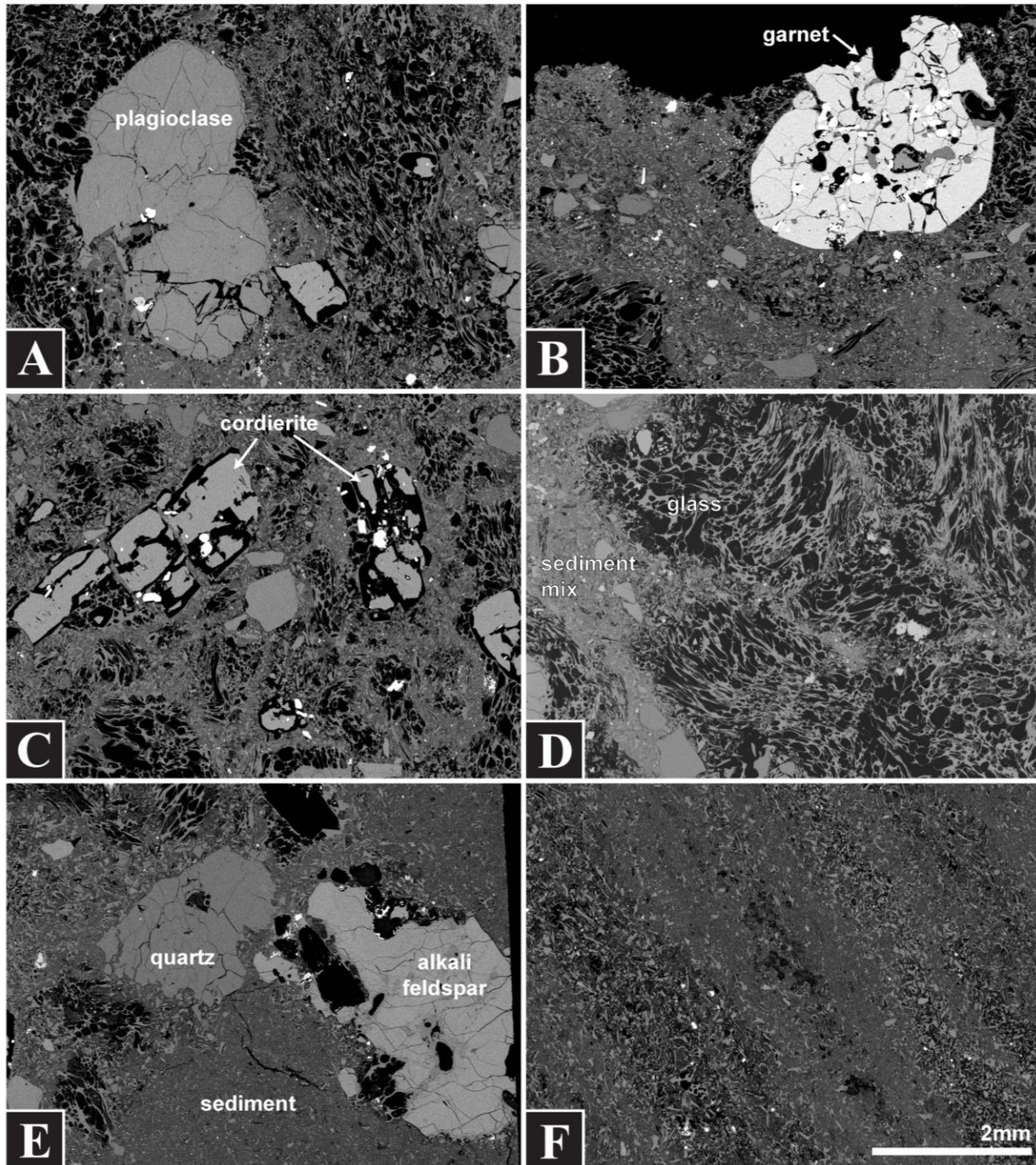
351 **Figure 3.** Total alkali-silica (TAS) diagram showing the bulk rock compositions of the U1570
 352 dacite (pink stars); analyses performed on clear glass (red squares) are interpreted as fresh dacitic
 353 melt and analyses performed on the brown glass interpreted as the peperitic matrix (brown
 354 squares). For comparison, hatched pentagon symbols are the bulk rock compositions of the ODP
 355 Leg 104 in Hole 642E dacitic rocks (Eldholm et al., 1989), and hatched triangles represent
 356 dacites from the Rockall Trough (RT) Well 163/6-1 A (Morton et al., 1988). Small open symbols
 357 represent compositions of melt inclusions (MI) in various solid phases (pl=plagioclase; grt =
 358 garnet; crd = cordierite) from the El Hoyazo, Spain enclaves; filled gray circles are the matrix
 359 glass compositions for the El Hoyazo lavas (Álvarez-Valero et al., 2005, 2007). The blue,
 360 crossed circle represents the matrix glass compositions of the Shiveluch Volcano (SV;
 361 Humphreys et al., 2008). (B) Trace element bulk concentrations normalized to the primitive

362 mantle (Sun and McDonough, 1989) of the two dacite samples compare with the average values
363 for the upper and lower continental crust (Rudnick and Gao, 2003).

364 3.2 Petrography and microstructures

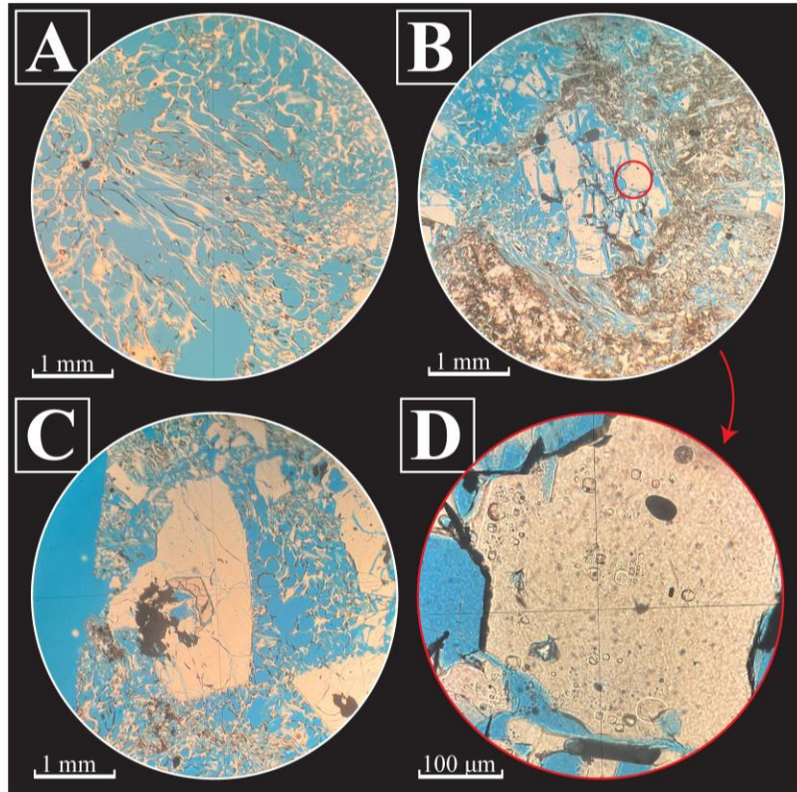
365 We estimated phase proportions using the GeoBalance mass balance calculation
366 spreadsheet based on a least squares problem (Li et al., 2020) for inputs of the mean bulk rock
367 composition, and the major element compositions obtained on solid phases and glass. The dacite
368 is composed of 67 ± 12 wt% glass with ~12 wt% cordierite, ~7 wt% plagioclase, ~5 wt% quartz,
369 ~2 wt% garnet, ~1 wt% alkali feldspar, ~1 wt% ilmenite, ~0.5 wt% apatite, and less than 6 wt%
370 of other accessory minerals (e.g., pyrite, zircon) and products of alteration (e.g., kaolinite).

371 The most abundant solid phase, cordierite, occurs as <1 mm to 4 mm large grains. These
372 are subhedral, with distinctive void space around the edges and along cracks (Fig. 4). Subhedral
373 almandine garnets were also identified in most samples. Both garnet and cordierite grains contain
374 abundant solid (ilmenite, pyrite, apatite, zircon) and melt inclusions (Fig. S3). Garnet is also
375 present in inclusion in cordierite (Fig. 5C). Large (2–6 mm) quartz grains are euhedral to
376 subhedral, often with very few or no inclusions (Fig. 4). Garnet, cordierite, and quartz are also
377 often highly fragmented (Figs. 4 and 5). Finally, plagioclase grains are subhedral with few
378 inclusions, sometimes with evidence of dissolution at the margins (Fig. 4A) and alkali feldspars
379 are mostly anhedral and sometimes contain large melt inclusions and embayments (Fig. 4E).
380 Smaller (<1mm) mineral phases disseminated in the samples include ilmenite, pyrite, apatite,
381 kaolinite, and graphite. Graphite is also observed both as a monomineral phase as thin deposits
382 on larger grains (Fig. 5). The matrix is rhyolitic glass (~77 wt% SiO₂; Fig. 3) in ash-sized glass
383 shards and micro-pumices. Mix domains exist where dark sand and silt are mingled within the
384 crystal-rich ash with a preserved grain structure (Figs. 4, 5).



385
 386 **Figure 4.** SEM-BSE images highlighting the various textures observed in the dacitic unit. (a)
 387 Plagioclase grain showing dissolution texture on the rim (sample U1570A-26R2-4). (b)
 388 Inclusion-rich garnet grain (sample U1570A-26R2-4). (c) Subhedral fractured cordierite grains
 389 surrounded by void space (sample U1570A-26R2-4). (d) Texture showing mingling between the
 390 vesicular glass and the sediments (sample U1570D-15R1-36). (e) Grains on the basal contact of

391 the dacite with sediments (sample U1570A-27R1-10). (f) Preserved layering of sediments less
 392 than a centimeter away from the basal contact of the dacite (sample U1570A-27R1-10). All
 393 images have the same scale.



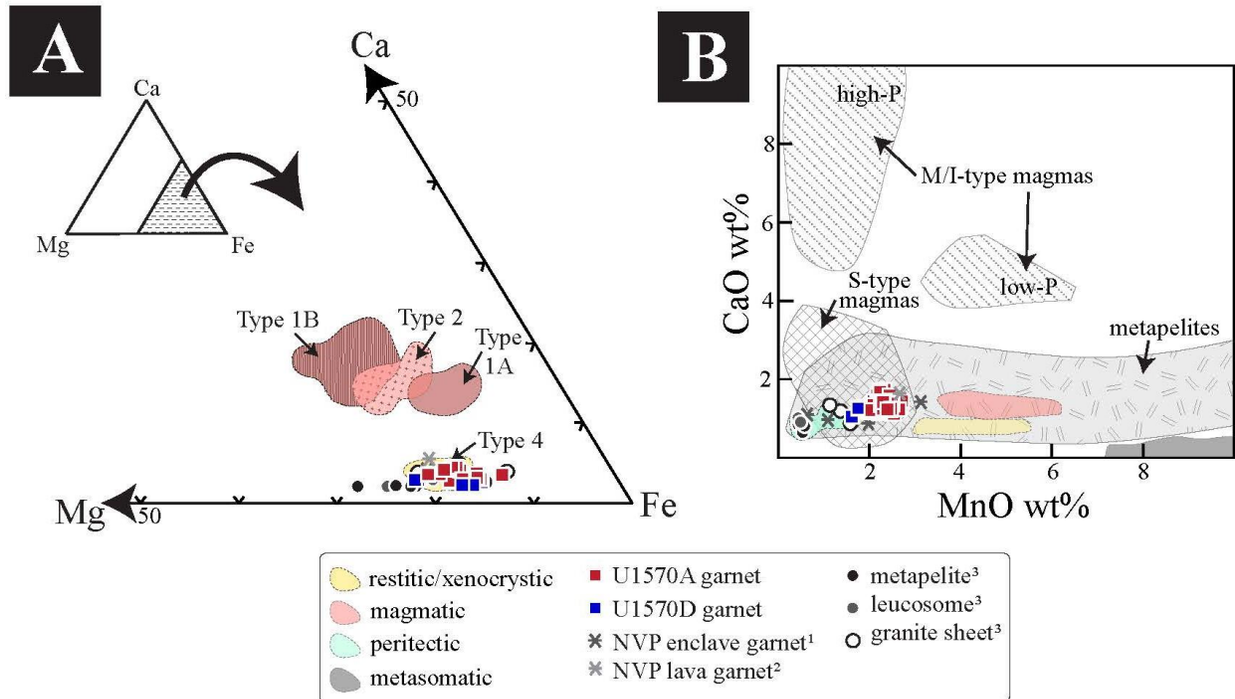
394
 395 **Figure 5.** Petrographic images of samples impregnated with blue epoxy to highlight textures and
 396 porosity. (a) Example of pumiceous glass texture with elongated and folded vesicles (sample
 397 U1570D-14R3-8). (b) Cordierite grain showing fragmented texture with associated graphite
 398 (sample U1570A-26R1-26). (c) Garnet and graphite in inclusion in a cordierite grain (sample
 399 U1570A-26R2-44). (d) Melt inclusions in cordierite grain from (b).

400 3.3 Mineral chemistry

401 3.3.1 Major and minor elements

402 The twelve garnets analyzed are all Fe-rich, almandine-pyrope solid solution (77–85%
 403 Alm, 7–15% Py; Fig. 6A; Table 1). They are MnO- and CaO-poor with fractions of spessartine
 404 (XSpss) and grossular (XGr) of 0.040 ± 0.003 (1σ) and 0.03 ± 0.01 (1σ), respectively (Fig. 6B).
 405 The garnets show a relatively constant Mg# (molar $\text{Mg}/[\text{Mg} + \text{Fe}] \times 100 = 12.7\% \pm 0.7\%$ (1σ).

406 Garnet crystals from samples across boreholes show no significant variations in major element
 407 concentrations. Ten of these garnets were large enough to acquire compositional profiles. There
 408 is no systematic zonation in the grains, and the compositional variability is limited with 0.05
 409 ± 0.04 wt% FeO and 0.21 ± 0.05 wt% MnO mean relative variability per grain (Fig. S4).



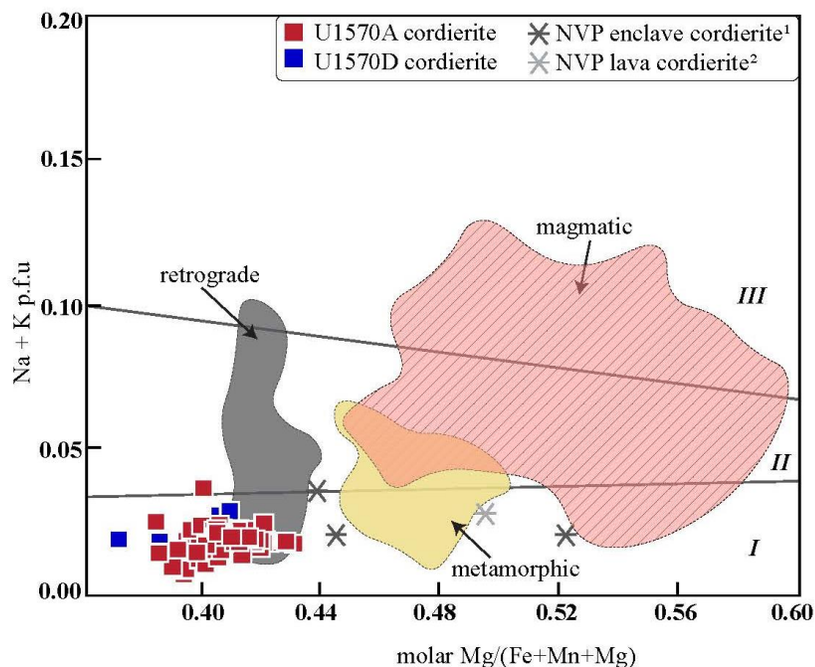
410
 411 **Figure 6.** (a) Mg-Ca-Fe ternary plot (Harangi *et al.*, 2001) with the compositions of garnets
 412 analyzed in enclaves found in the Neogene Volcanic Province (gray asterisks; Álvarez-Valero &
 413 Waters, 2010) and as separated grains in the associated lavas (black asterisks; Hiwatashi *et al.*,
 414 2021) and in Mkhondo Valley Metamorphic Suite rocks separated by the three stages of
 415 “selective peritectic phase entrainment”: 1) metapelite (black filled circles), 2) leucosome (gray
 416 circles), and granite sheet (black open circles; Taylor & Stevens, 2010). (b) CaO vs. MnO wt%
 417 plot; colored fields represent different types of garnet adapted from Khedr *et al.* (2022) and gray,
 418 patterned fields represent garnets from different sources adapted from Harangi *et al.* (2001). On
 419 both plots, restitic and xenocrystic-type garnet compositions are represented by yellow fields and
 420 magmatic garnet compositions are represented by light red fields. In (b), peritectic-type garnets
 421 are represented by the light blue field and metasomatic garnets by the solid dark gray field.
 422 U1570A (red squares) and U1570D (blue squares) garnet compositions are plotted. See text for
 423 details.

424 **Table 1.** Major element summary for major phases in the dacite samples.

	garnet <i>n</i> = 229		cordierite <i>n</i> = 94		plagioclase <i>n</i> = 60		alkali feldspar <i>n</i> = 26		quartz <i>n</i> = 61		ilmenite <i>n</i> = 62		kaolinite <i>n</i> = 15		apatite <i>n</i> = 5		glass <i>n</i> = 35	
	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ	< >	1 σ
SiO₂	35.89	0.69	47.22	0.50	56.22	1.58	65.12	0.45	99.84	0.05	0.21	0.31	78.84	2.10	0.00	0.00	76.60	0.33
TiO₂	0.05	0.07	0.01	0.01	0.02	0.01	0.03	0.01	0.03	0.02	51.45	0.66	0.14	0.04	0.04	0.01	0.12	0.01
Al₂O₃	21.40	0.42	33.09	0.51	27.72	1.14	19.78	0.57	0.08	0.01	0.15	0.03	13.23	0.23	0.01	0.00	13.05	0.10
FeO	36.66	0.99	13.83	0.39	0.10	0.03	0.03	0.02	0.03	0.02	47.08	0.66	2.03	0.21	1.61	0.16	1.45	0.24
MnO	1.78	0.13	0.23	0.03	0.02	0.01	0.01	0.01	0.01	0.01	0.46	0.04	0.05	0.01	0.21	0.08	0.03	0.02
MgO	2.99	0.14	5.38	0.12	0.00	0.00	0.00	0.00	0.01	0.02	0.67	0.02	0.09	0.02	0.23	0.01	0.09	0.01
CaO	1.05	0.08	0.01	0.01	9.08	1.23	0.09	0.05	0.01	0.00	0.01	0.01	0.31	0.04	54.91	1.46	0.32	0.02
Na₂O	0.02	0.04	0.08	0.02	6.10	0.59	3.20	0.07	0.01	0.02	0.02	0.02	2.26	0.82	0.00	0.00	3.28	0.15
K₂O	0.01	0.04	0.13	0.01	0.65	0.18	11.67	0.22	0.01	0.01	0.01	0.01	2.76	1.60	0.01	0.00	5.04	0.20
Total	100.74	3.53	100.65	0.81	0.11	0.02	98.60	1.23	99.26	1.13	101.36	1.61	93.55	3.33	98.25	0.71	92.37	2.64
	(12 oxygens)		(18 oxygens)		(8 oxygens)		(8 oxygens)		(2 oxygens)		(3 oxygens)		(9 oxygens)		(4 oxygens)			
Si	3.3477	0.0592	5.9169	0.0408	2.9955	0.0622	3.3588	0.1580	0.9991	0.0002	0.0010	0.0047	4.0754	0.0295	0.0000	0.0000		
Ti	0.0034	0.0036	0.0009	0.0009	0.0006	0.0005	0.0006	0.0010	0.0002	0.0001	0.9305	0.0099	0.0053	0.0014	0.0004	0.0005		
Al	1.1759	0.0248	2.4430	0.0461	0.8685	0.0440	0.6306	0.0966	0.0005	0.0001	0.0027	0.0020	0.4032	0.0126	0.0001	0.0001		
Fe	2.8527	0.1032	1.4536	0.0558	0.0107	0.0473	0.0388	0.1389	0.0002	0.0002	1.1041	0.0184	0.0880	0.0091	0.0500	0.0042		
Mn	0.1400	0.0107	0.0247	0.0035	0.0006	0.0006	0.0007	0.0023	0.0001	0.0001	0.0107	0.0010	0.0020	0.0006	0.0064	0.0025		
Mg	0.4157	0.0203	1.0032	0.0396	0.0001	0.0003	0.0254	0.0948	0.0001	0.0002	0.0120	0.0016	0.0061	0.0028	0.0127	0.0004		
Ca	0.1085	0.0446	0.0014	0.0010	0.5170	0.0749	0.0040	0.0034	0.0000	0.0000	0.0001	0.0005	0.0174	0.0020	2.1841	0.0226		
Na	0.0016	0.0048	0.0119	0.0255	0.3155	0.0282	0.1524	0.0337	0.0001	0.0001	0.0006	0.0013	0.1140	0.0425	0.0000	0.0000		
K	0.0005	0.0028	0.0103	0.0021	0.0221	0.0059	0.3627	0.0916	0.0000	0.0000	0.0003	0.0008	0.0920	0.0549	0.0002	0.0001		
% Alm	81.11	1.21																
% Spss	3.99	0.34																
% Grs	3.07	1.10																
% Py	11.83	0.69																

425 Oxides concentrations are given as wt. % and elemental concentrations are given as molar fractions. Molar fractions are normalized to 12, 18, 8, and 3 for garnet, cordierite, feldspars,
426 and ilmenite, respectively. Values are the averages of all the measured grains (< >) and the associated standard deviation (1 σ). Garnet solid solution end-member proportions are also
427 provided: Alm = almandine, Spss = spessartine, Grs = grossular, Py = Pyrope. Individual analyses are provided in Table S3.

428 The cordierite crystals have an $Mg\# = 40.8 \pm 1.6$. They have low abundances of Na and K
 429 with total alkali (Na + K) molar fraction of 0.02 ± 0.03 (Fig. 7). Plagioclase grains classify as
 430 labradorite, with 60.2% anorthite and 37.2% albite. Alkali feldspars are 46–72% orthoclase and
 431 28–46% albite (Fig. S5A). No significant variability across samples or boreholes is observed. All
 432 ilmenite crystals are mostly pure ilmenite with a $\leq 0.01\%$ pyrophanite ($MnTiO_3$) component
 433 (Table 1).

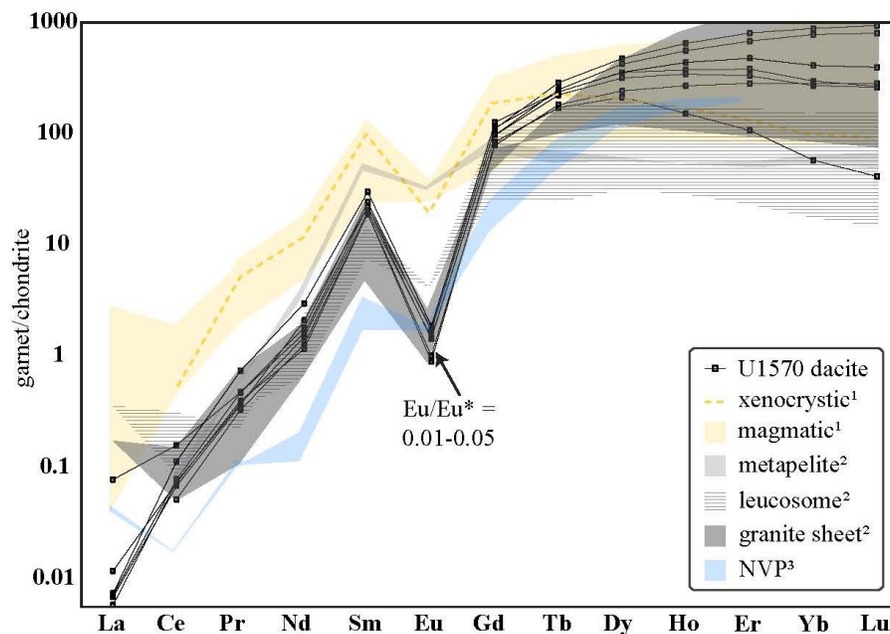


434
 435 **Figure 7.** (Na + K) p.f.u. (assuming 18 Oxygens) vs. molar $Mg/(Fe+Mn+Mg)$ ratios of cordierite
 436 grains from our study: samples from U1570A (red squares) and U1570D (blue
 437 squares). Cordierite compositions from enclaves (gray asterisks; Álvarez-Valero & Waters,
 438 2010) and separate grains in lavas (black asterisk; Hiwatashi et al., 2021) in the Neogene
 439 Volcanic Province volcanic rocks are shown for comparison. Adapted from Pereira and Bea
 440 (1994), the magmatic (red), metamorphic (yellow), and retrograde (gray) fields were classified
 441 by petrographic textures of cordierite grains. Solid lines define fields based on the chemical
 442 compositions of metamorphic (I), anatectic (II), and magmatic (III) cordierite grains.

443 3.3.2 Trace elements

444 Chondrite-normalized (McDonough & Sun, 1995) garnet REE patterns show notable
 445 LREE depletion ($Ce_N/Yb_N = 0.0001-0.0009$), with strong negative Eu anomalies ($Eu/Eu^* =$

446 0.01–0.04; Fig. 8). HREE slopes are variable, measured by Lu_N/Gd_N ratios ranging from 0.1–
 447 20.2 with a difference up to $\text{Lu}_N/\text{Gd}_N = 16.9$ within a single grain (Fig. S6). While some garnets
 448 show core-rim variation with steeper HREE slopes in their cores (Fig. S6), others show more
 449 complex patterns. Using the mathematical approach described by Anenburg and Williams
 450 (2022), no clear distinction can be made between the garnet grains. Garnet concentrates Sc, Ti,
 451 V, and Cr relative to other phases (Table 2). Y concentrations are highly heterogeneous, ranging
 452 from 170 to 1323 ppm, with intragranular variability as high as 858 ppm and are generally
 453 positively correlated with HREE concentrations.



454
 455 **Figure 8.** Chondrite-normalized (McDonough & Sun, 1995) garnet REE concentration diagrams.
 456 The black lines represent the mean composition of individual garnet grains analyzed in our
 457 samples (individual analyses on each garnet are plotted in Fig. S6B), compared with magmatic
 458 (yellow field) and xenocrystic (yellow, dashed line) garnets from Northern Pannonian Basin
 459 calc-alkaline volcanic rocks. The gray fields represent REE concentrations of garnets from three
 460 representative stages of “selective peritectic phase entrainment”, (1) the metapelite garnets (light
 461 solid gray), (2) leucosome garnets (horizontal gray lines), and (3) granite sheet garnets (solid
 462 dark gray). Neogene Volcanic Province volcanic rock garnets are represented by the blue field.
 463 Subscripts 1, 2, and 3 in the legend correspond to Harangi et al. (2001), Taylor & Stevens
 464 (2010), and Álvarez-Valero & Waters (2010), respectively.

465 **Table 2.** Mean trace element concentrations ($\mu\text{g}/\text{kg}$) for major solid phases in the dacite.

	garnet \pm		cordierite \pm		plagioclase \pm		alkali feldspar \pm		quartz \pm		zircon \pm	
	$n=25$		$n=28$		$n=22$		$n=18$		$n=62$		$n=88$	
Li									42	8		
Sc	240	112	1.12	0.74	0.16	1.17	0.8	3.0				
Ti	194	50	32.2	15.1	76.97	17.88	108	24	148	41	15	43
V	616	110	6.76	1.45	0.99	0.33	0.2	2.0			3	17
Cr	641	356	2.01	2.12	3.30	2.86	0.4	13.4				
Co	37.4	1.2	26.1	1.3	0.20	0.72	0.32	1.47				
Zn	178	13	277	18	8.55	13.86	6.7	27.4				
Ga	13.3	1.1	47.3	3.1	34.82	3.59	16.71	3.31				
Ge	11.6	2.3	0.50	2.58	1.47	2.51	1	8	1.00	0.36		
Rb									0.21	0.43		
Sr	0.04	0.03	0.07	0.13	662	72	472	68	0.29	0.55	0.9	1.45
Y	606	268	0.02	0.05	2.30	0.53	0.14	0.16			1650	686
Zr	23.9	10.0	0.30	1.07	0.06	0.13	0.00	0.27	0.31	0.59		
Ba									0.44	0.88		
La	0.01	0.02	0.01	0.04	33.67	7.56	4.48	0.75			5	33
Ce	0.06	0.04	0.03	0.11	59.8	13.6	4.23	0.72			19	78
Pr	0.05	0.02	0.00	0.01	5.66	1.44	0.26	0.10			1	5.7
Nd	0.88	0.41	0.01	0.03	19.1	5.5	0.48	0.24			5	21
Sm	3.57	0.81	0.01	0.02	3.10	0.69	0.06	0.09			7	7
Eu	0.09	0.04	0.00	0.00	6.02	0.60	5.08	0.84			0.52	0.74
Gd	21.7	4.5	0.03	0.06	1.83	0.54	0.02	0.07			43	17
Tb	8.92	1.84	0.00	0.00	0.20	0.06					14.09	4.84
Dy	91.3	28.1	0.01	0.02	0.81	0.31					162	62
Ho	24.0	12.1	0.00	0.01	0.10	0.05					56.43	24.1
Er	77.5	55.2	0.00	0.01	0.17	0.11					242	117
Yb	78.4	79.6	0.00	0.00	0.05	0.06					432	244
Lu	11.8	13.5	0.00	0.00	0.01	0.03					87	50
Hf	0.46	0.18	0.01	0.04	0.00	0.07	0.02	0.05			11094	1200
Ta	0.01	0.00	0.00	0.02	0.00	0.00	0.07	0.04			0.73	0.97
Th	0.01	0.00	0.01	0.04	0.01	0.02	0.01	0.04			118	354
U	0.01	0.01	0.01	0.03	0.00	0.02					271	275

466 Values are the averages of all the measured grains (< >) and the associated standard deviation (1σ). Individual analyses are
467 reported in Table S4.

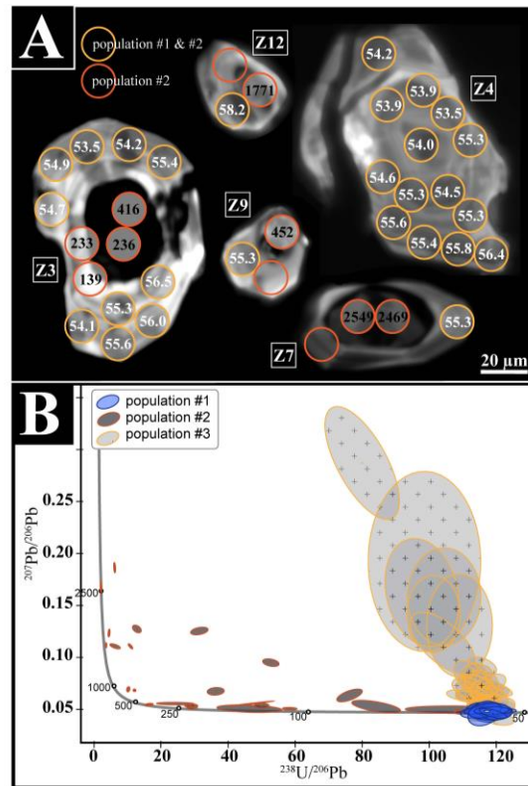
468 Trace element analysis for cordierite crystals shows they are very REE-depleted (ΣREE_N
469 < 1.5 ppm). Cordierites are Zn-enriched and Sr-depleted relative to other phases ($\text{Zn} = 277 \pm 18$
470 ppm, $\text{Sr} = 0.07 \pm 0.13$ ppm; Table 2). Plagioclase is LREE-enriched relative to HREE (La_N/Sm_N
471 $= 5.6\text{--}12.0$). They are relatively enriched in Sr, Ti, and Cr with small positive Eu anomalies
472 ($\text{Eu}/\text{Eu}^* = 5.0\text{--}10.0$; Fig. S5B; Table 2). In comparison, alkali feldspars are significantly more
473 LREE-depleted, but present more fractionated patterns ($\text{La}_N/\text{Sm}_N = 10.1\text{--}125.0$). They also

474 concentrate more Ti and less Sr than plagioclase and have strong positive Eu anomalies (Eu/Eu*
475 189.2–521.0; Fig. S5B; Table 2).

476

477 3.4 U-Pb zircon ages

478 Zircon grains have only been observed as inclusions in garnet and cordierite and are
479 dominantly subhedral with rare euhedral crystals. Many of the zircons are fractured and range
480 from 10 to 100 μ m in diameter (Fig. 9A). Zircon analyses were plotted on a Tera-Wasserburg
481 concordia plot ($^{238}\text{U}/^{206}\text{Pb}$ vs. $^{207}\text{Pb}/^{206}\text{Pb}$; Tera & Wasserburg, 1972) in the IsoplotR toolbox
482 (Vermeesch, 2018). The data define a ternary mixing field in Tera-Wasserburg space with the
483 endmembers interpreted as (1) dominant younger populations, with a subset of concordant
484 analyses and (2) sparse older inherited population of zircon. Non-anchored, uncertainty-weighted
485 linear regression (i.e. model-1 fit) of the young, concordant analyses with the array of young
486 analyses containing variable amounts of common Pb (population 1, Fig. 9B) yielded an initial
487 lead composition of 0.79 ± 0.1 ($^{207}\text{Pb}/^{206}\text{Pb}$) and lower intercept of 54.8 ± 1.1 Ma (MSWD =
488 1.7). The lower intercept of a linear regression anchored to Stacey-Kramer initial Pb value of
489 0.85 also yielded a 54.8 ± 1.1 Ma (mean square weighted deviation, MSWD = 1.7; Ludwig,
490 1997; Andersen, 2002; Chew et al., 2014). Spot dates were calculated using the Stacey-Kramer
491 anchored ^{207}Pb correction (Fig. 9A). Analyses with $^{238}\text{U}/^{206}\text{Pb}$ less than 105 and concordance
492 outside of 0.9–1.05 were excluded from the weighted mean calculation. The weighted mean date
493 of 54.6 ± 1.1 Ma, calculated using concordant analyses from population 1, is interpreted as the
494 magmatic age of the dacite. The analyses interpreted as inherited (#2 above) yielded spot dates
495 ranging from 61–2500 Ma (Fig. S7).



496

497 **Figure 9.** (a) Representative cathodoluminescence images of zircon grains from sample
 498 U1570A-26R-2-4. Circles represent LA-ICP-MS analysis locations, and the numbers are the
 499 calculated spot age for that analysis. Empty circles show spot ages that did not fall on the
 500 regression from the weighted average and were too discordant (concordance >0.9) to calculate
 501 ages. Populations #1 and #3 (magmatic) and #2 (inherited) are represented in yellow and orange,
 502 respectively. (b) U-Pb isotope results for zircon on Tera-Wasserburg Concordia plot. Only
 503 analyses in blue were used to calculate the formation age of the dacite (see text for details).
 504 Population #1 includes samples that fit a regression from the weighted average to the expected
 505 original $^{207}\text{Pb}/^{206}\text{Pb}$ and represent magmatic young zircons. Samples that do not fit the regression
 506 are included in population #2, representing older inherited zircon.

507

508 3.5 Sedimentary mercury concentration and organic carbon maturation data

509 The stable carbon isotope analyses of bulk organic matter ($\delta^{13}\text{C}_{\text{org}}$), total organic carbon
 510 (TOC) and mercury (Hg) contents in the sediments collected in borehole U1570A and U1570D

511 are plotted in Figure 2 and reported in Tables S5 and S6. Sedimentary mercury concentrations
512 are ~190 ppb (mean) for the sediments analyzed for U1570D (81.23 meters below surface, mbsf,
513 CSF-A to 192.28 mbsf). Two strongly Hg enriched samples (> 1 ppm; 87.25 and 172.4 mbsf)
514 were found but these occur 10s of meters from the dacitic unit. For other samples, mercury
515 concentrations are above average shale values (~ 62 ppb, Grasby et al., 2019) throughout the
516 succession but no clear trends in Hg concentration were observed towards the dacite unit.

517 Organic-carbon maturation data obtained through RockEval analyses indicates the
518 organic-matter preserved in the sediments is immature; Tmax averages ~ 405 °C. RockEval
519 parameters HI (mean ~50) and OI (mean ~130) signal a dominant terrestrial organic-matter
520 composition. Total organic carbon concentration measurements yielded ~0.6 wt% for the
521 Paleocene interval and ~1.1 wt% for the lower Eocene and, similar to Hg concentrations, none of
522 the RockEval parameters (TOC, HI, OI, Tmax) show trends towards the dacite unit.

523

524 **4 Interpretation of texture and mineral chemistry of garnet and cordierite**

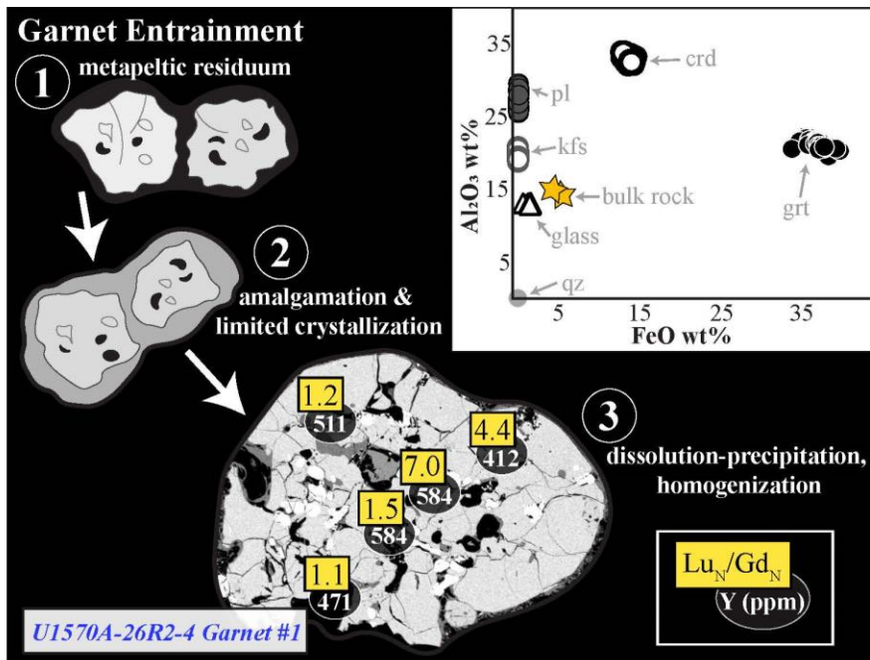
525 Our samples contain mineral phases commonly found in crustal metapelites and restites,
526 particularly garnet and cordierite produced during crustal anatexis (Fig. 4, 5; Harley &
527 Carrington, 2001; Vry et al., 1990; White & Chappell, 1977; Stevens et al., 1995, 2007;
528 Weinburg & Hassalova, 2015). In this context, these minerals could be (1) liquidus phases
529 produced during the crystallization of a magma as phenocrysts; (2) disseminated xenocrysts from
530 the original protolith; or (3) peritectic phases entrained during magma transport. A magmatic
531 chemical signature would indicate they are liquidus phases, while a metamorphic signature
532 would indicate they are either disseminated xenocrysts or peritectic phases which preserve the
533 chemistry of the protolith. Below, we use several proxies to distinguish magmatic and
534 metamorphic major and trace element compositions of garnet and cordierite.

535 Harangi et al. 2001 and Bach et al. (2012) presented similar classifications based on the
536 Ca, Mg, and Fe (\pm Mn) components of the garnet solid solution. Both studies show that magmatic
537 garnets are systematically more enriched in grossular than metamorphic garnets (Fig. 6). Garnet
538 compositions in our study are relatively depleted in grossular, overlapping the compositions of
539 xenocrystic garnets described in Harangi et al. (2001) and plotting near metamorphic garnet

540 compositions in Bach et al. (2012)'s classification, but with a more significant almandine-
541 spessartine enrichment in comparison to the garnet compositions reported in their study.
542 According to the classification defined by Pereira and Bea (1994) based on alkali content and
543 Mg# (= molar Mg / [Mg+Fe+Mn]), the cordierite grains in our samples also show a metamorphic
544 major element signature (Fig. 7).

545 Harangi et al. (2001) did not report trace element analyses on pure xenocrystic garnets
546 but rather on composite garnets, interpreted as xenocryst cores surrounded by magmatic rims
547 (Fig. 8). Both the magmatic and composite garnets from their study show REE patterns similar to
548 those observed in our samples, displaying substantial LREE depletion and variable HREE slope.
549 While none of them have negative Eu anomalies as prominent as those observed in the garnet
550 grains from the Site U1570 dacite, the Eu anomalies for the xenocrystic cores of composite
551 garnets ($\text{Eu}/\text{Eu}^* = 0.14$) are stronger than those reported for the magmatic garnets ($\text{Eu}/\text{Eu}^* =$
552 $0.17\text{--}0.68$; Fig. 8). Harangi et al. (2001) also reported depleted Y (277 ppm) and enriched Zr and
553 Hf (178 and 2.75 ppm, respectively) for the xenocrystic core compositions relative to magmatic
554 compositions. In comparison, garnets from our study have more variable Y contents (170–1323
555 ppm) and lower Zr and Hf concentrations (9–47 and 0.20–0.84 ppm, respectively; Table 2).
556 However, trace elements concentrations of magmatic garnets reported in Bach et al. (2012) show
557 Y, Zr, and Hf variability covering the entire range of those presented for magmatic and
558 xenocrystic garnets in Harangi et al. (2001), indicating that these elements are not reliable
559 proxies for garnet origin.

560 In their study of mid-crustal anatectites from the Mkhondo Valley Metamorphic Suite,
561 Swaziland, Taylor and Stevens (2010) outlined how garnet can undergo several stages of
562 entrainment that result in modifications to both their major and trace element compositions (Fig.
563 10). By analyzing almandine garnet from metapelites, leucosomes, and granite sheets, they
564 proposed these garnets represent a sequence of processes related to “selective peritectic phase
565 entrainment,” in which only solid peritectic products become entrained by associated melts
566 (Stevens et al., 2007). This model provides a mechanism to produce hot, water-undersaturated,
567 granitic magmas. Preferential entrainment of garnet and ilmenite during high-temperature,
568 incongruent, fluid-absent melting of biotite metapelites results in an increase in A/CNK, Mg#,
569 and Ca content and a decrease in Si and K in the magma.



570
 571 **Figure 10.** Summary of thermobarometric calculations. The gray dashed line represents the
 572 mean temperature calculated for Ti concentration in quartz using Wark and Watson (2006)’s
 573 original TitaniQ thermometer with no pressure correction (standard deviation represented by the
 574 gray field). Black circles represent the pressure-dependent thermometer for Ti concentration in
 575 quartz for Huang and Audéat (2012). For comparison the pressure correction from Thomas et al.
 576 (2010) is also plotted (gray circles). The vertical red line and red field represent the mean
 577 temperature and standard deviation calculated using the Fe-Mn exchange between garnet-
 578 ilmenite (Grt-Ilm; Pownceby et al., 1991). The red triangles show the pressures calculated with
 579 the garnet barometer (Wu, 2019) and the blue squares represent the temperatures calculated with
 580 the garnet-cordierite (Grt-Crd) thermometer (Kaneko & Miyano, 2004). The pressure and
 581 temperature field obtained using 1wt% H₂O and the bulk rock composition in the MAGEMin
 582 package (Riel et al., 2022) for the assemblage observed in our samples is shown by the hatched
 583 triangular area. Subscripts in legend represent references: (1) Wark and Watson (2006), (2)
 584 Thomas et al. (2010), (3) Huang and Audéat (2012), (4) Pownceby et al. (1991), (5) Kaneko &
 585 Miyano (2004), (6) Wu (2019), (7) Riel et al. (2022).

586 From their study, Taylor and Stevens (2010) describe three stages of the “selective
 587 peritectic phase entrainment model. Peritectic garnet growth in rapid partial melts of metapelites
 588 represents the first stage of entrainment and these garnets have Mg# = 20–27, XSpss = 0.01, and

589 homogenous compositions. The second stage of entrainment is present in the peritectic garnets
590 from leucosomes, interpreted to have undergone limited recrystallization and some
591 amalgamation at high temperatures with $Mg\# = 20\text{--}22$ and $XSpss = 0.01$. Finally, the garnets
592 from the granite sheets are rounded and euhedral, resulting from dissolution-precipitation
593 reactions, amalgamation, and re-equilibration with the magma. This third stage is further divided
594 into large (0.8–4 mm) garnets with $Mg\# = 18\text{--}21$ and $XSpss = 0.02$ and very small (75–600 μm)
595 garnets with $Mg\# = 12\text{--}14$ and $XSpss = 0.02\text{--}0.03$. Both large and small garnets from granite
596 sheets have rims with slightly lower $Mg\#$ and higher $XSpss$ contents than their cores (Taylor &
597 Stevens, 2010). With homogenous $Mg\# = 12\text{--}13$ and $XSpss = 0.04$, the garnets in our samples
598 are compositionally most similar to those from the granite sheet (Fig. 6). Additionally, the
599 presence of melt inclusions in both garnet and cordierite (Fig. 5) provides evidence for rapid
600 crystal growth (Cesare et al., 2011).

601 Taylor and Stevens (2010) also presented chondrite normalized REE data for the garnets
602 in their study. Representing the first stage of garnet entrainment, metapelite garnets have flat,
603 relatively depleted HREE patterns and weak negative Eu anomalies ($Lu_N/Gd_N = 0.98\text{--}0.16$,
604 $Eu/Eu^* = 0.5$). Leucosome garnets from the second stage have slightly steeper HREE slopes and
605 much stronger negative Eu anomalies ($Lu_N/Gd_N = 1.59\text{--}0.56$, $Eu/Eu^* = 0.07$). Garnets from the
606 third stage, the granite sheets, have the steepest HREE slopes and the most pronounced negative
607 Eu anomalies ($Lu_N/Gd_N = 1.15\text{--}28.38$, $Eu/Eu^* = 0.03$). The strongly negative Eu anomalies of
608 the U1570 dacite garnets overlap with the latter stages of entrainment, where evidence of
609 dissolution-precipitation reactions is prominent. Taylor and Stevens (2010) also reported changes
610 in HREE slope from steeper in the core to flatter in the rim of garnet in the granite sheets, and
611 complex zonation of yttrium (Y) concentrations, which they attribute to the amalgamation of
612 grains. The HREE slopes in the Site U1570 dacite garnets cover the range presented in Taylor
613 and Stevens (2010), though most overlap with the stage three granite sheet garnets, and some
614 show evidence of the same zonation patterns from core to rim (Fig. 10). The garnets from our
615 study also have similarly complex Y zoning, with a broad pattern of Y enrichment in the cores
616 relative to the rims (Fig. 10). Finally, the bulk composition of our samples shows depletion
617 in HREE, Zr and Hf (Fig 3B, S2) consistent with the presence of residual garnet in the source.
618 Peritectic garnets are expected to be enriched in HREE (Fig. 8; Taylor and Stevens, 2010). While
619 present as large grains, the fraction of garnet in our samples is relatively small (< 2 wt.%).

620 Hence, if some of these newly formed grains were not entrained in the magma, they might
 621 explain the HREE (and Hf and Zr) depletion of the bulk rock.

622 In summary, garnet compositions in our samples have notable similarities in major and
 623 trace element compositions with peritectic phases entrained in a melt. Notably, they are
 624 chemically similar to entrained garnets resulting from dissolution-precipitation reactions grains
 625 in a magma. Through re-equilibration with the magma and amalgamation of new grains formed
 626 by peritectic reaction, dissolved metapelite garnets can be re-precipitated with their major
 627 elements homogenized (Taylor & Stevens, 2010). These processes can produce garnets with
 628 magmatic textures, xenocrystic, homogenous, major element chemistry, and complex trace
 629 element zonation through peritectic growth in a hot, water-undersaturated magma during
 630 metapelite melting (Clarke, 2007). At Mimir High, the main solid peritectic products which are
 631 transported within the dacite magma toward the surface are Fe-rich garnet and cordierite,
 632 resulting in a bulk rock composition with high A/CNK and Fe contents and relatively low Si and
 633 K contents (Figs. 3, 10).

634 **5 Geothermobarometry and pseudosections**

635 The mineral textures (subhedral morphology, melt inclusions) and mineral chemistry of
 636 the garnet and cordierite grains support their origin as products of peritectic melt reactions that
 637 have been entrained in the dacitic melt. During evolution and transport the dacite may have been
 638 impacted by variable degrees of re-crystallization and chemical re-equilibrium. As such, the
 639 observed mineral assemblage and bulk rock chemistry can be used to constrain the pressure,
 640 temperature, and water saturation conditions of melt generation of the magma.

641 **5.1 Ti in quartz, TitaniQ thermometer**

642 Wark and Watson (2006) formulated the TitaniQ thermometer based on the temperature-
 643 dependent substitution of titanium for silicon in the quartz matrix over a temperature range of
 644 600–1000 °C at 10 kbar. Their equation for rutile-undersaturated rocks is:

$$645 \quad T(^{\circ}\text{C}) = \frac{-3765}{\log\left(\frac{X_{\text{qtz}}^{\text{Ti}}}{a_{\text{TiO}_2}}\right) - 5.69} - 273 \quad (1)$$

646 where X_{Ti}^{qtz} is the concentration (ppm) of Ti in quartz and a_{TiO_2} is the activity of TiO_2 in the
 647 melt.

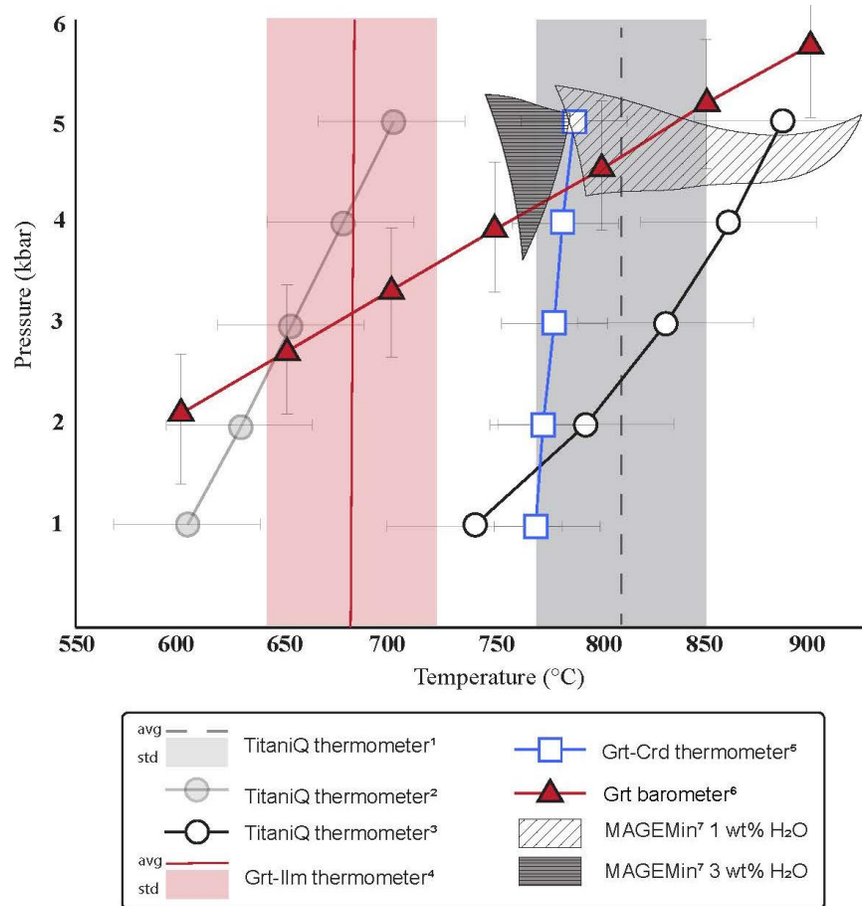
648 For silicic igneous rocks that are TiO_2 undersaturated, indicated by the lack of rutile as a
 649 stable phase, the activity of titanium in the melt < 1 and must be estimated to apply TitaniQ.
 650 Ghiorso and Gualda (2013) calculated the activity of titania in the liquid relative to rutile
 651 saturation (a_{TiO_2}) for the Shiveluch Volcano dacite using two independent methods: one based
 652 on the composition of coexisting spinel and ilmenite in the magma, and one calculated with
 653 rhyolite-MELTS (Gualda et al., 2012) using the rhyolitic matrix glass composition from
 654 Humphreys et al. (2008). This melt composition is comparable to the rhyolite glass in our sample
 655 with 78.0 wt% SiO_2 , 7.4 wt% alkali content, 12.4 wt% Al_2O_3 , and 0.3 wt% TiO_2 (Fig. 3). The
 656 two sets of calculations intersect for a_{TiO_2} between 0.75 and 0.90 for 730–800 °C (see Fig. 3 in
 657 Ghiorso & Gualda, 2013).

658 Using equation (1), variation of a_{TiO_2} within ± 0.1 only results in temperature variation
 659 within $\pm 20^\circ C$. The mean Ti concentration and its one standard deviation in quartz hosted by the
 660 dacite is 142.04 ± 37.8 ppm. Assuming $a_{TiO_2} = 0.8$, we obtain a mean crystallization temperature
 661 of 817 ± 42 °C (Fig. 11; Table 3).

662 A few studies have reformulated TitaniQ to include the effect of pressure (e.g., Thomas
 663 et al., 2010; Huang & Audétat, 2012). In their experimental study, Thomas et al. (2010) showed
 664 that the solubility of Ti in quartz decreases with increasing pressure. They present a pressure-
 665 dependent TitaniQ thermometer calibrated between 5 and 20 kbar. Huang and Audétat (2012),
 666 however, argued that Thomas et al. (2010)'s thermometer overestimates Ti solubility, and
 667 consequently the effect of pressure, in their experiments. Using new rutile-saturated experiments
 668 performed between 1 and 10 kbars, the authors proposed the following equation:

$$669 \quad \log Ti = -0.27943 \left(\frac{10^4}{T} \right) - 660.53 \left(\frac{P^{0.35}}{T} \right) + 5.6459 \quad (2)$$

670 where Ti is the rutile-saturated concentration of Ti (in ppm) in quartz, T the temperature in
 671 Kelvin, and P pressure in kbar. For natural samples that are not saturated in rutile, $X_{Ti}^{qtz-sat}$ must
 672 be calculated using a_{TiO_2} , such as $X_{Ti}^{qtz-sat} = X_{Ti}^{qtz} / a_{TiO_2}$. Temperatures calculated in our
 673 study using the Huang and Audétat (2012) thermometer and $a_{TiO_2} = 0.8$ range from 886 ± 45 °C
 674 at 5 kbar down to 740 ± 39 °C at 1 kbar (Fig. 11; Table 3).



675

676 **Figure 11.** Schematic diagram for selective peritectic phase entrainment model for garnet grains.

677 (1) Grains of peritectic phases from metapelitic residuum become entrained in the melt. (2)

678 Residual grains experience limited new crystallization and amalgamate in the second stage. (3)

679 BSE image of garnet from sample U1570A-26R2-4 representing the third stage in which

680 amalgamated grains dissolve and re-precipitate as single grains with homogenized major element

681 compositions but variable trace element compositions. Adapted from Taylor and Stevens (2010).

682 LA-ICP-MS analysis locations are represented by yellow squares; values on yellow squares are

683 the corresponding HREE slopes (i.e., Lu_N/Gd_N , chondrite-normalized ratio). Values in black684 circles are the Y (ppm) concentrations at the same location. The Al_2O_3 vs. FeO wt% plot shows

685 bulk rock compositions (yellow stars) compared to major phases including garnet (black filled

686 circles), cordierite (black open circles), plagioclase (dark gray filled circles), alkali feldspar (gray
687 open circles), quartz (light gray filled circles), and matrix glass (black open triangles).

688 **Table 3.** Summary of calculated pressure and temperature conditions of the dacite sample
689 formation estimated from thermodynamic calculations and thermobarometers on mineral phases.

		H ₂ O	T (°C)		P (kbar)
MAGEMin	Riel et al., 2021	1 Wt%	725-800		1.5-3.5
pseudosection		3 Wt%	710-720		2.9-3.2
TitaniQ	Wark & Watson, 2006		817	± 42	1
thermometer	Thomas et al., 2010		604	± 35	1
<i>a</i> TiO ₂ = 0.8			652	± 37	3
			701	± 39	5
	Huang & Audétat, 2012		740	± 39	1
			831	± 43	3
			886	± 45	5
Grt-Ilm	Pownceby et al., 1991		686	± 37	
thermometer					
Grt-Crd	Kaneko & Miyano, 2004		769	± 45	1
thermometer			778	± 45	3
			787	± 45	5
Grt	Wu, 2019		600		2.1 ± 1.3
barometer			700		3.3 ± 1.3
			800		4.6 ± 1.3
			900		5.8 ± 1.3

690 Pseudosection results are given as ranges for which the observed phase assemblage is stable. Thermometer
691 results are given as an average and one standard deviation or temperature with corresponding input pressures
692 if applicable. Barometers results are given as an average and one standard deviation for each input
693 temperature.

694 5.2 Garnet-ilmenite and garnet-cordierite thermometers

695 Xenocrysts present in the dacite allow for the use of the garnet-ilmenite geothermometer
696 based on Fe-Mn exchange reactions and the garnet-cordierite geothermometer based on Fe-Mg
697 exchange reactions.

698 The Fe-Mn exchange between garnet and ilmenite was investigated by Pownceby et al.
699 (1987), who developed a pressure independent geothermometer. Pownceby et al. (1991)
700 reinvestigated the original thermometer to include the influence of grossular content in garnet
701 and presented the thermometer:

$$T = \frac{14918 - 2200(2X_{\text{Mn}}^{\text{Ilm}} - 1) + 620(X_{\text{Mn}}^{\text{Grt}} - X_{\text{Fe}}^{\text{Grt}}) - 972X_{\text{Ca}}^{\text{Grt}}}{R \ln K_d + 4.38} - 273.15 \quad (3)$$

where R is the gas constant, T is the temperature in Celsius, $X_{\text{Mn}}^{\text{Ilm}}$ is the molar concentration of Mn in ilmenite, and $X_{\text{Mn}}^{\text{Grt}}$, $X_{\text{Fe}}^{\text{Grt}}$, $X_{\text{Ca}}^{\text{Grt}}$ are the molar concentrations of Mn, Fe, and Ca in garnet, respectively. K_d is the distribution coefficient defined by:

$$K_d = \frac{(X_{\text{Mn}}^{\text{Grt}}) \times (X_{\text{Fe}}^{\text{Ilm}})}{(X_{\text{Fe}}^{\text{Grt}}) \times (X_{\text{Mn}}^{\text{Ilm}})} \quad (4)$$

Using experimental results performed at 600–1000 °C and 10 kbar and FeTiO_3 – MnTiO_3 activity data, their thermometer has a resolution of ± 30 – 50 °C. In our study we only used this thermometer on ilmenite inclusions in garnet to account for the rapid diffusion of Fe–Mn in ilmenite. In fact, the relatively slow Fe–Mn diffusion in garnet effectively shields the ilmenite from re-equilibration and records temperatures for garnet growth (Pownceby et al., 1991). Garnet-ilmenite temperature results for seven garnets in the dacite give an mean of 686 ± 37 °C (Fig. 11; Table 3).

The garnet-cordierite geothermometer is commonly used to determine temperatures in high-grade amphibole and granulite facies metamorphic rocks (Spear & Selverstone, 1983). However, it is considered to be a poorly calibrated thermometer because cordierite is prone to severe alteration in high-grade metapelites. We note, however, alteration is not observed in our sample (Figs. 4, 5). In addition, the presence of a garnet inclusion in cordierite (Fig. 5C) supports chemical equilibrium between the two phases. Several calibrations have been proposed for this thermometer (e.g., Thompson, 1976; Holdaway & Lee, 1977; Perchuck et al., 1985; Wells, 1979). Here we present results obtained with Kaneko and Miyano (2004)'s calibration:

$$T = \frac{-26144.7 + (-0.122 + W_V^{\text{Grt}})(P - 1) + W_H^{\text{Grt}} - 80.44(\text{Mg}^{\text{Crd}} - \text{Fe}^{\text{Crd}})}{-12.7094 - R \ln K_d + W_S^{\text{Grt}} + 1.642(\text{Mg}^{\text{Crd}} - \text{Fe}^{\text{Crd}})} - 273.15 \quad (5)$$

where T is Celsius, and P is in bar. The W_H^{Grt} , W_S^{Grt} , and W_V^{Grt} are the non-ideal mixing terms for the garnet solid solution based on Berman (1990)'s expression (see equations A5, A6, and A7, respectively in Kaneko and Miyano, 2004). The terms Fe^{grt} , Mg^{cd} , etc. denote the molar fractions of Fe and Mg in the garnet and cordierite solid solutions such as $\text{Mg}^{\text{Crd}} = \text{Mg}/[\text{Mg} + \text{Fe}]$, and $\text{Mg}^{\text{Grt}} = \text{Mg}/[\text{Mg} + \text{Fe} + \text{Ca} + \text{Mn}]$, with Mg, Fe, Ca and Mn in mol. %. Finally, the exchange coefficient (K_d) defined by:

$$729 \quad K_d = (Fe^{Grt})/(Mg^{Grt}) \times (Mg^{Crd})/(Fe^{Crd}) \quad (6)$$

730 The estimated absolute error for Kaneko and Miyano (2004)'s calibration is $\pm 45^\circ\text{C}$.
 731 Calculated temperatures using the compositions of garnet and cordierite grains in our samples
 732 and equation (5) and (6) show significantly higher temperatures ($\sim 770\text{--}790^\circ\text{C}$, Fig. 11; Table 3)
 733 than the garnet-ilmenite thermometer, with an effect of pressure smaller than the estimated
 734 uncertainty on the temperature.

735 5.3 Garnet barometer

736 While commonly used to determine pressures for metapelites, the garnet- Al_2SiO_5 -
 737 plagioclase-quartz (GASP) barometer (Holdaway, 2001) cannot be applied to our samples due to
 738 the absence of an aluminosilicate and biotite in the assemblage. Additionally, the garnet-rutile-
 739 ilmenite-plagioclase-silica (GRIPS) barometer (Bohlen & Liotta, 1986) is not recommended for
 740 pressures below 6 kbar (Wu & Zhao, 2006). Instead, we use the garnet barometer developed by
 741 Wu (2019) for metapelitic assemblages to estimate the pressure recorded by the phase
 742 assemblage in our samples. In fact, Wu (2019)'s barometer was initially developed because
 743 plagioclase is often CaO-poor or absent in metapelites, making barometers such as GASP and
 744 GRIPS less precise or unusable. It is empirically calibrated using natural metapelite samples with
 745 pressure and temperature conditions of 1–15 kbar and 430–900 $^\circ\text{C}$ for garnets with molar
 746 fractions of calcium and iron in the solid solution of 0.02–0.29 and 0.42–0.91, respectively:

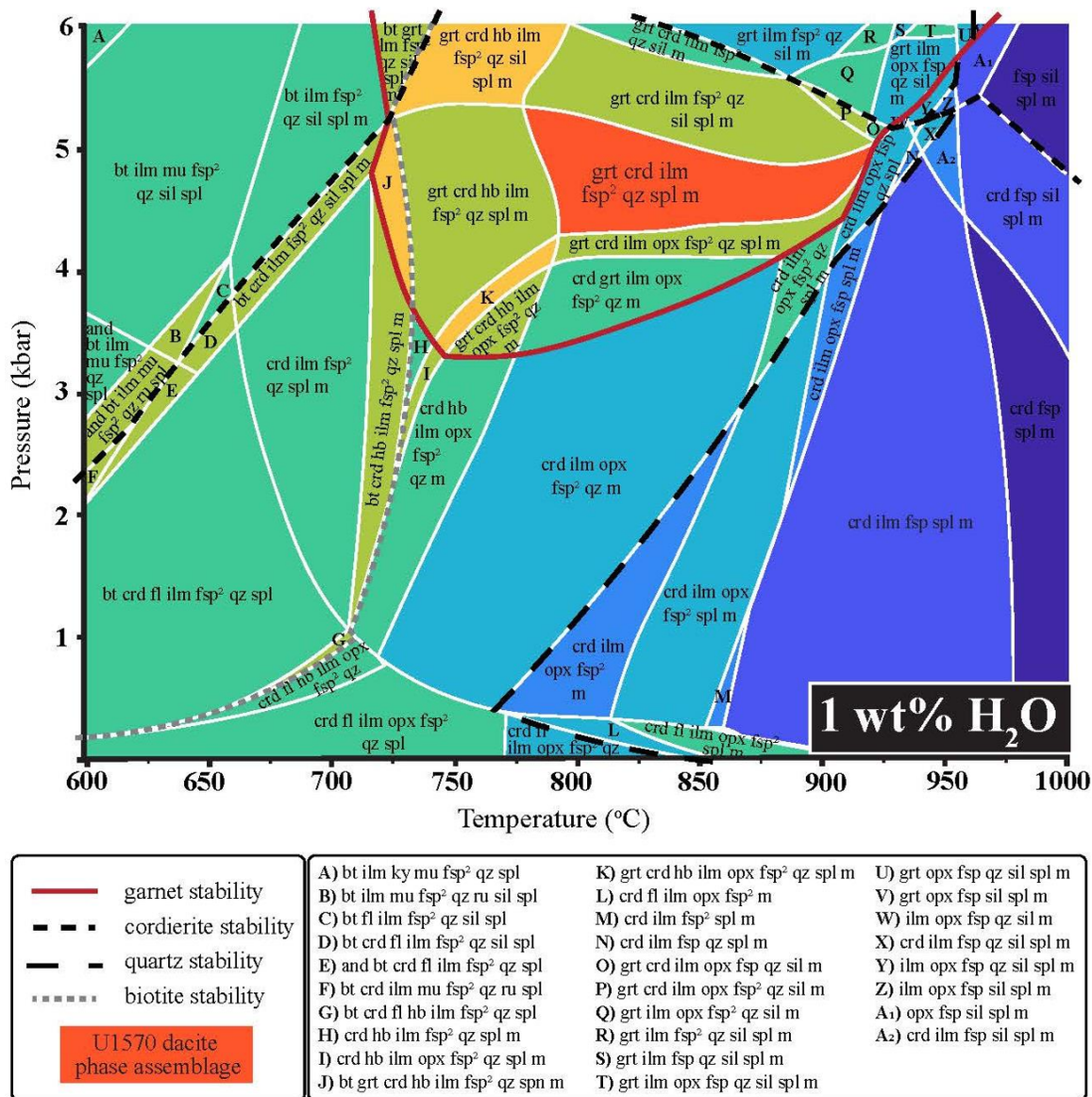
$$747 \quad P \text{ (bar)} = \frac{-8904.5 + 24.542T + 0.45RT \ln(Ca^{Grt}/Fe^{Grt}) + 0.15aT + 0.15c}{1 - 0.15b} \quad (7)$$

748 T is in Kelvin and a, b, and c are polynomial coefficients (see equations (5), (6), and (7),
 749 respectively in Wu (2019), which account for differences of excess chemical potential between
 750 grossular and almandine components in the garnet solid solution; Holdaway, 2001). The
 751 estimated error for pressures calculated using equation (7) is less than ± 1.3 kbar. Using our
 752 sample's mean garnet composition (Table 2), we obtain a range of pressures from ~ 2.1 kbar at
 753 600 $^\circ\text{C}$ to ~ 5.7 kbar at 900 $^\circ\text{C}$ (Fig. 11; Table 3).

754 5.4 Pseudosections

755 The phase assemblage calculated with MAGEMin (Riel et al., 2022) that includes garnet
 756 (grt), cordierite (crd), ilmenite (ilm), two feldspars (fsp^2), quartz, (qz), spinel (spl), and silicate

757 melt (m) best matches the assemblage observed in the Site U1570 dacite. The pseudosection
 758 calculated with 1 wt% H₂O shows this assemblage being stable at 775–915 °C for pressures
 759 between 4.3–5.3 kbar (Fig. 12; Table 3). With 3 wt% H₂O, the range of temperatures over which
 760 the phase assemblage observed in our samples is stable is much narrower (750 °C and 790 °C)
 761 but the pressure range is expanded (3.7–5.2 kbar; Fig. S8; Table 3).



762
 763 **Figure 12.** Pseudosection calculated from the bulk rock composition acquired on the sample
 764 U1570-26R2-51-54 (Fig. 2D), assuming 1 wt % H₂O contents, using the MAGEMin
 765 thermodynamic package (Riel et al., 2022). Colored fields ranging from blue to yellow represent
 766 an increasing number of phases in an assemblage; the dark orange field represents the one that
 767 reproduces the phase assemblage observed in the U1570 dacite (grt-crd-ilm-fsp²-qz-spl-m).

768 Garnet stability field is represented by the solid red line, cordierite stability field with a black,
769 short-dashed line, quartz stability field with a black, long-dashed line, and biotite stability field
770 with a gray, dashed line. Phases include: andalusite (and), biotite (bt), cordierite (crd), ternary
771 feldspar (fsp, fsp² indicates two compositions of ternary feldspar), garnet (grt), hornblende (hbl),
772 ilmenite (ilm), muscovite (ms), olivine (ol), quartz (qz), sillimanite (sil), spinel (spl, spl²
773 indicates two compositions of spinel), silicate melt (m), and aqueous fluid (H₂O). Calculations
774 with 3 and 5wt% H₂O in the bulk compositions are shown in Fig. S8.

775 Finally, this phase assemblage is not reproduced in the pseudosection calculated with 5 wt%
776 H₂O. The pressure and temperature range where garnet, cordierite, and quartz overlap (~3.3 kbar,
777 ~675–700 °C) also includes amphibole in the assemblage and lacks the two feldspars (Fig. S8;
778 Table 3). In the following, we therefore assume that the bulk water content of the Site U1570
779 dacite is low (< 3 wt% H₂O).

780 **6 Discussion**

781 **6.1 Dacite origin and emplacement**

782 To understand the potential implications that dacite from Site U1570 may have for large
783 igneous province emplacement, continental breakup, and thermogenic gas production, it is
784 necessary to consider its petrogenesis. Below we discuss (1) the emplacement of the dacite using
785 textural observations and stratigraphic constraints and (2) the origin of the dacite with results
786 from thermodynamic calculations, thermobarometry, and phase composition analyses.

787 *6.1.1 Stratigraphic constraints*

788 The highly vesicular nature of the unit is consistent with emplacement of the dacite in a
789 shallow environment at low pressure (Fig. 4, 5). For evolved melts, highly vesicular pyroclasts
790 (>80% vesicularity) only form above water depths of 500 m, unless the dissolved magmatic
791 water content is unrealistically high (>7 wt%; Wright et al., 2003). Dacitic melts containing a
792 few wt% water may be expected to have viscosities of 10⁷ to 10⁶ Pa.s for pure melt between 750
793 and 850 °C (Giordano et al., 2008). Such melts would be prone to explosive fragmentation
794 during transport to the surface, with exsolution of the volatiles and formation of vesicles
795 resulting in increasing bulk viscosity and further promoting fragmentation. Such rapid
796 decompression and explosive fragmentation is also supported by the fragmentation of the large

797 mineral grains observed in the samples (Figs. 4,5; e.g., Best & Christiansen, 1997; Bindeman,
798 2015).

799 In order to assess the emplacement conditions of the dacite unit, several temperature and
800 chemical proxies were measured from the surrounding sediments. The total organic matter
801 (TOC) record shows no systematic loss of TOC in the sediments surrounding the dacitic unit.
802 Moreover, no significant variations in $\delta^{13}\text{C}_{\text{org}}$ (Fig. 2) occur near the dacite unit and no
803 appreciable organic-matter maturation appears to have taken place near the dacite based on the
804 RockEval parameters (Tmax), which would be expected even at relatively low temperature
805 (<100 °C; Svensen et al., 2023; Bédard et al., 2023). Analyses performed on sediment samples
806 from directly overtop the dacite in borehole U1570D show high Hg concentrations (~350 ppb),
807 however, these concentrations remain within the variability of the rest of the analyzed samples.
808 Svensen et al. (2023) demonstrated that thermal alteration in intrusive aureoles can result in a
809 loss of sedimentary Hg, likely because the most common sedimentary Hg phases (organic-matter
810 bound Hg and HgS) rapidly volatilize at temperatures of 200–300 °C, therefore, elevated Hg is
811 not consistent with thermal impacts from the dacite (e.g., Biester & Scholz, 1997).

812 Finally, samples were collected from directly at and within the mixing zone at the top and
813 base of the unit, as well as in a transect away from the contact and prepared for palynological
814 analyses. Samples in the mixing zone and adjacent to the dacite yielded abundant thermally
815 immature assemblages of acritarchs, dinocysts, pollen, and spores. These assemblages yielded a
816 Thermal Alteration Index (TAI; Gutjahr, 1966; Staplin, 1969) of 1+, values that were also
817 recorded from similar palynofloras 20 m further down the core. These TAI values are indicative
818 of temperatures of <45°C and confirm that the dacite did not thermally alter the sediments into
819 which it was recovered. This combined evidence from the sediments, along with the pyroclastic
820 nature of the unit, strongly argues against emplacement of the dacite as a liquid magmatic
821 intrusion. Although, emplacement as an invasive lava flow could explain the mingling textures
822 evident on the upper margin (Famelli et al., 2021), it is also inconsistent with the pyroclastic
823 nature of the dacite. The alternative, and preferred, interpretation is that the unit comprises a
824 deposit associated with an explosive pyroclastic density current (PDC; Branney et al., 2021).
825 Explosive pyroclastic density currents are known to be capable of flowing subaqueously if their
826 density exceeds that of the water in which they are erupted (Sparks et al., 1980a, b; White, 2000).
827 The flow of PDC's below water is in large part density-controlled, and, as such, the potential for

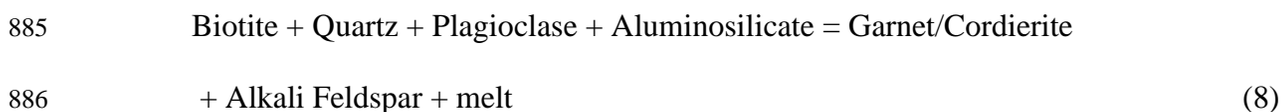
828 a PDC to flow into other fluids, such as muddy unconsolidated basin floor sediments, should be
829 equally controlled by density and other rheological properties, such as shear strength and
830 cohesion of water saturated sediments which will restrict the depth to which a PDC could
831 become invasive (e.g., Owen, 1987). The relatively sharp contact between the base of the dacite
832 and sediment below (Fig. 2E) with preserved sediment structures and local incorporation of
833 sediment clasts indicates some level of basal erosion and incorporation during the emplacement
834 of the dacite as a PDC (e.g., Roche et al., 2013). The emplacement dynamics of the dacite clearly
835 merit further study, however, we tentatively propose emplacement as a subaqueous PDC, which
836 exceeds the density of the uppermost layer of unconsolidated, mud-rich basin sediments,
837 promoting it to be partially invasive and resulting in local mingling with sediments.

838 *6.1.2 Thermobarometric and H₂O constraints*

839 Ideally, we would construct the pseudosections and perform thermobarometric
840 calculations using the composition of the protolith (e.g., Álvarez-Valero and Waters, 2010) from
841 which the dacite was derived to estimate the pressure and temperature conditions and water
842 contents of formation. In our case, however, we do not have access to the composition of the
843 protolith. Because the textural and chemical characteristics of garnet, cordierite and alkali
844 feldspar that these minerals are peritectic phases (see equation 9), they should be in equilibrium
845 with the melt. The anorthite (An) contents of the plagioclase are more variable (Fig. S5a). We
846 compare plagioclase found in our samples with plagioclase grains found in lavas from El
847 Hoyazo, Spain (Hiwatashi et al., 2021). The occurrence of high K-volcanics in this province can
848 be considered as an analog for the formation of the dacitic unit studied here (see below).
849 Plagioclases in the lavas from El Hoyazo show a large compositional spread with An-rich
850 plagioclase (>80%) showing resorbed or patchy textures, intermediate compositions showing
851 oscillatory zoning, and An-poor (<50%) compositions occurring in homogeneous subhedral
852 grains. Hiwatashi et al. (2021) interpret this variability as evidence for magma mixing between
853 high-SiO₂ dacite and andesite. Plagioclase observed in our samples overlap with the
854 compositions of the homogeneous grain interpreted as in equilibrium with a dacitic magma.
855 Hence, using the bulk rock composition, we consider that the pressure-temperature field that
856 reproduces the phase assemblage observed in our sample provides a good approximation for the
857 conditions of formation of the magma.

858 Partial melting of metapelitic rock during metamorphism commonly produces almandine
 859 garnet, which has commonly been interpreted as a high-pressure (>7 kbar) peritectic or liquidus
 860 phase in a hydrous, Al-rich magma (Green & Ringwood, 1968; Clemens & Wall, 1988; Harangi
 861 et al., 2001; Sieck et al., 2019). Green (1977), however, demonstrated that the grossular and
 862 spessartine content of almandine garnets are sensitive to pressure and that Ca and Mn-rich (> 4
 863 wt% CaO and MnO) almandine garnets could have crystallized at pressures less than 5 kbar.
 864 However, this cannot explain low-pressure signature recorded by the almandine garnet in our
 865 samples as they are Ca-poor (~1 wt% CaO) and Mn-poor (~1.8 wt% MnO). Aranovich and
 866 Podlesskii (1983) experimentally investigated the cordierite-garnet-sillimanite-quartz
 867 equilibrium at 4–8 kbar to determine the coexisting garnet and cordierite equilibrium
 868 compositions. At 4 kbar and 700°C, their experimental products have Fe-rich compositions
 869 consistent with natural samples from their study. They also showed that the Mg content of garnet
 870 and cordierite increases significantly with pressure from $Mg^{grt} = 0.10$ mol% and $Mg^{crd} = 0.44$
 871 mol% at 4 kbar to $Mg^{grt} = 0.51$ mol% and $Mg^{crd} = 0.89$ mol% at 8 kbar. Compared to the Mg
 872 content measured in garnet and cordierite grains in our study ($Mg^{grt} = 0.10 \pm 0.1$ mol% and Mg^{crd}
 873 $= 0.41 \pm 0.02$ mol%), their results support the relatively low equilibration pressure calculated for
 874 our samples. Additionally, Clemens and Wall (1981) observed garnet crystallization down to 1
 875 kbar for temperatures 700–900 °C in their experiments for water-undersaturated, peraluminous
 876 magmas, and in the last decade, several studies, through thermodynamic calculations and
 877 characterization of melt inclusions (e.g., Acosta-Vigil et al., 2010; Bartoli et al., 2013a, 2013b),
 878 have shown that peritectic garnet can also grow at low temperatures ($\leq 700^\circ\text{C}$) and low pressures
 879 (≤ 5 kbar).

880 Growth of peritectic garnet, cordierite, and alkali feldspar during anatexis of metapelites
 881 is thought to result from the breakdown of biotite during dehydration melting (e.g., Cesare et al.,
 882 2009; Ferrero et al., 2012; Stevens et al., 2007). Weinburg and Hasalová (2015) cite the
 883 following reaction in metapelites and variable pressure and intermediate temperature (700–
 884 800°C; Le Breton & Thompson, 1988; Spear, 1993):



887 in which cordierite results from the subsequent breakdown of garnet (Stevens et al., 1995). The
888 absence of biotite in our sample, the dissolution texture observed on the margins of plagioclase
889 grains, and the existence of a garnet inclusion in cordierite support the proposed reaction for
890 peritectic garnet growth (Eq. 8). Our results are consistent with biotite dehydration melting,
891 because its absence in our samples suggests that the magma formed at temperatures higher than
892 the field of stability of biotite (<350 °C for <3 wt% H₂O).

893 The phase assemblage observed in our samples is only reproduced for bulk water
894 contents ≤ 3wt%. A water-undersaturated composition is additionally supported by the presence
895 of graphite in the assemblage and anhydrous cordierite grains suggested by high analytical totals
896 (Table 1). In fact, for any H₂O-bearing phase, such as melt or cordierite, to coexist with graphite,
897 it must have an $a_{\text{H}_2\text{O}} < 1$ (Bartoli et al., 2013c). Although metapelitic anatexis at high temperature
898 (700–800°C) is facilitated by biotite dehydration melting and the release of water, experimental
899 studies by Holtz et al. (1992) show that the melt remains water-undersaturated. Additionally,
900 these water-fluxed, undersaturated melts are particularly buoyant, supporting rapid migration to
901 the surface (Holtz et al., 1992).

902 Using the pseudosection results for a magma with 1–3 wt% H₂O, the observed
903 phenocryst assemblage for our samples is stable at pressures of ~4–5 kbar and temperatures
904 higher than 750 °C, consistent with results obtained from thermobarometric calculations (Fig.
905 11). The pressures and temperatures recorded by the dacite phase assemblage are considered to
906 approximate the conditions of the last equilibrium state of the magma prior to rapid transport and
907 quenching. The limited compositional variability of the glass and mineral phases, as well as the
908 highly vesicular and glassy texture of the matrix, are interpreted to be consistent with rapid
909 emplacement and quenching. Therefore, we highlight that the recorded pressure-temperature
910 conditions may not represent the original melting conditions and the sample may have re-
911 equilibrated at depth along its ascent path (e.g., Álvarez-Valero & Waters, 2010).

912 We note that while crystallization and heat loss from the breakdown of garnet during the
913 decompression of magma should result in a lower temperature for Grt-Crd pairs than for Grt-Ilm
914 pairs (Sykes & Holloway, 1987), our samples record significantly higher temperatures for the
915 Grt-Crd pairs (Fig. 11). Hence, these results are consistent with a progressive melting process
916 happening at a relatively constant low pressure and variable high temperatures, with garnet-

917 ilmenite pairs recording the early stage of melting and garnet-cordierite pair the final stage of
918 melting. Therefore, we consider ~4.5 kbar, ~800 °C, and 1–3 wt% H₂O to be a good
919 approximation of the conditions for the average peak temperature during the formation of the
920 dacitic magma. Because seismic profiles of the Mimir High show the sediment cover is relatively
921 thin where the dacite was collected (Fig. 2), we can assume this pressure range is more likely to
922 result from generation at depth within the crust rather than from intrusion into the sediments
923 (Abdelmalak et al. 2017; Polteau et al., 2020).

924 *6.1.3 Age constraints*

925 McKenzie et al. (2018) suggest that the REE abundances in zircon preserve a record of
926 the composition of silica-rich parental melts since REE diffusion in zircon is slow and Zr is
927 abundant and incompatible in silica-rich melts. This means that variation in REE patterns, like
928 Eu anomalies, are representative of different parental magmas. Compared to population #1,
929 population #2 zircons show significantly more variable Eu anomalies (Fig. S9), indicating they
930 come from different sources and are therefore inherited rather than crystallized directly from the
931 dacite. The higher variability of population #2 is also observed on the shape components of the
932 trends (i.e., lambda coefficients; Anenburg & Williams, 2022), even when the Ce and Eu
933 anomalies are not included (Fig. S9C). The more restricted range of concentrations observed in
934 population #1 is consistent with crystallization from a single melt phase, likely during peritectic
935 growth of garnet and cordierite. The spatial distribution of these populations on CL images of the
936 individual zircons indicates that population #1 analyses are more often located on the rims of
937 grains, while population #2 analyses are either in or on the edge of darker cores (Fig. 9A). This
938 suggests that the population #1 analyses likely represent new growth from the dacite magma
939 reaching zircon saturation and that the calculated weighted average of these analyses (54.6 ± 1.1
940 Ma) is interpreted as the magmatic age. We note that as the analyzed zircons were found as
941 inclusions in garnet and cordierite, this age corresponds to the maximal age (not storage at depth)
942 for the eruption age.

943 The kernel density estimate (KDE) plot of the spot ages from population #2 shows zircon
944 as old as 2500 and 1700 Ma, with additional peaks at 100–400 Ma (Fig. S7A). This is in range of
945 what is expected for inherited grains on the Vøring Plateau, with possible contributions from
946 East Greenland sources (450–350 Ma, 2000–1700 Ma, and 3800–2500 Ma) and Norwegian

947 sources (450–350 Ma, 1250–900 Ma, and 1750–1500 Ma; Fonneland et al., 2004). However, it
948 should be noted that our calculations only include 11 analyses for population #2 and are
949 therefore unlikely to be a statistical representation of the entire spectrum of inheritance ages.

950 Zircon tends to incorporate HREEs relative to LREE and middle REEs (MREEs) from
951 their host melt (McKenzie et al., 2018). Chondrite-normalized (McDonough & Sun, 1995) REE
952 spectra for zircon measured in our samples are broadly consistent with this magmatic zircon
953 trend with steep positive slopes (Fig. S9A). There are a couple of analyses that have elevated
954 LREE concentrations. Because REE diffusion in zircon is slow, this may be the result of
955 overlapping inclusions during analysis or can be an effect of hydrothermal alteration as an influx
956 of LREE (Hoskin, 2005; McKenzie et al., 2018). The trace element concentrations for these
957 analyses do not suggest analytical overlap with other minerals and both are in older population
958 #2 grains. Therefore, we suggest their LREE concentration results from hydrothermal alteration,
959 and neither of these zircon grains are used in further interpretations.

960 6.2 Regional implications

961 6.2.1 *The evolution of rifting on the Vøring Plateau*

962 Crustal anatexis at magma-rich margins during rifting is often considered to be a lower
963 crustal process, facilitated by a mafic magmatic underplate. Crustal magmas are then transported
964 to an upper crustal reservoir or to the surface with varying degrees of re-equilibration and crustal
965 assimilation during transport (e.g., Vissers et al., 1995; Pederson et al., 1998 and references
966 therein; Platt et al., 1998; Duggen et al. 2004; Álvarez-Valero & Kriegsman, 2007; Thybo &
967 Artemieva, 2013 and references therein; Neumann et al., 2013). Dacite generation through
968 underplating has been described by Luo et al. (2018) as a result of crustal anatexis from heat
969 provided by mantle-derived basaltic magmas. The basaltic magmas mix with the crustal melts to
970 produce andesitic magmas, which can evolve to strongly peraluminous dacitic magmas by
971 stalling, fractionating, and the assimilation of anatectic melts of surrounding metasedimentary
972 rocks. The presence of underplated material at the base of the crust is often supported by seismic
973 reflection images of thick, lower crustal bodies with high velocities ($V_P > 7$ km) that are
974 characteristic of magma-rich margins (Kelemen and Hobbrook; 1995; White et al., 1987, 2008;
975 Gac et al., 2022). In the Jameson Land Basin in East Greenland and in the Arctic Basin of North

976 Greenland, the underplated material has also been proposed as a source of melt generation
977 (Thorarinsson et al., 2011; Eide et al., 2022).

978 Our thermobarometric calculations suggest, however, that the dacite collected at site
979 U1570 was not produced in the lower crust but was instead generated in the upper crust (< 18 km
980 depth assuming a continental crust density of 2.8 g/cm³). The bulk trace elements composition of
981 the samples (Fig. 3B) also supports an upper crust origin. The enrichment in Cs, in particular, has
982 been proposed as a discriminating factor between the upper and the lower crust (Meyers et al.,
983 2009). No evidence of deeper crustal processes is observed. This may be explained by chemical
984 re-equilibration in an upper crustal reservoir of a magma formed at depth. For instance, this
985 scenario has been favored to explain the formation of the high-K volcanic lavas in the NVP, SE
986 Spain. Thermobarometric calculations performed on enclaves of the high-K volcanic rocks from
987 the El Hoyazo and Mazarrón volcanoes suggest melting pressures of 7 kbar (Cesare et al., 1997;
988 Cesare & Gómez-Pugnaire, 2001) and 4 kbar (Cesare et al., 2003), respectively. The difference
989 in pressure recorded at the two volcanoes is attributed to varying depths of magma stagnation in
990 the crust. The dacite from our study presents mineral phase compositions that match the mineral
991 phases of the NVP high-K volcanic rocks (Figs. 6, 7, 8) and records pressure and temperature
992 conditions of anatexis similar to those reported for the Mazarrón volcano, consistent with an
993 upper crustal origin. Additionally, in comparison to the NVP, the glass composition (Fig. 3) in
994 the Site U1570 dacite is more homogenous, which suggests that magma storage for our samples
995 was long enough to approach chemical equilibrium. However, temperature recorded by the
996 garnet-ilmenite and the garnet-cordierite pairs suggest they formed as a result of heating at a
997 constant pressure rather than through decompression processes as we would expect for migration
998 from the lower to upper crust. We propose the Site U1570 dacite is the product of low pressure
999 (<5 kbar) partial melting of crustal metapelites on a thinned lithosphere, consistent with previous
1000 assumptions on the origin of silicic magmatism in the NE Atlantic (e.g., Abdelmalak et al., 2016;
1001 Eldhom et al., 1987; Sinton et al., 1998; Viereck et al., 1988) as the result of crustal anatexis at
1002 low pressure. This scenario is also consistent with a study of the Isle of Rum, in which
1003 rhyodacites were interpreted to be produced by melting amphibolite rather than deep crustal
1004 granulite (Meyer et al., 2009b), and with ODP Site 642E dacite trace element data that are
1005 compatible with melting Caledonian metasedimentary rocks (Viereck et al., 1988). Our results
1006 also provide evidence for

1007 Previously published Ar-Ar dating on plagioclase from the dacite collected by ODP in
1008 hole 642E did not provide a precise age of eruption of the silicic magmatism (57–51.5 Ma;
1009 Sinton et al., 1998). It is, however, consistent with the age obtained on dacitic lava recovered
1010 from the North Rockall Trough (Darwin Complex; Fig. 3; Morton et al., 1988a) indicating an age
1011 ~55 Ma (Sinton et al., 1998 and references therein) and both are coincident with the magmatic
1012 age reported in this study. Importantly, the weighted mean U-Pb date interpreted as the age of
1013 zircon growth in this study (54.6 ± 1.1 Ma) postdate the age of the PETM (56–55.8 Ma; Jones et
1014 al., 2023).

1015 Dacite recovered at ODP Site 642E is thought to have been emplaced before the peak of
1016 magmatism (Abdelmalak et al., 2017), and stratigraphic relationships indicate that the basalt
1017 overlying the silicic lavas from the North Rockall Trough and the Vøring Plateau erupted shortly
1018 after the silicic magmas (Sinton et al., 1998 and references therein). This resulted in
1019 interpretations suggesting the end of silicic magmatism is coeval with the onset of the main
1020 basaltic phase, potentially due to a change of extension rate (e.g. Abdelmalak et al., 2016; Sinton
1021 et al., 1998). Interestingly, the dacite recovered at Site U1570 is not overlain by basaltic
1022 magmatism but was rather emplaced at the surface in a shallow marine setting. At Mimir High,
1023 the overall palynofacies of the sediments is dominated by terrestrial elements (pollen, spores, and
1024 phytodebris) in addition to marine elements (dinoflagellate cysts), suggesting that the
1025 depositional environment was coastal for most of the Paleocene and Eocene (Planke et al., 2023).
1026 This demonstrates that the coastal environment was sustained despite being an active depocenter,
1027 consistent with substantial subsidence of the continental crust during extension. The
1028 emplacement of the dacite within these sediments indicates this extension remained active during
1029 and after the emplacement of this dacite.

1030 Several studies (e.g., Peron-Pinvidic et al., 2013, Tugend et al., 2018; Gillard et al., 2019;
1031 Ferrand, 2020) have proposed that magma-rich rift margins may go through a period of
1032 hyperextension, suggesting that extreme crustal thinning does not preclude the emplacement of a
1033 high volume of magmatism. Tugend et al. (2018) suggested that when the distinction between
1034 magma-poor and magma-rich margins reflect the magmatic productivity, the overall evolution
1035 before breakup may only reflect the timing of decompression melting relative to thinning of the
1036 margin. In this context, Ferrand (2020) proposed that the Vøring margin was initiated as a
1037 magma-poor margin and was subsequently affected by hotspot-induced magmatic processes and

1038 overprinted by volcanic activity. Evidence for silicic magmatism in the Northeast Atlantis as old
1039 as ~62 My ago, with tuff interlayers between basalt flows on the Isle of Muck (Emeleus et al.,
1040 1996) dated at 62.8–62.4 Ma (Pearson et al., 1996) and ash layers in sediments of the North Sea
1041 (Knox & Morton, 1998; Morton et al., 1998 that indicate mixing between silicic and basaltic
1042 magmatism dated at 62–60 Ma. Together, these ages suggest a longer period of crustal anatexis
1043 off the coast of Norway during rifting of the Northeast Atlantic. Additionally, the occurrence of
1044 dacitic magmatism on the mid-Norwegian margin, either overlain by subaerial magmatism (ODP
1045 hole 642E and the North Rockall Trough) or emplaced in a sediment of an early Eocene age
1046 (IODP Expedition 396, Site U1570), indicates that magmatism not only occurred at the zone of
1047 plate separation, but also landward of the ocean-continent transition on the European side. We
1048 propose that the pre-breakup phase during the development of the Vøring margin was associated
1049 with significant continental lithospheric extension. This extension resulted in low-pressure
1050 melting of metasedimentary rock to generate silicic magmas that erupted later.

1051 *6.1.2 Implications for the Paleocene-Eocene carbon cycle perturbations*

1052 Excess magmatism, thermogenic gas release and sediment melting from contact
1053 metamorphism around sills have been invoked to explain the carbon cycle perturbations that
1054 occur throughout the late Paleocene and early Eocene, most notably the PETM, the largest of a
1055 series of global warming episodes (e.g., Svenson et al., 2004; Westerhold et al., 2020). The
1056 PETM was marked by rapid surface warming (~5 °C), ocean acidification, and the release of
1057 ¹³C-depleted carbon into the oceans and atmosphere, evidenced by a sharp decrease in
1058 sedimentary $\delta^{13}\text{C}$, a negative Carbon Isotope Excursion (CIE; Kennet & Scott, 1991; Zeebe et
1059 al., 2009). Estimations provided by mass balance calculations suggest the PETM CIE was the
1060 result of the injection of 3,000–14,900 gigatons (Gt) of ¹³C-depleted carbon (Zeebe et al., 2009;
1061 Gutjahr et al., 2017; Haynes and Hönisch 2020) most likely over a geologically rapid (few
1062 thousand years) time frame (Zeebe et al., 2016; Kirtland-Turner & Ridgeway, 2016). While
1063 sedimentary proxy and stratigraphic evidence support a contribution from volcanic outgassing
1064 (e.g., Frieling et al., 2016; Jones et al., 2019; Kender et al., 2021), it remains unclear whether the
1065 volcanic source would provide the timing, composition, magnitude, and rate of outgassing
1066 required to generate the CIE (Haynes & Hönisch, 2020; Gutjahr et al., 2017). It has been
1067 proposed that mafic intrusions related to the excess magmatism of the mid-Norwegian margin

1068 resulted in hydrothermal vent complex formation and direct emission of CO₂ into the atmosphere
1069 (e.g., Gernon et al., 2022) or contact metamorphism of organic-rich ocean sediments and
1070 subsequent explosive release of large amounts of methane with a ¹³C-depleted isotope signature
1071 (δ¹³C; e.g., Svensen et al., 2004; Frieling et al., 2016). Rampino (2013) further suggested that the
1072 NAIP emplacement may have contributed more thermogenic gases by anatexis melting of
1073 organic-rich sediments rather than pure contact metamorphism. The authors proposed that dacitic
1074 flows observed throughout the Norwegian continental margin may be the direct product of this
1075 process. Our dacitic sample from the Mimir High does not support Rampino (2013)'s hypothesis.
1076 In fact, the sediments in contact with the dacite indicate no thermal alteration, which suggests
1077 that its emplacement did not mobilize or volatilize significant amounts of carbon from
1078 surrounding sediments. While the graphite film observed on some grains suggests that minor
1079 amounts of fluid-deposited carbon may be present (e.g., Luque et al., 1993), the presence of
1080 graphite grains (as inclusion or disseminated in the sample) is consistent with a restitic phase that
1081 has been entrained in the melt (Zeck 1970, 1992). Hence, the presence of graphite is likely
1082 inherent to the nature of protolith of our samples, a graphitic metapelite, rather than a product of
1083 melting or burning organic matter.

1084 However, this does not imply the formation of the Site U1570 dacite did not release CO₂.
1085 Cesare et al. (2005) showed that in relatively reducing conditions (as in graphitic metapelites),
1086 biotite is the only source of Fe³⁺, with the melting reaction of biotite is important to iron
1087 reduction. As the melt produced during the reaction is relatively iron-poor (Table 1), biotite
1088 breakdown is accompanied by carbon oxidation and the production of CO₂. The amount of CO₂
1089 produced in this way, however, is likely too small to be a significant contribution to the PETM.
1090 Additionally, the zircon inclusions record post-PETM U-Pb magmatic ages. Sill intrusions and
1091 hydrothermal vent complexes can release large volatile concentrations into the atmosphere and
1092 may have contributed to the onset and longevity of the PETM (Frieling et al., 2016; Berndt et al.
1093 2023), but the dacite sample outlined by this study is not an example of this. In fact, the long
1094 period of silicic magmatism (~62–54.6 Ma) in the Northeast Atlantic highlighted here warrants
1095 caution in including NAIP-related thermogenic carbon released from eruptions into wet
1096 sediments in carbon emission calculations.

1097 **5 Conclusions**

1098 The garnet and cordierite-bearing dacite sampled during IODP Expedition 396 in
1099 boreholes U1570A and U1570D was likely emplaced as a high-density pyroclastic current into
1100 wet, unconsolidated sediments in a shallow coastal environment after the onset of the PETM.
1101 This is evidenced by the folded, elongated glassy pyroclastic textures, local mingling with
1102 sediment and preserved sediment clasts, and highly fragmented phenocrysts. The chemical
1103 signatures of the peritectic garnet and cordierite grains suggest an upper crustal, metapelitic
1104 protolith. This is further supported by the inheritance patterns observed in the zircon U-Pb ages
1105 that match inherited grains on the Vøring Plateau. Our thermodynamic results suggest the
1106 magma is water undersaturated (≤ 3 wt% H₂O) and indicate low pressure (<5 kbar), high
1107 temperature (~800 °C) anatexis in the upper continental crust.

1108 We propose this dacite is a crustal anatectic product of an intense and long (> 7 Myr)
1109 phase of continental stretching that preceded the main phase of basaltic magmatism
1110 emplacement, that was later ($< 54.6 \pm 1.1$ Ma) emplaced on top of unconsolidated wet sediments.
1111 Implications of lithospheric thinning in excess-magmatic margins should be considered in
1112 tectonic reconstructions for the evolution of the North Atlantic rift margin.

1113

1114 **Data Availability Statement:** All the data are shared in tables in the main text and as
1115 supplementary tables attached with this manuscript and will be archived on Zenodo upon
1116 acceptance.

1117

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1129

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