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Evidence for polybaric fractional crystallization in a continental arc: Hidden Lakes mafic complex, Sierra Nevada batholith, California

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Evidence for polybaric fractional crystallization in a continental arc: Hidden Lakes 1 mafic complex, Sierra Nevada batholith, California 2 3 Madeline J. Lewis^{*1}, Claire E. Bucholz¹, Oliver E. Jagoutz² 4 5 1. Division of Geological and Planetary Sciences, California Institute of 6 Technology 7 2. Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts 8 Institute of Technology 9 10 *Correspoding Author: Madeline J. Lewis, maddie@purdue.edu 11 12 13 ABSTRACT 14 15 Although the voluminous granitoids that constitute the upper crust of the 16 17 Sierra Nevada batholith (California) have been investigated in detail, comparatively few studies focus on the origin of mafic bodies at similar crustal 18 levels. Here, we present field and petrographic observations, geochronology, and 19 geochemistry of the Hidden Lakes mafic complex in the central-eastern Sierra 20 Nevada batholith. Our results show that the complex comprises norites, gabbros, 21 22 monzondiorites, and monzonites that record fractional crystallization of a hydrous 23 (~3 wt.% H₂O), non-primitive basalt within the upper crust (0.3 GPa) at c. 95-96 Ma. To quantitatively model the generation of the observed lithologies, we 24 25 construct a two-stage polybaric crystallization model based on cumulate and meltlike bulk-rock compositions. In the first step, we model fractionation of a 26 27 primitive, mantle-derived basalt at >30 km depth, generating dominantly

28 pyroxenite cumulates. The evolution of the derivative melt (67% of melt mass remaining) is then modeled to fractionate at 12 km depth to produce the observed 29 lithologies within the Hidden Lakes mafic complex. Extension of this model to 30 higher-silica melt compositions (>65 wt.% SiO₂) replicates observed granodiorite 31 32 compositions in the batholith, suggesting that polybaric crystallization could be an important process for the generation of arc granitoid melts. The depth of 33 differentiation in continental arcs is debated, as field observations indicate 34 abundant lower crustal fractionation while experimental data suggests that high-35 pressure crystallization of hydrous basalts cannot produce the non-peraluminous 36 granitoid compositions observed in continental arc batholiths. Our model supports 37 polybaric differentiation as one potential mechanism to resolve this inconsistency. 38

39

40 Introduction

41 The generation of batholith-scale granodiorite intrusions in continental arc settings is an unresolved problem, as differentiation mechanisms produce only 42 small volumes of granitic melt (~10-20% of original melt mass) in comparison to 43 much greater volumes of ultramafic and mafic cumulates or restites (Kay and Kay 44 1985; Soesoo 2000; Jagoutz and Klein 2018). However, lower crustal mafic and 45 ultramafic cumulates are ubiquitous in magmatic arcs exhumed to 30 to 55 km 46 depths (Debari and Sleep 1991; Greene et al. 2006; Jagoutz et al. 2009; Jagoutz 47 2010; Otamendi et al. 2012; Jagoutz 2014; Walker et al. 2015; Guo et al. 2020) and 48

| 49 | may be brought to the surface as xenoliths in active volcanic arcs (Debari et al. |
|----|--|
| 50 | 1987; Arai and Ishimaru 2008). Therefore, magmatic differentiation by lower |
| 51 | crustal fractional crystallization of hydrous basalts is considered an important |
| 52 | process in the formation of granodiorites and granites characteristic of the upper |
| 53 | crust (Greene et al. 2006; Otamendi et al. 2012; Jagoutz 2014; Walker et al. 2015; |
| 54 | Guo et al. 2020). Experimental studies (Müntener et al. 2001; Sisson et al. 2005; |
| 55 | Nandedkar et al. 2014; Müntener and Ulmer 2018; Ulmer et al. 2018) and |
| 56 | thermodynamic modeling (Annen et al. 2006; Jagoutz and Klein 2018) also |
| 57 | support the hypothesis that deep (>0.7 GPa) differentiation of hydrous basalts can |
| 58 | produce high-silica melts. |

However, detailed experimental high-pressure crystallization studies suggest 59 that differentiation in the deep crust cannot be solely responsible for the 60 compositional stratification observed in arcs. Crystallization of hydrous primitive 61 basalts in the lower crust (i.e. >0.7 GPa) produces peraluminous andesites, which 62 are rarely observed in arc batholiths (Cawthorn and Brown 1976; Cawthorn and 63 O'Hara 1976; Blatter et al. 2013; Nandedkar et al. 2014; Melekhova et al. 2015; 64 Müntener and Ulmer 2018; Ulmer et al. 2018). Though the experimental basalts 65 used in some of these studies are Al-enriched, their starting compositions are 66 within the range of observed primitive arc basalts, as compiled by Schmidt and 67 Jagoutz (2017). At lower crustal pressures and high (≥ 2 wt.%) water contents, 68

suppression of plagioclase and abundant early clinopyroxene crystallization drives 69 residual melts towards peraluminous compositions (i.e., ASI > 1.1 where ASI =70 molar Al/(Ca-1.67×P+Na+K). In contrast, at lower pressures (<0.7 GPa) and/or 71 72 lower water contents plagioclase and orthopyroxene (in the case of lower water activities) will saturate in a basalt at higher temperatures, limiting the increases in 73 ASI in a crystallizing melt. As supported by high-pressure experimental studies, 74 differentiation in arcs is not a simple story of high-pressure differentiation. 75 Thus, there are two observations that must be reconciled: (1) lower crustal 76 arc sections dominated by ultramafic to mafic cumulates indicating the importance 77 of deep crustal fractional crystallization and (2) experimental crystallization studies 78 suggesting that although high-silica melts can be produced at these pressures, they 79 will become peraluminous at relatively low SiO_2 contents (<60 wt.%), a feature 80 rarely observed in arc batholiths. Three processes have been proposed to explain 81 this apparent contradiction: (1) polybaric differentiation (Hamada et al. 2014; 82 Melekhova et al. 2015), (2) mixing between low-Mg basaltic and granitic melts 83 (Sisson et al. 1996; Grove et al. 2003; Sisson et al. 2005; Blatter et al. 2013; 84 Müntener and Ulmer 2018), or (3) mixing of dry and wet magmas to produce 85 "damp" (<2 wt.% H₂O) basalts to andesites that then further differentiate in the 86 lower crust (Rezeau et al. 2021) (e.g., Sisson et al. 2005; Müntener and Ulmer 87 88 2018). All three of these processes require the presence of evolved (low-MgO)

89 basalts produced through lower crustal differentiation of more primitive, high-MgO basalts. However, each is not without its shortcomings. In scenario (1), the 90 low-MgO basalts can ascend into the upper crust to differentiate further, producing 91 92 non-peraluminous silicic melts. The absence of significant volumetrically significant mafic plutons in the upper crust of exposed arcs, however, suggests that 93 this may be relatively minor process. In scenario (2), the low-MgO basalts mix 94 with higher-silica melts (formed through fractional crystallization, partial melting, 95 or both) to produce granodioritic magmas. However, mixing of two magmas with 96 strongly contrasting viscosities is mechanically unviable (Sparks and Marshall 97 1986). Last, in scenario (3), the low-MgO basalts with variable water contents mix 98 to produce a damp (<2 wt.% H₂O) basalt that then differentiates in the lower crust 99 without significant alumina enrichment due to early orthopyroxene and plagioclase 100 crystallization. However, although crystallization of damp basalts in the lower 101 crust can ameliorate issues with ASI increases during differentiation, this model 102 103 cannot explain other features of arc batholiths. For example, dry to damp differentiation of basalts will follow a tholeiitic trend (increasing FeO/MgO with 104 SiO_2) which is not observed in the calc-alkaline batholiths of continental arcs (e.g. 105 106 Villiger et al. 2004; Melekhova et al. 2015; Müntener and Ulmer 2018). 107 To test whether these processes play an important role in the generation of 108 arc batholiths, this study focuses on an upper crustal mafic complex in a

109 continental arc batholith. Although less common than voluminous silicic plutons, upper crustal mafic intrusions potentially preserve information on the nature and 110 origin of low-MgO basalts in the formation of arc batholiths. Here, we undertake a 111 112 case study of the Hidden Lakes mafic complex (HLMC), an exposure of norite through monzonite emplaced into the upper crust of the Sierra Nevada batholith. 113 We present detailed field mapping to document intrusive relationships and the 114 range of mafic lithologies, and U-Pb zircon geochronology to determine the timing 115 of crystallization of the HLMC relative to surrounding high-silica intrusions. Using 116 bulk-rock and mineral chemistry data, as well as comparisons to experimental 117 studies, we aim to constrain crystallization pressures, temperatures, H₂O content, 118 and oxygen fugacity (fO_2) of the crystallizing magma. As field, petrographic, and 119 geochemical data suggest a fractional crystallization origin for the HLMC, we 120 construct a two-stage polybaric fractionational crystallization model to replicate 121 the composition of the evolving melt. In light of our findings, we discuss the 122 123 importance of evolved basalts in the formation of high-silica arc batholiths.

124

125 Geologic Setting

The HLMC is located in the eastern central Sierra Nevada batholith (SNB),
the largely granodioritic intrusive product of the Sierra Nevada paleo-continental
arc. From the late-Triassic through late-Cretaceous, magmatism related to

6

| 129 | subduction of the Farallon plate beneath North America produced felsic intrusions |
|-----|---|
| 130 | that now form the bulk of the batholith. Magmatism ceased by 75 Ma due to |
| 131 | shallowing of the subduction angle (English et al. 2003; Kirsch et al 2016). |
| 132 | Presently, most of the SNB is exposed to upper crustal depths (0.2 to 0.4 GPa), |
| 133 | with the exception of the southernmost lower crustal section which extends to 1.0 |
| 134 | GPa pressures (Ague and Brimhall 1988; Pickett and Saleeby 1993). The late- |
| 135 | Jurassic (145-160 Ma) and late-Cretaceous (85-100 Ma) periods in the SNB are |
| 136 | characterized by enhanced magmatic activity, during which of 85% of the batholith |
| 137 | volume was generated (DeCelles et al. 2009; Ducea et al. 2015; Kirsch et al. 2016). |
| 138 | Numerous mafic bodies throughout the SNB upper crust were noted during |
| 139 | early studies of the batholith (Mayo 1941; Bateman and Eaton 1967; Frost 1987), |
| 140 | but have been less comprehensively studied as compared to the felsic batholith due |
| 141 | to their relatively small areal fraction of the exposed SNB. Geochronological |
| 142 | studies on a few of the mafic bodies demonstrate that they are coeval with the |
| 143 | surrounding granitoids and therefore may play a critical role in the formation of the |
| 144 | granitoid batholith (Coleman et al. 1995; Ratajeski et al. 2001). Several mafic |
| 145 | bodies have been investigated in previous studies (Frost 1987; Frost and Mahood |
| 146 | 1987; Coleman et al. 1995; Sisson et al. 1996; Ratajeski et al. 2001; McCarthy and |
| 147 | Müntener 2016), all of which document the presence of hydrous, low-Mg basalts |
| 148 | in the upper crust of the SNB. |

149

150 Field Relationships

| 151 | The Hidden Lakes mafic complex (HLMC) is exposed in the central SNB |
|-----|--|
| 152 | near Bishop, CA, and covers an area of 2.5 km ² . Our field mapping (Fig 1) refines |
| 153 | the outer margins of the mafic complex where it is in contact with surrounding |
| 154 | granodiorite plutons, as previously mapped by Bateman (1992), and displays the |
| 155 | variation in rock types and the nature of contacts within the mafic complex. |
| 156 | Amphibole- and biotite-bearing medium- to fine-grained gabbros are the dominant |
| 157 | lithology throughout HLMC. Norites occur as small pods within the gabbro in the |
| 158 | eastern section of HLMC. In the northeastern section of HLMC, near its contact |
| 159 | with the Round Valley Peak granodiorite, lithologies transition gradationally from |
| 160 | gabbro to monzodiorite to monzonite toward the northern edge of the mafic |
| 161 | complex. The field occurrences and characteristics of each rock type in the mapped |
| 162 | area are described in detail below. |
| 163 | |

163

164 Norites

Norites are the most primitive lithology found in HLMC and are present as small, rounded pods (up to 3 m) within the gabbro unit in the eastern section of the complex (Fig S2a). Their volumetric abundance within the gabbro is difficult to ascertain due to exposure on steep cliff faces, however, they weather out and occur

as large boulders in the scree fields to the west of the cliffs. Norite boulders are

170 significantly less abundant in scree fields than gabbro boulders, so it is presumed

171 that norite is less common than gabbro in the cliff exposures.

172

173 Anorthosites

Anorthosite was observed only in rare boulders below gabbro-dominated cliffs in the southeastern section of HLMC (Fig 1). The anorthosite is medium-fine grained (0.5-2 mm), isotropic, and contains 90 to 95% dark grey plagioclase with 5 to 10% biotite.

178

179 Gabbros

Amphibole-biotite gabbro is the volumetrically dominant rock type in HLMC and is present in most areas of the mafic complex, except for the northern margin characterized by more evolved lithologies (Fig 1). The gabbros contain abundant plagioclase and amphibole, along with common clinopyroxene and biotite, and minor orthopyroxene. Based on their macroscopic textures, the gabbros are divided into three main groups: (1) medium-fine grained gabbro, (2) coarsegrained gabbro, and (3) "spotted" gabbro (Fig S2).

187 Medium-fine grained gabbro is the most common rock type in HLMC and 188 contains dark gray plagioclase, amphibole, and pyroxene (1-3 mm) with evenly

| 189 | distributed finer-grained biotite (~1 mm). The coarse-grained gabbro is present in |
|-----|--|
| 190 | elongate bodies within the medium-fine grained gabbro (Fig 1). The mineral |
| 191 | assemblage in the coarse-grained gabbro is identical to the medium-fine grained |
| 192 | gabbro, with slightly less plagioclase. The "spotted" gabbro is characterized by 1 |
| 193 | to 2 cm clots of poikilitic amphibole+biotite enclosing pyroxene and minor |
| 194 | plagioclase, which give it a spotted surface appearance (Fig S2). Clots are enclosed |
| 195 | in a finer-grained matrix that is mineralogically very similar to the medium-fine |
| 196 | grained gabbro. |
| 197 | Contacts between the different textural gabbroic units are gradational over 1 |
| 198 | to 5 m, or can exhibit mingling textures. Elongate intrusion geometry is evident in |
| 199 | the western and southern areas of HLMC, while the eastern section appears to be a |
| 200 | single large intrusion composed mainly of medium-fine grained gabbro that grades |
| 201 | into monzodiorite to the northeast. |
| 202 | |
| 203 | Monzodiorites and Monzonites |
| 204 | In the northeastern section of HLMC, two bodies of monzodiorite and two |
| 205 | elongate bodies of monzonite are in largely gradational contact with one another |

206 over >5 m. Mingling textures between the monzonite and monzodiorite, as well as

207 dikes of monzonite intruding into monzodiorite, and vice versa, are present. Thus,

these are synmagmatic lithologies that were mobilized shortly after crystallizationof the northeastern section of gabbro.

The monzodiorites are medium-fine grained, dominated by plagioclase, and contain biotite, K-feldspar, amphibole, and interstitial quartz in order of decreasing abundance. Rare clinopyroxene is present in the cores of amphibole grains. The monzonites are typically medium to coarse grained, dominated by plagioclase and clusters of biotite + amphibole. K-feldspar is more common than in the monzodiorites, and quartz is typically minor and interstitial.

217 Diabase Bodies

Generally small, irregularly shaped bodies of diabase, as well as diabase dikes, are present within and near HLMC. Intrusions are fine grained and occasionally contain blocks of HLMC gabbro. Contacts between the diabase and the mafic complex are sharp, in contrast with the generally gradational contacts between the coarser-grained lithologies, suggesting that the diabase is significantly younger than much of HLMC.

224

225 Surrounding Felsic Plutons and Outer Contacts

Three felsic intrusions are in contact with the HLMC around its perimeter: the granite of Chickenfoot Lake to the west and south, the Lake Edison

granodiorite to the east, and the Round Valley Peak granodiorite (Krv) to the north(Fig 1).

| 230 | The granite of Chickenfoot Lake (Jcf, 191.6 ± 1.0 Ma, this study) is one of |
|-----|--|
| 231 | few Jurassic-aged plutons in the central SNB. Where observed, the contact |
| 232 | between Jcf and HLMC is a 0.1 to 1 m wide gradational zone of dioritic |
| 233 | composition that occasionally includes coarse orthoclase xenocrysts, which are not |
| 234 | present in the adjacent gabbros and so presumably originated as crystals in Jcf (Fig |
| 235 | S3b). Diorite dikes originating from HLMC inrtude into the Jcf. The Lake Edison |
| 236 | granodiorite (Kle, 90.9 ± 0.1 Ma; Lackey et al. 2008) intrudes HLMC on its |
| 237 | eastern side, creating a subvertical and sharp contact with Kle truncating layering |
| 238 | in HLMC. The Round Valley Peak granodiorite (Krv) intruded at 88.8 ±0.02 Ma |
| 239 | (Lackey et al. 2008) and forms a complicated contact zone (the Heterogeneous |
| 240 | Intrusive Zone) where it intrudes the northeastern margin of HLMC. |

241

242 Heterogeneous Intrusive Zone

The east-west trending contact between Krv and HLMC is a ~150 m wide contact zone, which we term the Heterogeneous Intrusive Zone (HIZ). This zone consists of a magmatic breccia composed of 50 to 85% Krv granodiorite that has entrained angular to subangular blocks of gabbro, monzodiorite, and monzonite. Mafic blocks vary in size (5 cm-10 m) and angularity with the most angular blocks

| 248 | present on the southern margin of HIZ. Blocks become progressively more |
|-----|---|
| 249 | rounded and partially digested towards the north (Fig S3c). Meter to sub-meter |
| 250 | scale dikes of the Krv intrude the HLMC immediately south of the HIZ, and |
| 251 | sometimes contain angular xenoliths of gabbro, monzodiorite, and monzonite from |
| 252 | the HLMC (Fig S3d). |
| 253 | |
| 254 | Petrography |
| 255 | Descriptions of 28 representative samples in thin section are given below, |
| 256 | with at least three sections from each rock type. Modal percent of major minerals |
| 257 | were estimated visually (Table 1). |
| 258 | |
| 259 | Norites |
| 260 | The norites consist of orthopyroxene (15-75%), plagioclase (15-40%), |
| 261 | amphibole (0-20%), biotite (5-15%), clinopyroxene (1-5%) and olivine (<1%), |
| 262 | along with minor Fe-Ti oxides and accessory apatite and zircon. Norite samples |
| 263 | exhibit ad- and mesocumulate textures with rounded orthopyroxene grains that |
| 264 | range in size from 50 to 600 μ m (Fig 2a). All samples contain interstitial |
| 265 | plagioclase and few contain euhedral plagioclase crystals ranging from 0.2 to 3 |
| 266 | mm. Amphibole occurs either as large oikocrysts (1 cm) enveloping orthopyroxene |
| 267 | and plagioclase \pm olivine (Fig 2b), or as overgrowths on orthopyroxenes and |
| | |

| 268 | clinopyroxenes (Fig S4a). Late crystallizing biotite (0.4-2 mm) fills interstit | ial |
|-----|---|------|
| 269 | spaces and surrounds rounded orthopyroxenes. Olivine was observed in two |) |
| 270 | samples (0.2-1 mm). Minor Fe- and Fe-Ti-oxides are included in orthopyro | xene |
| 271 | and plagioclase. | |

- 272
- 273 Gabbros

The relative proportions of mineral phases in the gabbros are variable, with 274 the following ranges: plagioclase 45 to 75%, amphibole 5 to 50%, clinopyroxene 5 275 to 20%, biotite 5 to 10%, and orthopyroxene 0 to 5%. Gabbros with both 276 cumulative and non-cumulative textures are present in HLMC (Fig 2c,d). Coarse-277 grained and spotted gabbro samples, as well as some of the medium-fine grained 278 gabbros, preserve textural evidence for accumulation of plagioclase and 279 clinopyroxene. These meso- to adcumulate samples are dominated by equant, 280 nearly euhedral plagioclase (0.5-4 mm) and rounded clinopyroxene (0.5-3 mm) 281 282 crystals. Orthopyroxene is far less prevalent than clinopyroxene, and both pyroxenes commonly contain exsolution lamellae. Amphibole is present as a 283 primary magmatic phase, secondary magmatic rims nucleated on pyroxenes, and as 284 psuedomorphs after pyroxene (Fig 2c, Fig S4b). Biotite is a late crystallizing 285 286 interstitial phase containing inclusions of clinopyroxene, plagioclase, and Fe-Ti-287 oxides. Small, rare alkali feldspars are present interstitially.

289 Monzodiorites and Monzonites

| 290 | The monzodiorites and monzonites of HLMC are distinguished from the |
|-----|---|
| 291 | gabbros by a decreased abundance of pyroxene, though many monzodiorites |
| 292 | contain pseudomorphs of pyroxene altered to amphibole. These pseudomorphs |
| 293 | occasionally have ortho- and clinopyroxene cores. The monzodiorites consist of 60 |
| 294 | to 70% plagioclase, 15 to 20% biotite, 10 to 15% K-feldspar, 5 to 15% amphibole, |
| 295 | <2% quartz, and minor ortho- and clinopyroxene (0- 2%). All three occurrences of |
| 296 | amphibole are common (primary magmatic, secondary magmatic, and |
| 297 | pseudomorphic). Amphibole, biotite, and Fe-Ti-oxides are often found together in |
| 298 | 1-3 mm clots (Fig S4c). |
| 299 | The monzonites contain 40 to 60% plagioclase, 20 to 35% K-feldspar, 10 to |
| 300 | 20% biotite, 0 to 5% amphibole, and minor quartz (<5%). Euhedral to subhedral |
| 301 | primary magmatic amphibole grains (0.3-1.5 mm) lie between generally coarse |
| 302 | anhedral plagioclase and K-feldspars. Irregularly shaped flakes of biotite (0.5-2 |
| 303 | mm) are packed between plagioclase crystals and can be included in K-feldspars |
| 304 | (Fig S4d). Quartz (up to 0.5 mm) occurs interstitially. |

306 **RESULTS**

Zircon U-Pb isotopic measurements, bulk-rock major and trace element
concentrations, and mineral chemistry data are included along with sample GPS
locations in the Supplementary Data File (Tables S1-S3). Analytical methods are
described in the Supplementary Material.

311

312 Geochronology

313 Zircon from six total samples of monzodiorite and monzonite from the HLMC were dated using laser ablation inductively coupled mass spectrometry 314 (LA-ICPMS) at the University of California, Santa Barbara, and one granite 315 sample was dated using LA-ICPMS at the Arizona Laserchron Center. All dated 316 HLMC samples (not including the granite sample) are late Cretaceous in age. 317 Weighted mean crystallization ages and 2σ errors for monzondiorite samples are 318 95.1 ± 1.0 Ma for SNB-16-6, 95.6 ± 0.8 Ma for SNB-16-9, and 96.5 ± 0.8 Ma for 319 320 SNB-14-34. HL-17-2, an anorthosite, has an age of 95.7 \pm 0.7 Ma. The monzonite 321 ages are 96.3 \pm 0.8 Ma for SNB-16-12 and 90.5 \pm 0.8 Ma for SNB-14-40. These ages allow for up to 6 Ma of magmatic activity in HLMC and constrain 322 323 crystallization ages of the more mafic parts of the HLMC at 5 and 7 million years older than the Lake Edison granodiorite (90.9 \pm 0.1 Ma; Lackey et al. 2008) and 324 325 Round Valley Peak granodiorite (88.8 \pm 0.02 Ma; Lackey et al. 2008),

| 326 | respectively. HLMC is significantly younger than the granite of Chickenfoot Lake |
|-----|--|
| 327 | (at 191.6 \pm 1.0 Ma, this study). These ages are consistent with field relationships, |
| 328 | in which HLMC lithologies are observed intruding into Jcf and intruded by Kle |
| 329 | and Krv. Sample SNB-14-40, a monzonite from the northern edge of the mafic |
| 330 | complex near the contact with Krv, has a crystallization age ~5 Ma younger than |
| 331 | the other five mafic samples (90.5 \pm 0.8 Ma). This may represent a late stage melt |
| 332 | associated with differentiation of HLMC magmas that was remobilized by |
| 333 | intrusion of Kle, as their ages overlap within uncertainty. We presume that the |
| 334 | gabbros and norites crystallized within the range of monzodiorte ages (95.1-96.5 |
| 335 | Ma), though neither of these more mafic lithologies were directly dated due to the |
| 336 | rarity of zircon. |

337

338 Bulk-rock Major Element Chemistry

Major element bulk compositions from HLMC were analyzed via X-ray fluorescence (XRF) spectrometry at Caltech, following methods in Bucholz and Spencer (2019). Samples display a trend of magmatic evolution that straddles the boundary between a monzonitic series and granodioritic series (Bas et al. 1986; Middlemost 1994). For simplicity, we retain the lithology names norite, gabbro, monzodiorite and monzonite as classified based on mineralogy in previous sections, even if all samples of one lithology do not fit precisely within a single

| 346 | chemically defined field (Fig 3a). SiO_2 contents from the full suite of HLMC |
|-----|---|
| 347 | samples range from 43.6 to 62.0 weight percent (wt.%) and molar Mg numbers |
| 348 | $(Mg# = 100 \times Mg/[Mg+Fe])$ calculated using total Fe) range from 73 to 46 (Fig 3d). |
| 349 | Aluminum saturation index (ASI) values of HLMC samples range from 0.57 to |
| 350 | 0.96 (ASI= Al/ [Ca-1.67×P+ Na+ K]), indicating that all sampled lithologies are |
| 351 | metaluminous. |
| 352 | The norites represent the most primitive cumulates to crystallize from an |
| 353 | HLMC melt, based on their high Mg#s (73.1-64.6). The norites contain 49.8 to |
| 354 | 53.6% SiO ₂ , 10.2 to 20.0 wt.% MgO, and 9.8 to 14.8 wt.% FeO. Relatively |
| 355 | constant SiO ₂ content with decreasing MgO and FeO reflects increasing abundance |
| 356 | of plagioclase relative to orthopyroxene with progressive fractionation. Likewise, |
| 357 | co-variation in CaO (3.0-7.7 wt.%, Fig 3c) and Al_2O_3 (4.9-12.9 wt.%) is controlled |
| 358 | by the proportion of plagioclase relative to clinopyroxene, orthopyroxene, and |
| 359 | olivine. To highlight the differences in mineral abundances, we separate the norites |
| 360 | into two groups: high-Ca norites (CaO \geq 7.5 wt.%) and low-Ca norites (CaO < 6 |
| 361 | wt.%), with a compositional gap from 6 to 7.5 wt.% CaO (Figure 3b). In general, |
| 362 | the more differentiated norite samples (high-Ca group) have slightly lower SiO_2 |
| 363 | contents, significantly lower MgO and FeO contents, and elevated TiO ₂ , CaO, |
| 364 | Al_2O_3 , K_2O , Na_2O as compared to the low-Ca group. ASI varies from 0.57 to 0.72 |
| 365 | in the norites but does not clearly correlate with SiO ₂ . |

| 366 | The sequence of gabbros through monzonites follows a trend of increasing |
|-----|---|
| 367 | SiO_2 , K_2O , and Na_2O contents with decreasing MgO, FeO, CaO, Al_2O_3 , and TiO_2 |
| 368 | contents (Fig 3 and S10). ASI gradually increases from 0.70 to 0.96 from the |
| 369 | gabbros to the monzonites, in accordance with a decrease in the modal abundance |
| 370 | of calcic amphiboles relative to Al-bearing phases. Gabbros are the most |
| 371 | voluminous rock type in HLMC and fall into two groups based on major-element |
| 372 | concentrations: 1) low-SiO ₂ gabbros, with 43.9 to 46.8 wt.% SiO ₂ and 2) high-SiO ₂ |
| 373 | gabbros, with 49.2 to 52.5 wt.% SiO ₂ . The low-Si gabbros have higher proportions |
| 374 | of Fe-Ti oxides and pseudomorphic amphiboles relative to clinopyroxenes in the |
| 375 | high-SiO ₂ gabbros, while high-SiO ₂ gabbros contain a greater modal abundance of |
| 376 | plagioclase. Al ₂ O ₃ contents are significantly higher in the gabbros $(17.3-23.4)$ |
| 377 | wt.%) than the norites due to greater abundance of plagioclase and lower |
| 378 | abundance of orthopyroxene. The highest Al_2O_3 concentration (23.4 wt.%) is at |
| 379 | 52.2 wt.% SiO ₂ , then gradually decreases with increasing silica content through the |
| 380 | monzodiorites and monzonites. |

Two samples from cross-cutting diabase dikes and pods are shown to have very different Mg#s (64.1 and 42.6) despite similar SiO₂ content (47.0 and 49.2 wt.%). The lower Mg# sample has considerably higher alkali (Fig 3a) and P₂O₅ (0.84 wt.%) concentrations than other HLMC samples with similar silica contents (~0.25-0.4 wt.%), so this sample is unlikely to be co-genetic with HLMC. The

higher Mg# diabase is similar in composition to the least primitive norites or
highest Mg# gabbros, and therefore may be related to the HLMC system.

388

389 Bulk-rock Trace Element Chemistry

Trace element concentrations were collected with solution ICPMS, using 390 acid-digested chips of the glass disks used in XRF analyses. All norite samples 391 have low Sr concentrations (89-407 ppm) that increase with increasing modal 392 abundance of plagioclase (Fig 3e). Higher Sr contents (705-1030 ppm) in the 393 gabbros are reflective of higher plagioclase abundance. Sr contents in the 394 monzodiorites (659-921 ppm) and monzonites (604-934 ppm) overlap with and 395 extend to slightly lower values compared to the gabbros. Sr/Nd and Al/Si ratios, 396 and Eu anomalies are positively correlated and indicate the extent of plagioclase 397 accumulation. Primitive mantle normalized Eu anomalies (Eu/Eu*= $Eu_N / sqrt[Sm_N]$ 398 \times Gd_N]) are < 1 in the norites (0.45-0.78) and 0.72 to 1.46 in all other HLMC 399 lithologies. 400

The norites have relatively high Ni and Cr contents (97-504 ppm and 123551 ppm, respectively, Fig 3f and S11), which reflect the abundance of
orthopyroxene (up to 75%) in these samples, as well as the presence of minor
olivine and clinopyroxene. Ni and Cr contents are low in the low-SiO₂ gabbro

| 405 | group (3-24 ppm Ni, 2-21 ppm Cr), then increase slightly, on average, in the high- |
|-----|---|
| 406 | SiO_2 gabbros and remain relatively constant in the monzodiorites and monzonites. |
| 407 | On primitive-mantle normalized spider diagrams (Fig 4), all samples display |
| 408 | strong Nb and Ta depletions and Pb enrichment consistent with a subduction- |
| 409 | related origin for the HLMC parental melts (Miller et al. 1994; Stolz et al. 1996). |
| 410 | Most of the gabbros, particularly the low-SiO ₂ gabbros, exhibit Ti enrichments as |
| 411 | controlled by crystallization of Fe-Ti-oxides, biotite, and titanite. Significant |
| 412 | titanium depletions are observed in the norites, as these samples lack titanite. |
| 413 | Besides the norites, all lithologies display Sr enrichment, reflecting accumulation |
| 414 | of plagioclase in higher abundances than olivine and pyroxenes. Some norites are |
| 415 | considerably enriched in LIL elements, particularly Cs and Rb (up to 34 and 50 |
| 416 | ppm, respectively), likely due to biotite crystallization. In the more evolved rocks |
| 417 | and in two norites (Fig 4a, b, f-i), relative enrichments in Zr and Hf are observed |
| 418 | due to the presence of zircon. |
| 419 | Rare earth elements (REEs) and other incompatible element concentrations |

(Ba, Th, Nb, Pb, Hf, Ta, U, Rb, Zr) are positively correlated with SiO₂. HREE
elements are depleted relative to LREE elements in all rock types, and La/Yb ratios
increase with increasing silica content, though there is significant overlap between
rock types.

425 Classification of Cumulate and Liquid-Like Samples

Based on textural and compositional parameters, we classify each sample as 426 having a liquid-like or cumulate bulk composition. All norites are classified as 427 428 cumulates because cumulate textures were identified in all norite thin sections (Fig 2a), and their bulk-rock Mg#s are significantly higher than the rest of the HLMC 429 (Mg #= 64.6-73.1, with a gap in HLMC from 57.8-64.6). Gabbro textures are 430 variable from meso- or orthocumulates, to randomly oriented masses of 431 interlocking crystals (Fig 2d). The latter is suggestive of rapidly crystallized melt-432 like lithologies. In addition, we use Al/Si and Sr/Nd ratios as tracers of plagioclase 433 accumulation (Jagoutz 2010; Bucholz et al. 2014) and define cumulate gabbros as 434 those with AI/Si > 0.21 and cumulate monzodiorites and monzonites as those with 435 Sr/Nd > 30 (Fig 5a). These values place samples with textural observations into the 436 correct categories and allows us to classify samples for which we do not have 437 petrographic constraints. 438

When compared to the macroscopic rock textures, all liquid-like gabbros are medium-fine grained gabbros, and all coarse-grained and spotted gabbros are cumulates. Some medium-fine grained gabbros are also classified as cumulates based on our compositional criteria. Eu anomalies are variable within rock types, and either similar between liquids and cumulates of the same rock type (e.g., liquid-like monzodiorites and cumulate monzodiorites), or slightly elevated in the

445 cumulate group. Eu anomalies are <1 in the norites (0.45-0.69), reflecting minimal 446 plagioclase accumulation and incompatibility of Eu^{2+} in olivine and orthopyroxene 447 (Fig 5b).

448

449 Mineral Chemistry

Major element compositions of minerals were completed using electron probe microanalysis (EPMA) on roughly 3 to 10 grains per phase in each sample. In zoned crystals, at least 3 points were analyzed in each grain to assess core-torim compositional variation. Selected mineral compositions are summarized in Fig 6, and additional variation diagrams and geochemical data are included in the Supplementary Files (Fig S5-S8, Table S3).

456

457 Olivine

Olivine is rare in HLMC and present only in two low-Ca norite samples, SNB-14-62 and SNB-14-52. The highest Mg# olivine analyses are found as chadacrysts within in large amphibole oikocrysts. Olivine chadycrysts have Mg#s ranging from 80 to 81, while non-chadacrystic olivine grains have Mg#s ranging from 79 to 80, while olivines from cumulate xenoliths crystallized in the SNB lower crust extend to Mg# 92 (Dodge et al. 1986; Lee et al. 2001). Olivines have 0.28 to 0.35 wt.% MnO, which is negatively correlated with Mg# (Fig 6a).

Orthopyroxene

| 467 | Orthopyroxene is abundant in the norites, occasionally present in the |
|-----|--|
| 468 | gabbros, and rare in the monzodiorites and monzonites where it is found only in |
| 469 | amphibole cores. The Mg# of orthopyroxenes decreases from the norites (65-80) to |
| 470 | the gabbros (51-65). In the monzodiorites and monzonites, compositions |
| 471 | significantly overlap with those in the gabbros (Mg $\#$ = 49-59), suggesting that |
| 472 | these lithologies experience an early interval of orthopyroxene crystallization |
| 473 | followed by amphibole crystallization. |
| 474 | Orthopyroxene Cr_2O_3 contents generally decrease with decreasing Mg# and |
| 475 | vary from 0.01 to 0.11 wt.% in the norites and are below detection limit to 0.03 |
| 476 | wt.% in the gabbros, monzodiorites, and monzonites. Similarly, Al_2O_3 is positively |
| 477 | correlated with Mg#, ranging from 0.35 to 2.1 wt.% in the norites and 0.29 to 1.1 |
| 478 | wt.% in all other orthopyroxene-bearing lithologies (Fig 6b). MnO increases with |
| 479 | decreasing Mg# and varies between 0.30 and 0.55 wt.% in the norites, and 0.59 |
| 480 | and 1.1 wt.% in the gabbros and monzodiorites. The monzonites have slightly |
| 481 | higher orthopyroxene MnO concentrations of 0.95 to 1.2 wt.%. |
| 482 | There is a significant range in orthopyroxene Al_2O_3 contents in the SNB |
| 483 | xenoliths (0.7-5.4 wt.%; Dodge et al. 1986; Lee et al. 2001), likely due to |
| 484 | crystallization at a range of temperatures and pressures. Mg#s of these grains |

485 overlap with compositions in the norites and gabbros, as well as extend to higher
486 Mg#s (maximum 92).

487

488 *Clinopyroxene*

Clinopyroxene is a minor phase in the norites (<5% modal abundance), 489 common in the gabbros, and occasionally present in the cores of amphiboles in the 490 monzodiorites and monzonites. In the norites, clinopyroxene Mg#s range from 73 491 to 79 and decrease from individual grain cores to rims. Analogous to the 492 orthopyroxenes, clinopyroxene compositions in the gabbros, monzondiorites, and 493 monzonites are very similar to one another. In particular, the clinopyroxenes from 494 the cumulate gabbros and the liquid-like monzondiorites overlap significantly in 495 Mg#, which may indicate that these clinopyroxenes in HLMC may have 496 crystallized from a melt similar to the gabbros. Clinopyroxene Mg#s vary from 61 497 to 73 in the gabbros, 61 to 75 in the monzodiorites, and 60 to 68 in the monzonites. 498 499 Clinopyroxene Cr_2O_3 contents are roughly correlated with Mg#, ranging from 0.03 to 0.11 wt.% in the low-Ca norites, 0.02 to 0.03 wt.% in the high-Ca 500 norites, and commonly below detection limit (~0.015 wt. %) in all other rock 501 502 types, with a maximum of 0.04 wt.%. Clinopyroxene MnO content increases 503 consistently with decreasing Mg# (from 0.18 to 0.56 wt. %). Al₂O₃ content is 504 variable between 0.47 and 2.8 wt.%, commonly decreasing (~75% of grains) and

505 occasionally increasing ($\sim 25\%$ of grains) from crystal cores to rims. Al₂O₃ in 506 clinopyroxene cores increases from the norites (0.56-1.28 wt.%) and peaks at 1.79 507 wt.% in the liquid-like gabbros, then decreases through the gabbros, 508 monzodiorites, and monzonites to 1.09 wt.%. This peak corresponds to the maximum in bulk-rock Al₂O₃ concentration. 509 510 Amphibole 511 Amphibole is present in all rock types in HLMC and may be a primary 512 magmatic phase, secondary magmatic overgrowth on pyroxene crystals, or 513 pseudomorphic replacement of pyroxene due to reaction with an evolving melt. 514 There is no distinct compositional difference between these occurrences, however 515 pseudomorphic amphiboles contain abundant plagioclase micro-inclusions. 516 Amphibole compositions are dominated by magnesiohornblende based on the 517 classification of Leake et al. (1997). Analyses with <0.75 formula units of Al 518 519 (calculated on a 23-oxygen basis) are considered metamorphic alteration products and not further discussed. Nearly all amphibole in the high-SiO₂ gabbros (cumulate 520 and liquid-like) is actinolite, while all other rock types contain magmatic 521

522 amphibole.

523 There are two compositional populations of amphiboles in the low-Ca
524 norites, which are distinguishable by their aluminum contents and both fall under

525 the classification of magnesiohornblende (Fig 6c). Mg#s of both groups are comparable, with the high-Al population ranging from 71 to 77 and the lower-Al 526 527 population ranging from 72 to 79. The high-Al population (9.6-12.8 wt.% Al_2O_3) is 528 characterized by primary magmatic oikocrysts that surround olivine grains and is only present in samples SNB-14-52 and SNB-14-62 (c.f., Fig 2b). This population 529 is also high in TiO₂ (2.9-3.6 wt.%) and Na₂O (1.7-2.3 wt.%) compared to other 530 amphiboles in the norites with similar Mg#s (0.2-1.3 wt.% TiO₂ and 0.6-1.1 wt.% 531 Na₂O). The lower-Al population (4.5-5.9 wt.% Al_2O_3) appears to also be of 532 primary magmatic origin based on the euhedral to subhedral crystal shapes. These 533 two populations are not observed in the same samples and so may indicate slight 534 compositional variation between different norite pods. All amphiboles in the high-535 Ca norites are similar in composition with Mg#s ranging from 67 to 71 and Al_2O_3 536 varying between 4.8 and 6.3 wt.%. Amphibole Mg#s in the gabbros and 537 monzodiorites are negatively correlated with Al_2O_3 in a series of parallel trends, 538 539 which each represent within-sample compositional variation reflecting sub-solidus Tschermak exchange, a trend that is not evident in amphiboles from the norites. 540 541 Similarly, TiO₂ and Na₂O are negatively correlated with Mg# within individual 542 samples. Amphibole MnO content increases with decreasing Mg# and is relatively 543 constant within each rock type, ranging from 0.09 to 0.31 wt.% in the norites, 0.21 544 to 0.34 wt.% in the gabbros, 0.28 to 0.59 wt.% in the monzodiorites, and 0.48 to

545 0.62 wt.% in the monzonites. CaO content is negatively correlated with Mg#,

though there is considerable overlap between rock types.

547

548 Biotite

Biotite is present in all rock types, typically interstitially, and is thus 549 interpreted as a late crystallizing phase. The Mg# of biotite in the low-Ca norites 550 ranges from 54 to 76, with a compositional gap between 59 and 63. The high-Mg# 551 biotite group extends to higher TiO₂ (2.4-5.2 wt.%) and F (0.42-0.81 wt.%) than 552 the low-Mg# group (1.7-4.1 and 0.32-0.43 wt.%, respectively). There is no clear 553 textural difference between the low- and high-Mg# biotite, though individual 554 samples only contain biotite from one compositional group. Biotite Mg# decreases 555 to the high-Ca norites (52-60) and gabbros (47-61). The range of biotite Mg#s in 556 the monzondiorites (45-58) overlaps with that of the gabbros and the monzonites 557 (47-52). 558

559 MnO contents systematically increase with decreasing Mg#, from 0.02 to 560 0.18 wt.% in the norites to a maximum of 0.43 wt.% in the monzonites. Al_2O_3 561 content of biotite in the norites is variable (11.9-14 wt.%) and not clearly 562 correlated with Mg#. There is significant overlap in Al_2O_3 contents in biotite for 563 the remainder of the studied lithologies (13.2-15.8 wt.% in gabbros, 12.9-16.3 564 wt.% in monzondiorites, 12.5-15.6 wt.% monzonite). Within individual samples,

| 565 | there is a weak positive correlation between Mg# and Al ₂ O ₃ content of biotite, |
|-----|---|
| 566 | which is the opposite correlation from what is predicted by Tschermak exchange, |
| 567 | and suggests minimal subsolidus alteration. Na ₂ O contents reach 0.18 wt.% in |
| 568 | high-Mg# biotite in the norites, and is <0.11 wt.% in other lithologies, and often |
| 569 | below detection limit. Fluorine decreases with decreasing Mg#, following |
| 570 | established Fe-F avoidance trends (Fig 6d) (Mason 1992; Icenhower and London |
| 571 | 1997). Biotite F content ranges from 0.32 to 0.91 wt.% in the norites, and in other |
| 572 | rock types biotite contains between 0.19 and 0.54 wt.% F. |
| 573 | |

574 *Feldspars*

Plagioclase feldspar is dominantly interstitial in the norites, though 575 occasional euhedral crystals are present, and is a common fractionating phase in all 576 other HLMC rock types. The anorthite content (An) of interstitial plagioclase in 577 norites is highly variable, ranging from An₂₉ to An₇₂, while euhedral plagioclase 578 has approximately An₈₂. Euhedral to subhedral plagioclase in the gabbros can have 579 distinct core and rim compositions with An₇₆ to An₉₂ cores and An₄₀ to An₆₂ rims. 580 In crystals without a core-rim compositional difference, the composition is 581 582 comparable to the aforementioned rims. Distinct cores and rims are present in <5%of plagioclase crystals in the monzodiorite plagioclase and absent from the 583

| 584 | monzonites. Plagioclase is An_{27} to An_{58} in rims and $\sim An_{77}$ in cores in the |
|-----|--|
| 585 | monzodiorites, and An_{31} to An_{54} in the monzonites. |
| 586 | Alkali feldspar is found as small, interstitial grains in the norites and |
| 587 | gabbros, but becomes increasingly common in the monzodiorites and monzonites. |
| 588 | Orthoclase (Or) and albite (Ab) content is similar for all rock types, with variation |
| 589 | between Or ₇₂ Ab ₁₈ and Or ₉₈ Ab ₂ . |
| 590 | |
| 591 | Zircon |
| 592 | Zircon grains from HLMC have Ti concentrations ranging from 2 to 24 ppm |
| 593 | in the monzonites, 4 to 31 ppm in the anorthosites, and 5 to 27 ppm in the |
| 594 | monzodiorites. Zircon is rare in the gabbros and norites and was therefore not |
| 595 | separated for analysis. Hf concentration is negatively correlated with Ti |
| 596 | concentration within each sample, ranging from 8610 to 17100 ppm, with 2470 to |
| 597 | 7670 ppm of intra-sample variation. |
| 598 | REE concentrations of melts in equilibrium with the zircons were calculated |
| 599 | using the methods of Chapman et al. (2016) (Fig S9). Primitive mantle normalized |
| 600 | REE (plus Nb and Y) patterns of calculated melts are nearly parallel to the |
| 601 | corresponding whole-rock patterns. In cumulate samples, calculated melt |
| 602 | compositions are similar to the whole rock concentrations in the LREE, and |
| | |

slightly higher or lower in the HREE. Calculated melts have Eu anomalies <1

- 604 (0.70-0.81) that are slightly lower or equal within error (2 sigma of all analyses
- within a sample) than in the corresponding whole rock composition, implying that

606 zircon saturation occurred after some plagioclase fractionation.

607

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| 0 |

609 Crystallization Conditions

610 Out best estimates for crystallization pressures, temperatures, H_2O content, 611 and fO_2 and of HLMC lithologies are summarized in Table 2 and described below.

612 *Pressure*

We perform Al-in-hornblende barometry calculations using amphibole 613 analyses from monzonites, as this is the only HLMC lithology that contains the 614 requisite phase assemblage for this calibration (Mutch et al. 2016). Additionally, 615 these samples exhibit a lesser degree of Tschermak exchange than the gabbros and 616 monzodiorites, a process which could result in inaccurately low calculated 617 pressures. Using high-Al (7.5-8.8 wt.% Al₂O₃), low-Mg# (46-56) amphibole 618 analyses (i.e., those inferred to have experienced the least Tschermak exchange) 619 we calculate pressures of 0.30 to 0.34 GPa. Assuming an average density of 2.8 620 g/cm³ for the overlying largely granitic crust (Christensen and Mooney 1995), this 621 equates to an approximate depth between 11 and 13 km. 622

623

624 Temperature

We use a variety of mineral thermometers to constrain crystallization 625 626 temperatures across the range of HLMC lithologies. For these calculations, we utilize analyses from the rims of adjacent grains to give the best estimate of 627 equilibrium compositions. In the gabbros, orthopyroxene and clinopyroxene pairs 628 are out of Fe-Mg equilibrium, with partition coefficients ([Fe/Mg]_{cpx}/[Fe/Mg]_{opx}) of 629 0.65-0.86 as opposed to the equilibrium value of 1.09 ± 0.14 (Putirka 2008). This 630 is not surprising due to the presence of pervasive exsolution lamellae for samples 631 with coexisting ortho- and clinopyroxene. Thus, two-pyroxene equilibrium 632 temperatures of 760 to 950 °C (Köhler and Brey 1990; Putirka 2008) likely 633 represent cooling, rather than primary crystallization temperatures. The norites, 634 however, lack discernable pyroxene exsolution and contain little to no 635 clinopyroxene that could react with the available orthopyroxene. Orthopyroxene in 636 the norites has CaO contents ranging from 0.6 to 1.6 wt.%, yielding crystallization 637 temperatures of 900 to 1170 °C based on the Ca-in-orthopyroxene thermometer of 638 Köhler and Brey (1990). 639

Evidence of subsolidus chemical exchange is found in amphibole analyses in the gabbros, monzondioites, and monzonites, as intra-sample negative correlation between Mg# and Al_2O_3 in amphibole suggests alteration of the original magmatic compositions by Tschermak exchange (Fig 6c). However, in the norites, amphibole

is present in two distinct compositional populations that do not define a continuous 644 645 trend of decreasing Al_2O_3 with increasing Mg#, which we interpret as preserving 646 magmatic amphibole compositions. In the gabbro, monzonite, and monzodiorite 647 samples, we calculate crystallization temperatures based on amphibole compositions with realtively high Al_2O_3 contents (>7.5 wt.%) to avoid sites 648 649 significantly affected by subsolidus alteration. We do not report crystallization temperature estimates from the pseudomorphic amphibole grains, as these are 650 likely out of equilibrium due to incorporation of excess Ca from the reaction of 651 melt + clinopyroxene \rightarrow melt + amphibole into the plagioclase needed for 652 653 temperature calculation.

Thermometry based on the SiO₂ and anorthite contents of coexisting 654 655 amphibole and plagioclase, respectively, (Blundy and Holland 1990) defines a wide range of crystallization temperatures within HLMC depending on the rock 656 type. As analyses of amphiboles in the norites do not follow a trend of decreasing 657 658 Al₂O₃ content with increasing Mg# as expected if significant Tschermak exchange has significantly affected the mineral compositions, we believe the estimated 659 660 temperatures from the norite samples are representative of crystallization 661 conditions. Calculated crystallization temperatures of amphibole oikocrysts and 662 associated plagioclase in the norites are 865 to 950 $^{\circ}$ C, using amphibole SiO₂ 663 contents ranging from 40.5 to 42.9 wt. % and plagioclase compositions of An₅₈ to

| 664 | An ₆₁ . Estimates from non-oikocrystic amphibole in the same samples are |
|-----|---|
| 665 | significantly lower, ranging from 700 to 765 $^{\circ}$ C, using amphibole SiO ₂ contents |
| 666 | ranging from 46.5 to 50.5 wt. % and plagioclase compositions between An_{23} and |
| 667 | An ₅₂ . Locally elevated H_2O contents in some norite pods may have allowed for |
| 668 | earlier amphibole saturation in some norites, as the oikocrystic, high-Al |
| 669 | amphiboles are not found in all norite samples. The high-Al oikocrysts likely |
| 670 | crystallized earlier in the fractionation sequence than the low-Al amphiboles, as |
| 671 | they record higher crystallization temperatures and contain inclusions of olivine, |
| 672 | which are not found in the low-Al amphiboles. The lack of olivine inclusions in the |
| 673 | low-Al amphiboles may indicate that olivine was lost from the fractionating |
| 674 | assemblage in favor of orthopyroxene prior to crystallization of the low-Al |
| 675 | amphiboles. |
| 676 | In the gabbros, primary and secondary (i.e., rims on pyroxenes) magmatic |
| 677 | amphibole temperatures are indistinguishable from one another and range from |
| 678 | 740 to 880 °C (SiO ₂ in amphibole 44.3-50.6 wt.%, anorthite content An ₃₇₋₈₁). Using |
| 679 | the same method, monzodiorites and monzonites yield amphibole-plagioclse |
| 680 | crystallization temperatures of 815 to 870 $^{\circ}$ C (SiO ₂ in amphibole 42.6-48.6 wt.%, |
| 681 | anorthite content An ₃₅₋₅₂), and 800 to 850 °C (SiO ₂ in amphibole 41.8-45.2 wt.%, |

anorthite content An_{34-51}), respectively.
684 Water Content and Oxygen Fugacity

| 685 | Relatively early crystallization of amphibole, and to a lesser extent biotite, in |
|-----|---|
| 686 | the HLMC fractionation sequence requires a hydrous parental melt. We apply the |
| 687 | plagioclase-liquid hygrometer of Waters and Lange (2015) to plagioclase cores in |
| 688 | the liquid-like gabbros to estimate melt H ₂ O content. Plagioclase core |
| 689 | compositions are used because they represent the earliest crystalized sections, |
| 690 | which are most likely to be in equilibrium with the bulk-rock "melt" composition. |
| 691 | Using 990 to 1100°C as the temperature range for the onset of gabbro |
| 692 | crystallization in HLMC based on experimental studies at upper crustal pressures |
| 693 | and bulk compositions similar to the HLMC gabbros (Grove et al. 2003; Hamada |
| 694 | and Fujii 2008), we estimate H_2O contents of 2.9 to 3.5 wt.% in the liquid like |
| 695 | gabbros. |
| 696 | Oxygen fugacities (fO2) play a significant role in the Fe-Mg partitioning into |
| 697 | ferromagnesian minerals during crystallization. Oxygen fugacities elevated above |
| 698 | that for mid-ocean ridge basalts (near the fayalite-magnetite-quartz buffer) are |
| 699 | expected, as arc magmas are commonly characterized by fO_2 at NNO or higher |
| 700 | (Sisson et al. 2005; Kelley and Cottrell 2009; Rowe et al. 2009; Brounce et al. |
| 701 | 2014). Applying the oxybarometer of Loucks et al. (2020), which utilizes the |

concentrations of U, Ti, and Ce in zircon grains, we find a range of fO_2 values from

 $\Delta NNO - 1.6$ to 1.3 in the dated grains from monzodiorite and monzonite samples,

| 704 | with an average fO_2 of ΔNNO -0.5. We use biotite and bulk-rock compositions and |
|-----|--|
| 705 | MELTS thermodynamic modeling to calculate fO_2 as presented in Bucholz et al. |
| 706 | (2018) for melt-like monzonite SNB-16-9, which contains a relatively narrow |
| 707 | range of biotite compositions (Fig 6d and S6) and thus subsolidus reequilibration |
| 708 | was likely minimal. We find that the HLMC melts are oxidized to between NNO |
| 709 | and $\Delta NNO + 0.6 (\pm 1 \log unit)$. |
| 710 | |
| 711 | Comparison to Previous Studies |
| 712 | Our results from Al-in-hornblende barometry indicate a crystallization |
| 713 | pressure of between 0.30 and 0.34 GPa, which is consistent with previous studies |
| 714 | and show that the SNB around HLMC crystallized in the upper crust. The central |
| 715 | SNB exposes dominantly upper crustal intrusions that crystallized between 0.1 to |
| 716 | 0.4 GPa. Previous studies place the location of HLMC at pressures of 0.2 to 0.3 |
| 717 | GPa (Ague and Brimhall 1988; Chapman et al. 2012), and 0.3 to 0.35 GPa (Nadin |
| 718 | et al. 2008). |
| 719 | The following experimental studies were selected for comparison as they |
| 720 | produce the series of mineral assemlages observed in HLMC. Our upper |
| 721 | temperature estimate for the norites based on the Ca-in-orthopyroxene |
| 722 | thermometer (1170 °C) is similar to experiments by Hamada and Fuji (2008), |
| | |

which produce an olivine + orthopyroxene + plagioclase assemblage from a high-

| 724 | MgO basalt starting at 1160 $^\circ C, 0.4$ GPa, 1.6-2.7 wt.% H ₂ O, and one log unit |
|-----|--|
| 725 | above the Ni-NiO buffer (Δ NNO+1). Clinopyroxene appears in the experimental |
| 726 | assemblage at 1130 $^{\circ}$ C, and the assemblage of orthopyroxene + clinopyroxene + |
| 727 | plagioclase + Fe-oxides continues to 1100°C and the termination of the |
| 728 | experiment. The assemblage orthopyroxene + clinopyroxene + plagioclase + |
| 729 | amphibole first crystallizes at 990 °C, 0.2 GPa, NNO, and 4.5 wt.% H_2O in |
| 730 | experiments by Grove et al. (2003). Based on these two studies, which replicate the |
| 731 | mineral assemblages found in the norite and gabbro cumulates in HLMC and |
| 732 | bracket the pressure and H_2O content estimates for the HLMC and were performed |
| 733 | at similar fO_2 conditions, the onset of gabbro fractionation, characterized by |
| 734 | common clinopyroxene and minimal orthopyroxene, likely occurred between 1100 |
| 735 | and 990 °C. |
| 736 | Experiments by Blatter et al. (2013) using an arc basalt starting material |
| 737 | similar to HLMC melt-like gabbro compositions first crystallize amphibole + |
| 738 | orthopyroxene + clinopyroxene + plagioclase from a primitive high-Al arc basalt at |
| 739 | 0.4 GPa, 1000 °C, and 4 wt% H ₂ O. However, in these experiments orthopyroxene |
| 740 | and clinopyroxene are present in roughly equal proportions and olivine is not |
| 741 | present, unlike the HLMC norites. Orthopyroxene is part of the fractionating |
| 742 | assemblage of mid-ocean ridge dacites during 0.2 GPa crystallization experiments |
| 743 | under oxidizing conditions (NNO +1 to +2.1) with H_2O contents between 0.8 and 2 |

| 744 | wt.% (Erdmann and Koepke 2016). At higher water contents (>2 wt.%) in these |
|-----|--|
| 745 | experiments, the assemblage lacks orthopyroxene. The dominance of |
| 746 | orthopyroxene over clinopyroxene in the first crystallized HLMC cumulates |
| 747 | indicates a reduced H_2O content compared to Blatter et al. (2013), but the early |
| 748 | crystallization of amphibole suggests H ₂ O contents higher than in experiments |
| 749 | conducted by Erdmann and Koepke (2016) ($H_2O > 2$ wt.%). Experimental |
| 750 | evidence thus constrains the H_2O content of the HLMC parental melt between 2 |
| 751 | and 4 wt.%. |
| 752 | In the HLMC norites, amphibole oikocrysts containing olivine imply that |
| 753 | these phases coexist in the magma. Crystallization experiments of Krawczynski et |
| 754 | al. (2012) limit the stability of the crystallizing assemblage of olivine + amphibole |
| 755 | to between an fO_2 of NNO and Δ NNO +3 at 0.3 GPa at the calculated |
| 756 | crystallization temperature of the oikocrystic amphiboles (~950 °C). This study |
| 757 | complements our fO_2 calcultions in the monzodiorites and monzonites ($fO_2 \sim NNO$ - |
| 758 | Δ NNO+0.6), and shows that the parental melt to HLMC was similarly oxidized in |
| 759 | comparison to the more evolved lithologies. |
| 760 | |

761 A Cogenetic Fractionation Sequence

762 Field and Petrographic Evidence

The western limb of HLMC is composed of mainly medium-fine grained 763 764 gabbro with elongate, N-S trending texturally distinct gabbro bodies (coarse-765 grained and spotted gabbros, Fig 1). All coarse-grained and spotted gabbros, as 766 well as some of the medium-fine grained gabbros, exhibit ortho- to mesocumulate textures. The contacts between individual gabbro bodies and the medium-fine 767 grained gabbro are gradational over 1 to 5 cm, exhibiting mingling textures and 768 inclusions of gabbros with contrasting textures (Fig S2, S3). These characteristics 769 suggest that the gabbros of the western limb accumulated from multiple pulses of 770 magma that supplied compositionally similar melts within a relatively short period 771 of time, such that previously intruded gabbroic magma was still partially molten to 772 allow for mingled and gradational contacts. Due to their close temporal 773 association, these pulses of magma likely originated from the same source, and are 774 therefore part of a cogenetic sequence. The presence of both liquid-like and 775 cumulate gabbros in the western limb suggest that the parental melts to the gabbros 776 continued to fractionate in-situ. 777

The southeastern section of the HLMC consists of a single large mediumfine grained gabbro body containing norite and anorthosite cumulate pods, as well as a lens of coarse-grained gabbro, but is overall more texturally homogenous than the western section. The presence of well-defined cumulate textures in these rocks indicates that separation of crystals and liquid occurred within the mafic complex.

783 The contacts between gabbros, monzodiorite, and monzonite in the northern and northeastern section of the HLMC (Fig 1) are largely gradational over 5 to 10 m. 784 785 and there is no distinct contact between the medium-fine grained gabbros of the 786 western limb and this lithology in the eastern section. This suggests that eastern HLMC and the western medium-fined grained gabbros were emplaced as a single 787 large intrusion that differentiated *in-situ* to create the norite and gabbro cumulates 788 and melt was extracted from the gabbros to produce the more evolved 789 monzodiorites and monzonites. This is consistent with petrographic observations 790 suggesting that most of the gabbros are meso- and adcumulates, with the exception 791 of three liquid-like gabbro samples. 792

The presence of both cumulate and liquid-like gabbros, monzodiorites, and 793 monzonites suggests that the gabbroic HLMC parental intrusions fractionated in-794 situ. Multiple intrusions into the medium-fine grained gabbro are preserved as 795 gabbros with contrasting textures, and the bodies that represent separate intrusions 796 797 all have parental melts of similar composition. The gradational boundaries between the gabbros and monzonitic lithologies, as well as evolved pods within the 798 gabbros, indicate that the evolved lithologies were generated by extraction of melt 799 800 from the gabbroic cumulates. While some melts may have been extracted to 801 shallower levels in the crust, the liquid-like compositions present in HLMC 802 represent melts generated by fractionation within HLMC.

803 Evidence for mixing between HLMC and felsic melts at the level of emplacement is minimal and confined to the outer margins of the complex, and so 804 805 is unlikely to have caused the compositional variation within the mafic body. 806 Locally, the 90.5 to 96.5 Ma HLMC parental melts intruded into the much older granite of Chickenfoot Lake (Jcf, 191.6 Ma), which would have been fully 807 crystallized prior to the mafic intrusions. Any volume of Jcf-derived felsic melt 808 809 would need to be produced by heat from HLMC intrusions. This effect is visible along the HLMC-Jcf contact, where a dioritic gradational zone (Fig S3b) indicates 810 short-distance mixing between a HLMC parental melt and granitic melt produced 811 by partial melting of Jcf. However, this hybridized lithology does not extend more 812 than 1 m from the granite-gabbro contact. The contact between Kle and HLMC is 813 sharp and indicates little to no mixing or assimilation. Because Krv is younger than 814 HLMC, and HLMC-sourced blocks in the brecciated contact between these units 815 are angular with sharp contacts, HLMC was likely solidified by the time of Krv 816 intrusion and therefore could not mix. Field evidence precludes substantial magma 817 mixing origin for the compositional diversity within the HLMC and supports in-818 situ fractional crystallization. 819

821 *Geochemical Evidence*

The HLMC compositional trends suggest that the various rock types are 822 related to one another by fractional crystallization. Bulk-rock analyses preserve a 823 824 wide range of SiO₂ contents from 43.6 to 61.5 wt.% (Fig 3). Bulk-rock Mg# decreases from 73.1 to 35.8, though a compositional gap exists between the norites 825 and gabbros (Mg#= 57.8 to 64.6). Al₂O₃, TiO₂, CaO, MgO, and FeO contents are 826 variable over approximately constant SiO₂ content in the norites, then smoothly 827 decrease from the gabbros to monzodiorites to monzonites. Similar trends are 828 present in K₂O and Na₂O, but their concentrations increase with increasing SiO₂ 829 content. These compositional relationships are consistent with a fractional 830 crystallization liquid line of descent (LLD) and associated cumulates in arc settings 831 (Müntener and Ulmer 2018). 832 In general, incompatible trace element concentrations in liquid-like samples 833 increase with increasing SiO_2 content, indicating enrichment of these elements in 834 the remaining melt with progressive fractionation. Exceptions to this are Ni and Cr, 835 which are are rapidly removed from the melt during norite crystallization, and Sr 836 which decreases with increasing SiO₂ after saturation of significant amounts of 837 plagioclase. HREE elements are depleted relative to LREE elements in all rock 838

types (Fig 4), and this depletion, indicated by La/Yb ratio, increases with

840 increasing SiO_2 content. Though there is significant overlap in the degree of HREE

841 depletion between rock types, this pattern nevertheless supports a fractional

842 crystallization origin for the HLMC lithologies.

843 Mineral compositions are also consistent with a cogenetic fractionation sequence. Very similar trends are seen in ortho- and clinopyroxene, in which MnO 844 and Mg# are negatively correlated and higher Mg#s correspond to lithologies 845 crystallized earlier in the fractionation sequence. Cr₂O₃ and Al₂O₃ in orthopyroxene 846 (Fig 6b) decrease with decreasing Mg# and with increasing bulk-rock SiO₂ content, 847 indicating depletion of these components in the melt with olivine + pyroxene and 848 plagioclase fractionation. Euhedral plagioclase crystals in the norites have very 849 similar An content to the plagioclase cores in the gabbros and monzodiorites, 850 suggesting that these cores crystalized from a melt similar to the norite parental 851 melt and were progressively rimmed by lower-Ca plagioclase as the remaining 852 melt evolved. In biotite, the observed decreasing F (Fig 6d) and TiO₂ content with 853 decreasing Mg# correspond to increasing FeO content in the biotite and decreasing 854 crystallization temperature, respectively. 855

Thus, both the mineral compositions and modal mineralogy of the rocks indicate derivation by fractional crystallization from a common parental melt defined by a lithological progression of norites \rightarrow gabbros \rightarrow monzodiorites \rightarrow monzonites and a mineral crystallization sequence of olivine \rightarrow orthopyroxene + amphibole(high-

860 Al) \rightarrow clinopyroxene + Fe-Ti oxides \rightarrow plagioclase + amphibole(low-Al) \rightarrow 861 biotite \rightarrow K-feldspar \rightarrow quartz.

862

863 **Composition of the HLMC Parental Melt**

We define the HLMC parental melt as a representative, initial melt 864 composition that intruded into the upper crust at the location of HLMC and 865 fractionated to produce the range of observed lithologies. Based on our inferred 866 fractionation sequence, olivine is the first phase to crystallize from the parental 867 868 melt and is our best indicator of its composition. The most primitive analyzed olivine core in HLMC has a Mg# of 81. We calculate the Mg# of the parental melt 869 870 coexisting with this olivine to be 51, using the olivine-melt distribution coefficient of 0.32 (Blundy et al. 2020; Beattie 1993) and assuming an fO_2 of NNO and thus 871 Fe³⁺/Fe^T ratio of 0.2 (Kress and Carmichael 1991). 872

Primitive lavas erupted at continental arcs are thought to be representative of sub-arc mantle melts, and span a Mg# range of 67 to 73 based on a global compilation by Schmidt and Jagoutz (2017). The Mg# of the HLMC parental melt (51) is significantly lower than these values and therefore does not represent a (non-modified) partial melt of the mantle, but rather one that has experienced prior differentiation at deeper crustal levels. The major and trace element concentrations (Cr, Ni, Rb, Sr, Y, Zr, Nb, Ba, REEs, Hf, Ta, Pb, Th, U) of the HLMC parental

melt are determined via fractionation modeling as discussed below, and included in the Supplementary Data File (Table ST5). Though a basaltic dike exists in HLMC that appears to be a potential parental melt composition based on major element concentrations (SNB-14-39), this sample is significantly enriched in incompatible trace elements (particularly Nb, Rb, La, Ce, Lu, Ta, and Th) relative to the liquidlike samples at similar SiO₂ contents. Thus, we infer that this dike is not a parental melt and prefer the calculated parental melt composition.

887

888 Evidence for Lower Crustal Crystallization-Differentiation

Although the SNB is dominated by felsic plutons that crystallized at shallow 889 pressures (generally <0.5 GPa; Nadin et al. 2016), direct evidence for deep crustal 890 processes is present as ultramafic to mafic Cretaceous xenoliths in Miocene to 891 Pliocene basalts in the central SNB, sourced from 1 to 3 GPa (Lockwood and 892 Bateman 1976; Peselnick et al. 1977; Domenick et al. 1983; Dodge et al. 1986; 893 Dodge et al. 1988; Mukhopadhyay 1991; Mukhopadhyay and Manton 1994; Ducea 894 and Saleeby 1996; Ducea and Saleeby 1998; Lee et al. 2001; Lee et al. 2006). In 895 addition, southern exposures of the batholith in the Tehachapi Mountains 896 crystallized at pressures between 0.7 to 1.0 GPa (Ross 1985; Sams and Saleeby 897 1988; Pickett and Saleeby 1993; Lackey et al. 2005; Saleeby et al. 2008; Klein and 898 899 Jagoutz 2021, Rezeau et al. 2021). Orthopyroxene compositions from lower crustal

900 garnet-pyroxenite cumulate xenoliths range from Mg# ~80 to 90 (Fig 6b; Dodge et al. 1986; Lee et al. 2006). Using the orthopyroxene-melt distribution coefficient of 901 902 0.284 (Beattie 1993) and assuming an fO_2 of NNO (as above), the orthopyroxene 903 crystallized from melts ranging from a primitive mantle melt (Mg#= 69) to a fractionated basalt (Mg#= 48). These cumulates therefore record lower crustal 904 905 differentiation and the generation of low-Mg basalts that are consistent with the 906 HLMC parental melt composition. Additionally, the most primitive HLMC norite orthopyroxene compositions overlap with the least primitive compositions in the 907 xenoliths (Mg# ~80), indicating that both cumulates crystallized from similarly 908 909 fractionated melts.

Likewise, a lower crustal section of the southern SNB is exposed in the 910 Tehachapi Mountains, and exposes depths of up to 30 km (~1.0 GPa) (Ague and 911 912 Brimhall 1988; Pickett and Saleeby 1993). Within the Tehachapi Complex, the 913 Bear Valley Intrusive suite (BVIS, 0.4-1.0 GPa) consists of tonalite, norites, and 914 hornblende gabbros (Klein and Jagoutz 2021). The mineralogical similarities between the gabbroic cumulate rock types of the BVIS and the HLMC mafic 915 916 cumulates, particularly the prevalence of amphibole and orthopyroxene, suggest 917 that the melts that intruded into the upper crust are compositionally similar to those 918 that crystallized in the SNB mid-lower crust. Additionally, the most primitive bulk 919 rock analysis of gabbroid cumulates from the BVIS (Mg#=73.4; Klein and Jagoutz

2021) is similar to the most primitive HLMC norite (Mg#=73.1), suggesting that
the parental melts to both intrusions were similarly evolved basalts. Thus the
parental melts to both the BVIS and the HLMC may be generated by the same
process, namely through deeper crustal differentiation as recorded by the
ultramafic xenoliths.

In our case study of HLMC, differentiation of a primitive basalt is required 925 prior to the intrusion of the low-Mg parental melt in the upper crust. We favor 926 lower crustal fractional crystallization over partial melting of underplated basalts 927 as thermal modeling studies demonstrate that this latter process is more likely to 928 produce more silicic rather than basaltic or basaltic-andesite compositions (Annen 929 et al. 2006; Jagoutz and Klein 2018). Remaining possibilities for generating an 930 evolved basaltic melt include assimilation of felsic crust by a primitive basalt or 931 direct fractionation from a primitive basalt in the lower, mid- or upper crust. 932 Assimilation of crustal material is highly unlikely to produce the HLMC parental 933 melt composition from a mantle melt because assimilation of enough crustal melt 934 to decrease the basalt Mg# from ~70 to 51 would significantly increase the melt 935 936 SiO₂ content. Mid- and upper crustal fractionation of large volumes of primitive 937 melts is improbable because ultramafic cumulates are rarely observed at these 938 depths in the SNB. We therefore infer that the initial stage of fractional

- 939 crystallization occurred in the lower crust within the range of cumulate-dominated940 lithologies, as observed in accreted arc sections.
- 941
- 942 Polybaric Fractionation Model

Field relationships and geochemistry indicate that the HLMC lithologies 943 may be related by a common liquid line of descent (LLD) produced via fractional 944 crystallization, and their trace element signatures indicate that parental melts were 945 likely sourced from a sub-arc mantle melt. However, as discussed above, this 946 parental melt was fractionated relative to a primitive mantle melt. Thus, 947 fractionation must have occurred prior to the intrusion of the HLMC parental melts 948 at their current level in the crust (~0.3 GPa) where further differentiation occurred. 949 In agreement with studies of polybaric fractionation in arcs (Almeev et al. 2013; 950 Hamada et al. 2014; Melekhova et al. 2015), we construct a two-stage fractionation 951 model to test our fractional crystallization hypothesis. Stage 1 models fractional 952 crystallization of a primitive mantle melt in the lower arc crust using cumulate 953 compositions of xenoliths crystallized at depths of >30 km (Lee et al. 2006) to 954 produce a low-MgO basalt that becomes the parental melt to HLMC. Stage 2 955 956 tracks fractionation of this evolved basalt in the upper crust by removal of analyzed cumulate compositions from the HLMC from an evolving modeled liquid 957 composition. It is conceivable that multiple episodes of fractionation could have 958

| 959 | occurred prior to Stage 2 at different levels in the crust below 0.3 GPa. However, |
|-----|---|
| 960 | we choose the simplest approach and assume only one stage of crystallization prior |
| 961 | to intrusion at the level of HLMC exposure. The LLDs generated from our |
| 962 | modeling are illustrated in Figure 7, as well as Figures S10 and S11, and all results |
| 963 | of modeling calculations are included in the Supplementary Data File (Tables S4- |
| 964 | S6). |

965

966 Stage 1: Lower Crustal Fractionation Model Description

We model the evolution of a sub-arc mantle-derived melt by progressive 967 removal of bulk cumulate compositions from a melt composition, following the 968 methods of Jagoutz (2010). In this initial deep-crustal phase of differentiation, we 969 use representative compositions of cumulate xenoliths (Table S4) that crystallized 970 in the SNB lower crust between 1 and 3 GPa (Dodge et al. 1986; Lee et al. 2006), 971 using a fractionation sequence of orthopyroxenite and websterite \rightarrow garnet-972 pyroxenite. The starting melt composition was varied within the range of primitive 973 basalts from the Andes, Cascades, Central American, Mexican, and New Zealand 974 continental arcs (Schmidt and Jagoutz 2017). Stage 1 fractionation produces a 975 fractionated melt in equilibrium with the most primitive cumulates in HLMC at the 976 end of the Stage 1 and most successfully reproduces the HLMC liquid-like 977 compositions when input into the Stage 2 model. Stage 1 lower crustal 978

979 fractionation proceeds until the remaining melt reaches an Mg# of 51 after 33% crystallization, in equilibrium with the most primitive minerals in HLMC. 980 981 Because comprehensive trace element concentrations are not available for 982 all xenolith lithologies, we also model Stage 1 fractional crytallization using 983 alphaMELTS (Ghiorso and Sack 1995; Asimow and Ghiorso 1998) to generate a 984 trace element profile for the HLMC parental melt. A crystallization pressure of 1.1 985 GPa, or ~35 km depth, was used to replicate the lower crust of the Sierra Nevada arc as predicted by seismic studies (Jones and Phinney 1998; Ruppert et al. 1998; 986 Fliedner et al. 2000). Following our best estimates for oxygen fugacity, we 987 performed models at NNO. Initial water contents were assumed to be 2 wt.% H₂O, 988 which is within the range of primitive (Mg#= 65-74) melt inclusion concentrations 989 from basalts in the Mexican, Cascade, and Kamchatkan arcs (1.0-5.2 wt.% H₂O; 990 Sisson and Layne 1993; Cervantes and Wallace 2003; Portnyagin et al. 2007) and 991 992 reproduces the H₂O content (2.9-3.5 wt.%) of the HLMC gabbros after completion 993 of Stage 1 crystallization. Though the xenolith-based and alphaMELTS models produce the HLMC parental melt composition from slightly different primitive 994 995 starting compositions and diverge after the melt reaches the HLMC parental melt 996 composition, the orthopyroxenite and clinopyroxenite cumulates produced by the 997 MELTS model are consistent with xenolith compositions from the SNB lower 998 crust (Fig 7). After generation of the HLMC parental melt, we extend both the

- 999 xenolith fractionation and MELTS models to Mg#<51 to assess the melt evolution
- 1000 had it not been extracted from the lower crust (shown as dashed lines in Fig 7), but

1001 this is not included in our polybaric crystallization model.

1002

1003 Stage 2: Upper Crustal Fractionation Model Description

We use a similar stepwise cumulate subtraction approach to model the LLD 1004 of the HLMC parental melt as it fractionally crystallizes in the upper crust. 1005 Average cumulate compositions are calculated for 6 representative lithologies from 1006 HLMC (Table S5) following the fractionation sequence discussed above. The 1007 starting liquid composition is the HLMC parental melt as generated by our lower 1008 crustal fractionation model. We do not use alphaMELTS to model this stage for 1009 two reasons: 1) amphibole is a common fractionating phase in the HLMC and 1010 alphaMELTS currently lacks an amphibole model that accurately reproduces 1011 observed phase assemblages from experimental and natural studies, and 2) the 1012 fractionating assemblage and cumulate compositions can be constrained from field 1013 and geochemical observations. 1014

1015

1016 Proportions of Cumulates Crystallized

1017 Cumulate assemblages are introduced to the fractionation sequence based on1018 the calculated Mg# of melts in equilibrium with orthopyroxene and clinopyroxene

| 1019 | in the lower-crustal xenoliths, determined using mineral-melt Mg-Fe distribution |
|------|--|
| 1020 | coefficients of 0.284 and 0.23, respectively (Beattie 1993; Sisson and Grove 1993). |
| 1021 | Because the garnet-pyroxenite xenoliths span a significant range in bulk-rock Mg# |
| 1022 | (39-83), we split this lithology into four fractionating cumulate groups to more |
| 1023 | accurately replicate crystallization from an evolving melt: group A (Mg# 80.5-83), |
| 1024 | group B (Mg# 74-80), group C (Mg# 62-68), and group D (Mg# 39-59). |
| 1025 | Compositions of cumulates used in this model are averages for each group, and are |
| 1026 | included in the Supplementary Data File. Fractionation of a primitive arc basalt |
| 1027 | during Stage 1 fractionation of lower crustal cumulates produces 16.6% (by mass) |
| 1028 | of orthopyroxenite (\pm clinopyroxene), 10.2% of group A garnet-pyroxenites, 7.0% |
| 1029 | of group B garnet-pyroxenites, and 0.6% of group C garnet-pyroxenites. At this |
| 1030 | point, the melt reaches an Mg# of 51 and ascends into the upper crust where the |
| 1031 | HLMC cumulates begin to crystallize. Stage 1 differentiation results in 33% total |
| 1032 | crystallization of cumulates (by mass), closely replicated by 33% crystallization of |
| 1033 | pyroxenites produced in the alphaMELTS model. The resulting liquid composition |
| 1034 | is input into the Stage 2 model as the HLMC parental melt. |
| 1035 | In accordance with our inferred fractionation sequence, low-Ca norites are |
| 1036 | the first cumulates to crystallize in Stage 2 (3% crystallization of the remaining |

1037 melt mass), followed by high-Ca norite fractionation (9.3%), and low-SiO₂ gabbro

1038 crystallization (53%). As clino- and orthopyroxenes are consistently exsolved in

| 1039 | the gabbros they are unlikely to record the equilibrium melt composition, and we |
|------|---|
| 1040 | therefore infer that the Mg# of the melt was between that of the low-SiO ₂ (Mg#= |
| 1041 | 46) and liquid-like gabbros (Mg#= 49) at the onset of high-SiO ₂ gabbro |
| 1042 | crystallization. High-SiO ₂ gabbro is assumed to crystallize until the liquid-like |
| 1043 | monzonite compositions are replicated at 63 wt.% SiO_2 (20% crystallization by |
| 1044 | mass). Fractionation of cumulate monzodiorites continues from 63 wt.% SiO_2 and |
| 1045 | Mg# 44.7 to 70 wt.% SiO ₂ and Mg# 40 (7.3% by mass). Cumulate monzonites |
| 1046 | then fractionate until SiO ₂ reaches 78 wt.% (2.9% by mass), near the maximum for |
| 1047 | granites in the SNB. |

1048

1049 Model Results

Our polybaric fractionation model reproduces the range of major and trace 1050 element compositions of the HLMC, and when extended to higher SiO₂ contents is 1051 capable of replicating the compositional trends defined by the SNB felsic plutons 1052 (63-78 wt.% SiO₂, Fig 7). In Stage 1, differentiated melts in equilibrium with the 1053 most primitive HLMC minerals are produced by 33% fractionation by mass of a 1054 primitive arc basalt at pressures between 1 and 3 GPa (>30 km). The resulting melt 1055 composition is the HLMC parental melt, which has an H₂O content of 3.3 wt.% 1056 consistent with our estimates from plagioclase hygrometry in the liquid-like 1057 gabbros. During this crystallization stage, the Mg# of the melt drops from 68 to 51 1058

| 1059 | with little change in SiO_2 (49.9 to 48.9 wt.%) due to significant fractionation of |
|------|---|
| 1060 | orthopyroxene and clinopyroxene, with lesser contributions from garnet. Initially, |
| 1061 | 16.6% by mass of orthopyroxenite (with minor websterite) crystallizes, causing a |
| 1062 | decrease in Mg# from 68 to 59 (Fig 7a). Over this interval, MgO in the remaining |
| 1063 | melt decreases from 11.8 to 8.4 wt.%, while all other major oxides increase |
| 1064 | accordingly and ASI remains stable at 0.64. Garnet-pyroxenites (group A, Mg# |
| 1065 | 80.5-83) begin to fractionate at a melt Mg# of 59, in equilibrium with the highest |
| 1066 | Mg# clinopyroxene (Mg#= 65, Fe-Mg distribution coefficient 0.23; Sisson and |
| 1067 | Grove 1993). 10.2% of the melt crystallizes to form group A garnet-pyroxenites, |
| 1068 | causing an increase in melt ASI from 0.64 to 0.71 and decrease in Mg# from 59 to |
| 1069 | 55. Following, 6.3% resulting in a melt Mg# of 51 and ASI of 0.74. At this point, |
| 1070 | the melt is in equilibrium with the most primitive HLMC minerals. If the melt were |
| 1071 | to remain in the lower crust, the group C and D garnet-pyroxenites would proceed |
| 1072 | to crystallize, causing ASI to further increase, as is described below. Within Stage |
| 1073 | 1 crystallization, the MgO content of the melt decreases from 11 to 6.2 wt.% with a |
| 1074 | concurrent increase in FeO from 10 to 10.6 wt.%. This crystallization also causes a |
| 1075 | significant increase in Al_2O_3 from 13.5 to 17.2 wt.% with only a slight increase in |
| 1076 | CaO (9.4 to 10 wt.%), driving the observed rise in ASI. |
| 1077 | The derivative basalt produced at the end of Stage 1 is assumed to |

1077 subsequently ascend, intrude into the upper crust, and further differentiate. The

| 1079 | removal of norite cumulates, dominated by orthopyroxene with minor olivine and |
|------|--|
| 1080 | plagioclase, drives an initial decrease in the melt Mg# from 51 to 47 at relatively |
| 1081 | steady SiO ₂ . CaO, Al ₂ O ₃ , Na ₂ O, and Sr content increase in the melt during this |
| 1082 | interval due to the paucity of accumulated feldspar in the norites. |
| 1083 | Following the onset of low-SiO ₂ gabbro crystallization, MgO, CaO, FeO, |
| 1084 | and TiO ₂ smoothly decrease through the remaining LLD, while K_2O , ASI, and |
| 1085 | incompatible trace elements consistently increase. Fractionation of the low-SiO $_2$ |
| 1086 | gabbros causes a significant increase in the melt SiO_2 content from 49.9 to 56 |
| 1087 | wt.%. The low SiO ₂ (43.9-46.8 wt.%) and high FeO and TiO ₂ (9.6-13.6 and 1.2-1.6 |
| 1088 | wt.%, respectively) content of the low-SiO $_2$ gabbros is due to abundant amphibole |
| 1089 | and Fe-Ti oxides. As such, fractionation of these cumulates drives the SiO_2 |
| 1090 | enrichment and FeO and TiO_2 decrease in the melt. The liquid also decreases in |
| 1091 | MgO and CaO content over this fractionation interval. In contrast to norite |
| 1092 | fractionation, plagioclase is now a major fractionating phase and the Al ₂ O ₃ content |
| 1093 | of the melt increases only slightly during low-SiO ₂ gabbro fractionation, from 18.2 |
| 1094 | to 18.5 wt.%. Na ₂ O and P_2O_5 increase due to crystallization of high-Ca plagioclase |
| 1095 | (~An ₈₀) and a lack of apatite in the low-SiO ₂ gabbros. |
| 1096 | High-SiO ₂ gabbro fractionation continues the trend of increasing SiO ₂ |
| 1097 | content (56-63 wt.%) with a limited decrease in Mg# (47 to 45). Plagioclase is |

abundant in these cumulates and has lower An content (\sim An₅₀) than in the low-

| 1099 | SiO_2 gabbros, causing Al_2O_3 to decrease and the rate of Na_2O increase to shallow. |
|------|---|
| 1100 | Al ₂ O ₃ , Na ₂ O, and P ₂ O ₅ content continue to decrease for the remainder for the LLD. |
| 1101 | When gabbro fractionation ceases, the model has reproduced the most fractionated |
| 1102 | liquid-like monzonite in HLMC with 16.2% of the mass of HLMC parental melt |
| 1103 | remaining, or 10.9% for the primitive arc basalt. |
| 1104 | Fractionation of monzordiorite cumulates from the remaining melt continues |
| 1105 | the increasing trends in SiO ₂ (63-71 wt.%), K_2O (3.0-4.5 wt.%), and Na_2O (3.5-3.7 |
| 1106 | wt.%). The melt Mg# decreases from 45 to 40, along with decreasing MgO, TiO ₂ , |
| 1107 | and FeO primarily due to amphibole and biotite crystallization, and decreasing |
| 1108 | CaO and Al ₂ O ₃ which are depleted by plagioclase and amphibole fractionation. |
| 1109 | This crystallization interval replicates a representative SNB granodiorite |
| 1110 | composition (67.5 wt.% SiO ₂ , Table S5) with 9.9% of the parental melt and 6.6% |
| 1111 | of the primitive melt masses remaining. Monzonite fractionation continues the |
| 1112 | monzodiorite fractionation trends, causing the evolving melt composition to |
| 1113 | increase from 71 to 75 wt.% SiO ₂ . Mg# notably decreases from 40 to nearly 0 as |
| 1114 | MgO is consumed by fractionating amphibole and biotite. At this point, the |
| 1115 | remaining granitic melt mass is 5.8% of the HLMC parental melt and 3.9% of the |
| 1116 | primitive arc basalt. |
| | |

1117 To determine if the HLMC evolved rocks and SNB granodiorites could be 1118 produced by continued lower crustal fractionation, we extend both the lower

| 1119 | crustal (xenolith) crystallization model and the alphaMELTS model to Mg#s<51 |
|------|--|
| 1120 | (Fig 7). Though the starting compositions we use are within the low range of Al_2O_3 |
| 1121 | values found in primitive arc basalts (<13.5 wt. % Al_2O_3) (Schmidt and Jagoutz |
| 1122 | 2017), our models of lower crustal fractionation always produce peraluminous |
| 1123 | and esite liquids (i.e., ASI>1.1 when SiO ₂ <57 wt.%). In our model and in the |
| 1124 | Blatter et al. (2013) experiments, plagioclase does not become a significant |
| 1125 | fractionating phase at >0.9 GPa until 33 to 30% of the initial melt mass is |
| 1126 | remaining. In the xenolith cumulate crystallization model, Al is initially removal |
| 1127 | from the melt by minor incorporation into orthopyroxene and has little effect on |
| 1128 | ASI. Garnet is included in the fractionating cumulates after 17% crystallization, |
| 1129 | but the group A and B garnet-pyroxenites have low Al_2O_3 contents (7.6-8.8 wt.%) |
| 1130 | relative to CaO (12.1-15.7 wt.%), leading to a continuous increase in ASI. Thus, |
| 1131 | the HLMC compositions and non-peraluminous evolved arc melts are inconsistent |
| 1132 | with extensive lower crustal fractionation. |
| 1133 | |

1134 Comparison to Other Upper Crustal Mafic Bodies in the SNB

Numerous mafic bodies are preserved in the SNB upper crust and several
have been investigated for geochronology and geochemistry (Frost 1987; Frost and
Mahood 1987; Coleman et al 1995; Sisson et al. 1996; Ratajeski et al. 2001;
Holland et al. 2013). The Onion Valley (92.1 ± 0.3 Ma), Armstrong Canyon (91.5

| 1139 | \pm 0.1 Ma), and Lake Sabrina (91.1 \pm 0.3 Ma) mafic complexes in the eastern Sierra |
|------|---|
| 1140 | Nevada (Coleman et al. 1995) are broadly coeval with HLMC (90.5-96.5 Ma). |
| 1141 | Mafic magmatism in all four localities was contemporaneous with voluminous late |
| 1142 | Cretaceous felsic intrusions throughout the Sierra Nevada arc (Ducea 2001), |
| 1143 | demonstrating that mafic and felsic magmatism was coeval and possibly cogenetic. |
| 1144 | Geochemical data from the lower crustal Bear Valley Intrusive Suite (Klein |
| 1145 | and Jagoutz 2021) and 10 upper crustal SNB mafic complexes in addition to |
| 1146 | HLMC are presented in Figure 8. All localities are in the eastern SNB from Frost |
| 1147 | (1987) and Sisson et al. (1996) except for the Yosemite samples of Ratajeski et al. |
| 1148 | (2001) and the Ash Mountain complex of Holland et al. (2013). Amphibole- |
| 1149 | gabbros (\pm olivine) and amphibole-bearing diorites are common throughout the |
| 1150 | mafic complexes, indicative of crystallization from hydrous parental melts. In |
| 1151 | particular, the parental melt to the Onion Valley complex is estimated to have 6 |
| 1152 | wt.% H_2O (Sisson et al. 1996). The most primitive cumulates in the Onion Valley |
| 1153 | complex are olivine-hornblendites and the intrusion is generally pyroxene-poor. |
| 1154 | This is in contrast to the orthopyroxene-dominated norites of HLMC, which |
| 1155 | indicate crystallization from a less hydrous parental melt (estimated 3.1 wt.% |
| 1156 | H_2O). In both cases, elevated water contents likely facilitate the ascent of the low- |
| 1157 | Mg, high-Al basalts into the upper crust due to their decreased density and |
| 1158 | viscosity (Sisson et al. 1996). |

| 1159 | Layered cumulates and internal compositional ranges from gabbro to diorite |
|------|--|
| 1160 | within each body (Frost 1987; Sisson et al. 1996; McCarthy and Müntener 2016) |
| 1161 | suggest that in-situ upper crustal fractionation is common in upper crustal mafic |
| 1162 | complexes. Olivines from the Onion Valley complex have a maximum Mg# of |
| 1163 | 81.7 (in equilibrium with melt Mg# = 52), and sheeted sills interpreted as |
| 1164 | representative of melt compositions have Mg#s ranging from 41 to 54.5 (Sisson et |
| 1165 | al. 1996). The most primitive sill and the olivine-melt estimates are slightly more |
| 1166 | primitive, but still comparable to the HLMC parental melt (Mg# = 51), suggesting |
| 1167 | that the parental melts to these two bodies were fractionated to a similar degree in |
| 1168 | the lower crust prior to ascent. Mineral chemistry only exists for the Onion Valley |
| 1169 | and Hidden Lakes mafic complexes, rendering it is difficult to calculate the Mg# of |
| 1170 | parental melts for the other complexes. However, maximum bulk-rock Mg#s in |
| 1171 | each complex range from 53 to 76.5, and therefore did not fractionate from a |
| 1172 | primitive basalt (Mg# of cumulates > 80). Thus lower crustal differentiation is |
| 1173 | required to generate low-Mg basalts in order to produce these upper crustal mafic |
| 1174 | intrusions. Derivation of the upper crustal mafic complexes from evolved basalts |
| 1175 | and the presence of upper crustal cumulate lithologies demonstrate that polybaric |
| 1176 | fractionation is a common process in generating the range of compositions in SNB |
| 1177 | mafic complexes. |

| 1178 | All mafic complex samples have ASI values less than 1, indicating that the |
|------|--|
| 1179 | mafic melts that escape the lower crust are generally not peraluminous. Samples |
| 1180 | with high ASI relative to their SiO ₂ content (ASI = $\sim 0.8-0.97$, SiO ₂ = 40.7-50.3 |
| 1181 | wt.%) are likely cumulates, and thus reflect plagioclase accumulation rather than |
| 1182 | parental melts with initially high ASI values. Lower crustal fractionation of a |
| 1183 | primitive arc basalt creates peraluminous melts at 51 wt.% SiO_2 (our model) to 56 |
| 1184 | wt.% SiO ₂ (Blatter et al. 2013) by ~55% crystallization. Thus, melts that generate |
| 1185 | the mafic complexes must escape the lower crust (>0.7 GPa) before evolving to |
| 1186 | andesitic compositions. Because some lithologies found in these complexes have |
| 1187 | $SiO_2 > 56$ wt.%, upper crustal differentiation must occur to create these |
| 1188 | compositions after extraction from the lower crust. |
| 1189 | Though mineral compositional data is limited to two complexes, those that |
| 1190 | do exist define a narrow range of parental melt compositions (Mg# = $51-54.5$). The |
| 1191 | range of bulk-rock cumulate compositions from other complexes are similar in |
| 1192 | Mg# to the Onion Valley and Hidden Lakes mafic complexes, suggesting that all |
| 1193 | upper crustal mafic complexes may be derived from melts that are fractionated to a |
| 1194 | similar degree in the lower crust. Additionally, the SiO_2 content of hydrous |
| 1195 | parental melts cannot exceed 56 wt.% without becoming peraluminous during |
| 1196 | fractional crystallization at >0.7 GPa. Therefore, we suggest that there may be a |
| 1197 | narrow window of rheologic properties that allow these hydrous basalts to ascend. |

- 1198 For example, the hydrous, but still relatively $low-SiO_2$ nature of these basalts
- 1199 reduces their viscosity and decreases their density, enhancing their ability to
- 1200 ascend via diking (Sisson et al. 1996).
- 1201

1202 Production of Sierra Nevada Batholith Granitoids

Our polybaric fractionation model results support the hypothesis that the 1203 HLMC is an example of upper crustal fractional crystallization of a low-Mg basalt 1204 in a continental arc and that lower crustal fractionation is required to produce the 1205 HLMC parental basalt. We show that arc-like granodiorite and granite 1206 compositions can be generated by this two-stage crystallization process and thus 1207 polybaric crystallization can contribute to the generation of non-peraluminous 1208 granitoids in the SNB. However, volumes of mafic (noritic to gabbroic) cumulates 1209 at upper crustal depths (0.1-0.4 GPa) are relatively small. We posit that the 1210 fractionation of a low-MgO basalt deeper than the exposures of much of the SNB, 1211 yet at depths sufficiently low to allow for plagioclase saturation (<0.7 GPa), could 1212 generate evolved non-peraluminous melts. Crystallization of "damp" basalt to 1213 andesites and enhancement of orthopyroxene stability could also contribute to 1214 limiting the increase in ASI of differentiating melts at high pressures (Rezeau et 1215 al., 2021). This is supported by gabbro and gabbronorite lithologies with a 1216 maximum Mg# identical to the HLMC (Mg# = 73) in the $\sim 0.8-0.9$ GPa section of 1217

| 1218 | the Bear Valley Intrusive Suite (BVIS; Klein and Jagoutz 2021). Because the BVIS |
|------|--|
| 1219 | lithologies did not crystallize from a primitive mantle melt, an initial stage of |
| 1220 | fractionation is required below ~1 GPa, analogous to Stage 1 in our model for the |
| 1221 | HLMC. Throughout the batholith, Stage 2 fractionation may occur below |
| 1222 | exposures of the bulk of the SNB (>0.4 GPa) to produce granodioritic melts |
| 1223 | consistent with SNB compositions, which proceed to intrude into the upper crust. |
| 1224 | This reconciles the discrepancy between experimental and observational data, in |
| 1225 | which experiments indicate that melts produced by lower crustal fractionation |
| 1226 | rapidly become peraluminous (Cawthorn and Brown 1976; Cawthorn and O'Hara |
| 1227 | 1976; Blatter et al. 2013; Nandedkar et al. 2014; Müntener and Ulmer 2018; Ulmer |
| 1228 | et al. 2018), yet lower crustal ultramafic cumulates are ubiquitous in arc crustal |
| 1229 | cross sections (Debari and Sleep 1991; Greene et al. 2006; Otamendi et al. 2012; |
| 1230 | Jagoutz 2014; Walker et al. 2015; Guo et al. 2020). HLMC has captured a local |
| 1231 | example of upper crustal differentiation, indicating that granodiorite compositions |
| 1232 | consistent with the SNB can be produced by a polybaric fractionation processes |
| 1233 | with a range of possible depths for fractionation of a low-MgO basalt. |
| 1234 | The volumes of intermediate to evolved (>60 wt.% SiO ₂) melt produced by |
| 1235 | fractional crystallization will be small compared to the volumes of cumulates, so |
| 1236 | polybaric crystallization may not be the only mechanism for the generation of arc |
| 1237 | granodioites. Previously proposed mixing models for the SNB (e.g., Reid et al. |

1238 1983; Frost and Mahood 1987; Sisson et al. 1996), require both a high-SiO₂ melt, and an evolved basalt similar to the HLMC parental melt. Although the mechanical 1239 viability of mixing felsic and mafic melts is limited, these models also resolve the 1240 problem of peraluminous melt compositions generated by lower crustal 1241 fractionation, because the homogenized products between non-peraluminous, low-1242 Mg basalts and mildly peraluminous (i.e, ASI<1.1) granitic melts are consistent 1243 with batholith granitoid compositions (Blatter et al. 2013). Our modeling identifies 1244 a potential low-Mg basaltic end member composition (the HLMC parental melt) 1245 produced by lower crustal fractionation. We show that evolved basalts must be 1246 generated in the lower crust, and are thus available to participate in both mixing 1247 and polybaric crystallization processes that contribute to the production of 1248 batholith-scale volumes of granodiorite. 1249

1250

1251 Conclusions

1252 The Late Cretaceous Hidden Lakes mafic complex is a preserved example of 1253 intrusion of melt into the upper crust of the Sierra Nevada batholith 1254 contemporaneously with more voluminous felsic magmas. Bulk-compositions of 1255 lithologies ranging from olivine-norites to monzonites are consistent with 1256 fractional crystallization from a moderately hydrous basalt (2.9-3.5 wt% H₂O) at 1257 ~0.3 GPa, and oxidizing conditions (~NNO). The parental melt to the HLMC is a

1258 low-Mg basalt produced through differentiation in the lower crust prior to1259 intruding into the upper crust.

We model this polybaric fractional crystallization process in two-stages, 1260 using cumulate compositions from SNB deep crustal xenoliths, accompanied by 1261 alphaMELTS, to calculate the composition of an evolving primitive arc basalt in 1262 the lower crust (Stage 1) and constructing an upper crustal fractionation based on 1263 measured cumulate and liquid-like sample compositions (Stage 2). Our model 1264 indicates 33% mass fractionation in the lower crust, followed by extraction to 1265 upper crustal pressures and subsequent fractionation to form all HLMC cumulates 1266 and melt compositions. Polybaric fractionation of mantle-derived basalt to produce 1267 granodiorite resolves the disagreement between the prevalence of lower crustal 1268 cumulates in arcs and experimental data that suggest lower crustal fractionation 1269 will produce peraluminous melts inconsistent with arc batholith compositions. 1270 The HLMC contains rock types that are lithologically and compositionally 1271 similar to those other upper crustal mafic complexes of the SNB, suggesting that 1272 these intrusions crystallized from hydrous, fractionated basalts of comparable 1273 compositions. Consequently, their composition may be governed by unifying 1274 1275 rheologic characteristics that facilitate their ascent from the lower crust such as 1276 reduced density and viscosity.

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Table 1: Summary of Modal Mineralogies

| | nori te 0- | gabbro (coarse- grained) | gabbro (medium- fine grained) | monzodio rite | monzo nite | anortho site | |
|------------|------------------|--------------------------------|--|------------------|---------------|-----------------|----------|
| olivine | <1ª | _ | _ | _ | _ | _ | |
| orthopyro | 15- | | | | | | |
| xene | 75 | — | ~5 | — | 0-<1 | - | |
| clinopyro | | | | | | | |
| xene | 1-5 | 5-20 | 5-15 | 0-2 | 0-<1 | | <i>v</i> |
| plagioclas | 15- | | | | | | |
| e | 40 | 45-70 | 65-75 | 60-70 | 40-60 | 90-95 | |
| amphibol | 0- | | | | | | |
| e | 20 | 25-50 | 5-20 | 5-15 | 0-5 | _ | |
| | 5- | - 10 | - 10 | | 10.00 | - 10 | |
| biotite | 15 | 5-10 | 5-10 | 15-20 | 10-20 | 5-10 | |
| К- | 0- | | | | | | |
| feldspar | <1 | <1 | <1 | 10-15 | 20-35 | — | |
| quartz | — | — | - | 0-2 | 0-5 | — | |

^a all modal mineralogies are given in percent and estimated visually, with ranges reflecting variability within each lithologic group. Uncertainties are roughly 10% of reported values.

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| | Method | norites | gabbros | monzodiorite s | monzonite s |
|--------------------------------------|-------------------------------------|---|---|-------------------|----------------|
| Temperature (°C) | Ca-in-opx ^a | 900- 1170 | 930-1100 | 970- 1030 | 850-1010 |
| | plag-amph pairs ^b | 865- 950 (oikocrysts) 700- 765 (non- | 740- 815 (magmatic amph) 1040-1120 (pseudomorph | 815- 870 | 800- 850 |
| | | oikocrystic) | ic amph) | | |
| Pressure (GPa) | Al-in- amph ^c | [0.32] ^f | [0.32] | [0.32] | 0.32 |
| H ₂ O (wt.%) | plag hygrometr y ^d | | 2.9- 3.5 | 3 | 6-7 |
| $fO_2 (\Delta NNO, \pm 1 \log unit)$ | biotite ^e | | | | 0-0.6 |

Table 2: Calculated Crystallization Conditions

^a Method of Köhler and Brey (1990)

^b Method of Blundy and Holland (1990)

^c Method of Mutch et al. (2016)

^d Method of Waters and Lange (2015)

^e Method of Bucholz et al. (2018)

^f Values in brackets indicate those that were not calculated from that particular rock type due to mineral assemblage constraints, but are assumed to be constant throughout the fractionation sequence (e.g. pressure and fO_2)