

2 Phase Relations, Reaction Sequences and Petrochronology

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INTRODUCTION

At the core of petrochronology is the relationship between geochronology and the petrological evolution of major mineral assemblages. The focus of this chapter is on outlining some of the available strategies to link inferred reaction sequences and microstructures in metamorphic rocks to the ages obtained from geochronology of accessory minerals and datable major minerals. Reaction sequences and mineral assemblages in metamorphic rocks are primarily a function of pressure (P), temperature (T) and bulk composition (X). Several of the major rock-forming minerals are particularly sensitive to changes in P – T (e.g., garnet, staurolite, biotite, plagioclase), but their direct geochronology is challenging and in many cases not currently possible. One exception is garnet, which can be dated using Sm–Nd and Lu–Hf geochronology (e.g., Baxter et al. 2013). Accessory mineral chronometers such as zircon, monazite, xenotime, titanite and rutile are stable over a relatively wide range of P – T conditions and can incorporate enough U and/or Th to be dated using U–Th–Pb geochronology. Therefore,

34 linking the growth of P - T sensitive major minerals to accessory and/or major mineral
35 chronometers is essential for determining a metamorphic P - T - t history, which is itself critical
36 for understanding metamorphic rocks and the geodynamic processes that produce them (e.g.,
37 England and Thompson, 1984; McClelland and Lapen 2013; Brown, 2014).

38 Linking the ages obtained from accessory and major minerals with the growth and
39 breakdown of the important P - T sensitive minerals requires an understanding of the
40 metamorphic reaction sequences for a particular bulk rock composition along a well-constrained
41 P - T evolution. Fortunately, the phase relations and reaction sequences for the most widely
42 studied metamorphic protoliths (e.g., pelites, greywackes, basalts) can be determined using
43 quantitative phase equilibria forward modelling (e.g., Powell and Holland 2008). Comprehensive
44 activity-composition models of the major metamorphic minerals in large chemical systems (e.g.,
45 White et al. 2014a) allow the calculation of phase proportions and compositions for a given rock
46 composition along a metamorphic P - T path. For accessory minerals, subsolidus growth and
47 breakdown can be modelled in some cases using phase equilibria modelling (e.g., Spear 2010;
48 Spear and Pyle 2010). Suprasolidus accessory mineral behaviour can be investigated by coupling
49 phase equilibria modelling with the experimental results of accessory mineral solubility in melt
50 (Kelsey et al. 2008; Yakymchuk and Brown 2014b). This technique provides a basic framework
51 for interpreting the geological significance of accessory mineral ages in suprasolidus
52 metamorphic rocks.

53 In this chapter, we use phase equilibria modelling techniques to investigate the reaction
54 sequences for three common rock types (pelite, greywacke, MORB) along several P - T paths and
55 explore how these sequences relate to accessory mineral growth and dissolution with a particular
56 focus on zircon and monazite and to a lesser extent apatite. First, we review the important major

57 rock-forming minerals that are used to link metamorphic reaction sequences to trace element
58 chemistries in accessory minerals. Second, we summarize the current understanding of the
59 controls of accessory mineral growth and breakdown in metamorphic rocks during a P - T
60 evolution. Third, we use phase equilibria modelling to examine the reaction sequences for
61 common rock types along several schematic P - T paths and discuss the implications for
62 petrochronology. Finally, we examine some of the complicating factors for reconciling the
63 behaviour of accessory minerals in natural systems with the predictions from phase equilibria
64 modelling.

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MAJOR MINERALS

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Understanding the growth and consumption of the major rock-forming minerals is important in accessory mineral petrochronology for four reasons. First, major minerals may contain significant quantities of the essential structural constituents of accessory minerals commonly used as chronometers. In some cases, accessory mineral can grow directly from the breakdown of major minerals. Examples of this are zircon growth from the release of Zr during the breakdown of garnet (Fraser et al. 1997; Degeling et al. 2001) and ilmenite (Bingen et al. 2001). Second, these minerals may represent important repositories for trace elements, and thus the growth and breakdown of major minerals will influence the availability of these elements for incorporation by growing accessory minerals. Third, the major minerals are important hosts for inclusions of accessory minerals. The breakdown of the major minerals may liberate included accessory minerals into the reaction volume of the rock or alternatively may sequester these minerals away allowing their preservation when they would otherwise be consumed in a reaction

79 sequence (e.g., Montel et al. 2000). Fourth, the microstructural relationships between accessory
80 and major minerals provide context for delineating the P – T history of a metamorphic rock.

81 Garnet and plagioclase are the major minerals most commonly used in petrochronology
82 because of their distinctive trace element behaviour. Linking their growth and breakdown to ages
83 obtained from accessory mineral chronometers requires an understanding of the bulk
84 composition controls on their stabilities. Below, we outline the controls on the growth and
85 breakdown of garnet and plagioclase, which are of particular importance for petrochronology
86 studies. Our focus is on linking mineral growth to metamorphic reactions and we do not discuss
87 the minerals that are important for thermochronology studies (e.g., amphibole, biotite,
88 muscovite, K-feldspar).

89

90 **Garnet**

91 Garnet is one of the most useful minerals for constraining metamorphic grade and its high
92 density and strong partitioning of cations forms the basis of many useful thermobarometers (e.g.,
93 Spear 1995; Caddick and Kohn 2013). It is a common metamorphic mineral for many different
94 protoliths (pelites, mafic rocks, ultramafic rocks, calc-silicates) and has been extensively used in
95 petrochronology studies (e.g., Vance and O’Nions 1990; Vance and Mahar 1998; Harris et al.
96 2004). Coupling P – T estimates from garnet with monazite U–Pb geochronology can also be used
97 to constrain P – T – t paths (e.g., Foster et al. 2000; Gibson et al. 2004; Dragovic et al. 2016).
98 Garnet can also be directly dated using Sm–Nd and Lu–Hf geochronology (e.g., Baxter et al.
99 2013; Baxter et al. 2017, this volume). Although linking P – T information from garnet-bearing
100 assemblages with garnet geochronology is very powerful (e.g., Mulcahy et al. 2014; Dragovic et
101 al. 2012, 2015), direct dating of garnet is challenging and is not yet extensively used.

102 An important control on the stability of garnet during metamorphism is the bulk rock
103 MnO content (e.g., Symmes and Ferry 1992; Mahar et al. 1997; White et al. 2014b). For pelites,
104 higher bulk rock Mn concentrations stabilize garnet at lower temperatures in the greenschist
105 facies and at lower pressure in the amphibolite facies (White et al. 2014b). The ratio of Fe/Mg is
106 also an important control on garnet growth. For example, relatively magnesium-rich pelites have
107 restricted garnet stability fields (e.g., White et al. 2014a) and in very Mg-rich bulk rock
108 compositions garnet may not even grow along common P - T paths (e.g., Fitzsimons et al. 2005).
109 Therefore, depending on the bulk composition of the rock, garnet can be first stabilized in the
110 greenschist facies, the amphibolite facies or not at all.

111 Garnet is the most important major mineral sink for the heavy rare earth elements
112 (HREE) and yttrium in metamorphic rocks (Bea et al. 1994). Accessory minerals such as zircon,
113 monazite, xenotime, apatite, epidote and allanite are also important repositories for the HREE
114 and Y. Thus, the growth and breakdown of garnet plays an important role in the HREE and Y
115 budgets in rocks and links the trace element chemistry of accessory mineral chronometers to the
116 P - T information obtained from garnet-bearing metamorphic assemblages (Pyle et al. 2001;
117 Foster et al. 2002; Kohn et al. 2004; Rubatto 2002; Rubatto and Hermann 2007; Taylor et al.
118 2015). In general, the equilibrium distribution coefficients of the HREE between garnet and
119 zircon are close to unity for Gd to Lu at high temperature (Taylor et al. 2015). For Yttrium,
120 concentrations in monazite are ~ 1.5 orders of magnitude higher than in garnet (Bea et al. 1994).
121 Xenotime has Y concentrations that are ~ 2 orders of magnitude higher than in monazite (Pyle et
122 al. 2001). However, in most cases, garnet is substantially more modally abundant than monazite
123 and xenotime and thus the breakdown or growth of garnet will exert first-order controls on the Y
124 and HREE budget of the rock. Zircon and monazite crystallization during garnet growth will

125 result in relatively low concentrations of HREE and Y in these minerals because these elements
126 are partitioned into garnet. By contrast, garnet breakdown during accessory mineral growth can
127 result in HREE- and Y-enriched trace element concentrations of the new growth zones of
128 accessory minerals (e.g., Rubatto 2002; Foster et al. 2002, 2004).

129

130 **Plagioclase**

131 Plagioclase is extremely common in metamorphic rocks and is stable over a wide range
132 of P - T conditions. Plagioclase breakdown with increasing pressure is an important part of the
133 reaction sequences for high-pressure metamorphic rocks and eclogites. Similar to garnet,
134 plagioclase is commonly a key mineral in thermobarometers that provides important P - T
135 information in many different rock types (Ghent 1976; Wu 2015; Molina et al. 2015). For
136 petrochronology, plagioclase (and to a lesser extent K-feldspar) strongly partitions Eu and Sr
137 over the other major rock-forming minerals (e.g., Gromet and Silver 1983). The growth and
138 breakdown of plagioclase can be tied to Eu anomalies in rare earth element patterns of accessory
139 minerals (Rubatto et al. 2013; Holder et al. 2015; Regis et al. 2016). Strontium concentrations
140 have been used to link accessory mineral growth to the timing of plagioclase breakdown; this is a
141 particularly important relationship in ultrahigh pressure metamorphic rocks (Finger and Krenn
142 2007; Kylander-Clark et al. 2013; Holder et al. 2015).

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ACCESSORY MINERALS

145 Like the major minerals, accessory minerals are involved in the reaction sequence
146 experienced by metamorphic rocks. The behaviour of some accessory minerals used in
147 petrochronology such as epidote, titanite and rutile can be quantified along a P - T evolution using

148 phase equilibria modelling. However, the behaviour of zircon and monazite—the most
149 commonly used mineral chronometers—is more difficult to quantify with current phase
150 equilibria modelling techniques (e.g., Spear and Pyle 2010; Kelsey and Powell 2011) as they
151 contain key elements commonly not considered in model chemical systems.

152

153 **Epidote**

154 Epidote is common in low–medium pressure metabasites, metamorphosed intermediate
155 rocks and calc-silicates (e.g., Grapes and Hoskin 2004) as well as in high- to ultrahigh-pressure
156 metamorphic rocks (e.g., Enami et al. 2004). The chemical controls on epidote stability are
157 mainly bulk rock concentrations of Ca and Al as well as f_{O_2} (e.g., Enami et al. 2004). The
158 epidote-group minerals can be important repositories for Zr (e.g., Frei et al. 2004; Kohn et al.
159 2015) and LREE in metamorphic rocks (Frei et al. 2004; Janots et al. 2008). Therefore, the
160 breakdown of Zr-rich epidote may have the potential to generate metamorphic zircon. Coupled
161 allanite and epidote breakdown may produce new monazite (e.g., Janots et al. 2008) if there is
162 also a source of phosphorus such as xenotime or apatite.

163

164 **Titanite**

165 Titanite is common in mafic rocks and in some low-grade metasedimentary rocks and it
166 can be used as a barometer in metabasites (e.g., Kapp et al. 2009). An important factor in the
167 stability of titanite is the relative activities of Ca and Al as influenced by the bulk rock ratio of
168 Ca to Al and the accompanying mineral assemblage. High Ca activities favour titanite over
169 ilmenite and high Al activities favour anorthite (and ilmenite) over titanite (Frost et al. 2001).
170 Consequently, titanite is a common mineral in low-Al compositions such as calc-silicate and

171 mafic rocks whereas aluminous metasedimentary rocks generally have ilmenite as the main Ti-
172 bearing mineral above the greenschist facies.

173 Titanite can be directly dated using U–Pb geochronology, but there is some uncertainty
174 about the closure temperature for Pb, which is generally considered to be around 600°C (Warren
175 et al. 2012; Spencer et al. 2013; Stearns et al. 2015; Kirkland et al. 2016), depending on cooling
176 rate. Diffusion profiles of trace elements in titanite can be used to determine the timing and
177 duration of cooling (e.g., geospeedometry) and this is covered in more detail in Kohn (2017, this
178 volume). Titanite thermometry uses the Zr concentration of titanite to estimate temperature
179 (Hayden et al. 2008). However, application of this thermometer requires an estimate of titania
180 activity (a_{TiO_2}), which, in the absence of rutile, is difficult to constrain and can vary along a
181 metamorphic evolution depending on the reaction sequences and mineral assemblages as
182 discussed below.

183

184 **Rutile**

185 Rutile is commonly stable in relatively reduced bulk rock compositions (e.g., Diener and
186 Powell 2010) and is particularly useful in high-pressure metamorphic rocks (Zack and Kooijman
187 2017). Similar to titanite, rutile can be directly dated with U–Pb geochronology (e.g., Mezger et
188 al. 1989; Clark et al. 2000; Zack et al. 2011; Ewing et al. 2015) and it can be used as a
189 geospeedometer (Smye and Stockli 2014). One of the main uses of rutile is as a thermometer that
190 uses the Zr concentration of rutile to estimate temperature (Zack et al. 2004; Watson et al. 2006;
191 Ferry and Watson 2007; Tomkins et al. 2007; Hofmann et al. 2013; Taylor-Jones and Powell
192 2015). These applications are discussed in more detail by Zack and Kooijman (2017, this
193 volume).

194 Metamorphic reaction sequences with or without rutile have important implications for
195 applying the Ti-in-quartz thermometer (e.g., Chambers and Kohn 2012; Ashley and Law 2015)
196 and the Ti-in-zircon thermometer (e.g., Ferry and Watson 2007), both of which use a_{TiO_2} as a
197 variable. In rutile-bearing systems a_{TiO_2} is buffered at 1.0. In rutile-absent systems, application of
198 the thermometer requires an estimate of a_{TiO_2} that is less than 1.0. Figure 1 shows the difference
199 of calculated temperatures using the Ti-in-quartz thermometer (Wark and Watson 2006) and the
200 Ti-in-zircon thermometer (Ferry and Watson 2007) as a function of temperature for a range of
201 a_{TiO_2} values.

202 For example, consider an amphibolite-facies rock at 650°C with a true a_{TiO_2} value 0.6. If
203 an a_{TiO_2} value of 1.0 is assumed for Ti-in-quartz thermometry, the result would underestimate the
204 true temperature by ~50°C (Fig. 1a). Similarly, using an a_{TiO_2} value of 1.0 when the true value is
205 0.6, the Ti-in-zircon thermometer would underestimate the true temperature by ~40°C (Fig 1b).
206 Phase equilibria modelling provides one method of determining a_{TiO_2} in metamorphic rocks at
207 different stages of a P – T evolution in rutile-absent rocks (e.g., Ashley and Law 2015). We
208 explore this in more detail during our discussion of particular reaction sequences.

209

210 **Zircon**

211 Zircon is the main repository of Zr in most igneous and metamorphic rocks and is the
212 most widely used accessory mineral for U–Pb geochronology (Hoskin and Schaltegger 2003;
213 Rubatto 2017, this volume). Many metamorphic rocks contain relict zircon that may be detrital
214 or igneous in origin and may also be inherited from a previous metamorphic event. For
215 petrochronology, metamorphic zircon can be used to date different portions of a P – T evolution
216 depending on the growth mechanisms. The main processes that increase the mode of zircon in

217 metamorphic rocks include solid-state growth and the crystallization from anatectic melt. In
218 some cases, zircon may precipitate from hydrothermal fluids (e.g., Schaltegger, 2007).

219 Solid-state zircon growth can occur from the breakdown of other Zr-rich minerals such as
220 garnet (Fraser et al. 1997; Degeling et al. 2001), ilmenite (Bingen et al. 2001), rutile (Ewing et
221 al. 2014) and possibly amphibole (Sláma et al. 2007). This zircon can be used to date major
222 metamorphic assemblage changes during prograde (e.g., Fraser et al. 1997) or retrograde
223 metamorphism (e.g., Degeling et al. 2001) depending on when the Zr-rich mineral breaks down
224 during the reaction sequence. However, major minerals such as garnet, rutile and hornblende can
225 accommodate more Zr as temperature increases and, in general, zircon will be consumed and
226 prograde zircon growth is expected to be limited (Kohn et al. 2015).

227 The crystallization of anatectic melt typically drives new zircon growth in suprasolidus
228 metamorphic rocks (Watson 1996; Roberts and Finger 1997; Schaltegger et al. 1999; Vavra et al.
229 1999; Hermann et al. 2001; Kelsey et al. 2008; Yakymchuk and Brown 2014b). Experimental
230 studies show that the concentration of Zr in melt needed to maintain equilibrium with zircon
231 increases with temperature (e.g., Harrison and Watson 1983; Gervasoni et al. 2016) and the
232 compositional parameter M , which is the cation ratio of $[\text{Na} + \text{K} + 2\text{Ca}] / [\text{Al} \times \text{Si}]$ (Watson and
233 Harrison 1983; Boehnke et al. 2013). Generally, the concentration of Zr needed for saturation of
234 intermediate melts (high M values) is higher than for more felsic melts (low M values). In
235 suprasolidus metamorphic rocks, the most important factor is the amount of anatectic melt
236 present in the system (Kelsey et al. 2008; Kelsey et al. 2011; Yakymchuk and Brown 2014b). As
237 the fraction of melt increases during prograde metamorphism, zircon is expected to break down
238 to maintain Zr saturation of the melt in an equilibrated system. Consequently, prograde zircon
239 growth is expected to be limited above the solidus. In general, melt crystallization during cooling

240 from peak T is expected to be the main mechanism for zircon growth in suprasolidus
241 metamorphic rocks. This is supported by ranges of concordant ages that reflect protracted zircon
242 growth during melt crystallization from peak T in migmatites (e.g., Korhonen et al. 2013b,
243 2014).

244 New zircon growth can also occur at the expense of pre-existing zircon with no change in
245 the mode of zircon in the rock. Recrystallization of metamict zircon in the presence of a fluid
246 may be an important factor for some prograde zircon (e.g., Rubatto and Hermann 2003; Hay and
247 Dempster 2009). Ostwald ripening has been proposed as a mechanism that could produce
248 prograde growth of zircon in suprasolidus metamorphic rocks (Vavra et al. 1999; Nemchin et al.
249 2001; Kawakami et al. 2013). Ostwald ripening (or second phase coarsening) is a process
250 whereby small solids are preferentially dissolved and precipitate on existing larger solids to
251 reduce the total surface free energy (e.g., Tikare and Cawley 1998). While this process has been
252 studied and debated for the major minerals in metamorphic rocks (Miyazaki 1991, 1996; Carlson
253 1999, 2000) it has not been as extensively studied for accessory minerals—an exception being
254 Nemchin et al. (2001). Nonetheless, it is a possible mechanism for prograde growth of zircon in
255 suprasolidus metamorphic rocks, though involving no net modal increase in zircon.

256 Zircon can also be used as a thermometer in igneous and metamorphic rocks. The Ti-in-
257 zircon thermometer has been increasingly applied to high-temperature metamorphic rocks over
258 the last decade. Kelsey and Hand (2015) compiled Ti-in-zircon temperatures from UHT rocks
259 and noted that 62% of the results fall below the UHT threshold of 900°C. One of the possible
260 reasons for this discrepancy is that some UHT rocks do not contain rutile and are therefore
261 undersaturated in TiO_2 ($a_{\text{TiO}_2} < 1.0$). Values of a_{TiO_2} can range from 1.0 when rutile is in
262 equilibrium with growing zircon to values as low as 0.6 (e.g., Hiess et al. 2008). Using a value of

263 1.0 when rutile is absent will underestimate the true temperature. Again, the presence and
264 absence of rutile during a reaction sequence has important consequences for applying mineral
265 thermometers in metamorphic rocks.

266

267 **Monazite**

268 Monazite is generally more reactive than zircon in subsolidus metamorphic rocks and has
269 been extensively used for U–Pb geochronology in aluminous bulk compositions (e.g., Parrish
270 1990; Engi 2017, this volume). Monazite can be detrital in origin (Smith and Barreiro 1990;
271 Kingsbury et al. 1993; Suzuki et al. 1994; Rubatto et al. 2001), produced through solid-state
272 reactions (e.g., Rubatto et al. 2001; Wing et al. 2003), precipitated from a fluid (e.g., Ayers et al.
273 1999) or can crystallize from anatectic melt (e.g., Stepanov et al. 2012).

274 Solid-state monazite growth during prograde metamorphism occurs from the breakdown
275 of LREE-rich precursors, which can include: allanite for high-Ca bulk compositions (Spear and
276 Pyle 2010; Wing et al. 2003; Finger et al. 2016), LREE-rich clays (Copeland et al. 1971), Th or
277 LREE oxides and hydrous phosphates (Spear and Pyle 2002). The most studied solid-state
278 reaction is the growth of monazite at the expense of allanite in bulk compositions with sufficient
279 Ca and LREE to grow allanite at lower grade (e.g., Janots et al. 2008). In natural examples, this
280 reaction has been spatially correlated with the garnet-in isograd (Catlos et al. 2001; Foster et al.
281 2004), the staurolite-in isograd (Smith and Barriero 1990; Kohn and Malloy 2004; Corrie and
282 Kohn 2008) and the kyanite or sillimanite-in isograds (Wing et al. 2003; Štípská et al. 2015).

283 Spear (2010) used phase equilibria modelling to examine the P – T conditions of the
284 allanite to monazite transition in several bulk rock compositions relative to the average pelite of
285 Shaw (1956). The reaction boundaries for monazite-in at the expense of allanite for these various

286 compositions are summarized in Figure 2. A higher bulk rock CaO concentration allows allanite
287 to persist to higher temperatures whereas a higher Al₂O₃ concentration stabilizes monazite at
288 lower temperatures. However, the results of this modelling of monazite growth in rocks with
289 variable bulk rock Al₂O₃ concentrations by Spear (2010) are different to those observed in some
290 natural examples. For example, Wing et al. (2003) found that pelites from New England with
291 relatively elevated Ca and/or Al concentrations can preserve allanite to higher temperatures.
292 Gasser et al. (2012) found no link between bulk rock concentrations of CaO and Al₂O₃ and the
293 timing of monazite growth. Therefore, the bulk compositional controls of monazite growth at the
294 expense of allanite are not always clear, but it appears that monazite can grow at a range of
295 temperatures and pressures along a prograde *P–T* path.

296 The relative amounts of iron and magnesium in a bulk rock composition may also play an
297 important role in the reactions that produce subsolidus monazite. For example, Fitzsimons et al.
298 (2005) showed that monazite in pelitic schists from Western Australia was generated during
299 garnet-breakdown to staurolite and was interpreted to record the timing of peak metamorphism.
300 However, samples that were too magnesium-rich to grow garnet or too iron-rich to grow
301 staurolite at the expense of garnet yielded older monazite ages. Fitzsimons et al. (2005)
302 interpreted these older ages to represent greenschist-facies monazite growth.

303 In suprasolidus metamorphic rocks, monazite breakdown and growth is controlled mainly
304 by dissolution into and crystallization from anatectic melt. Experimental studies demonstrate that
305 monazite dissolution into melt is a function of temperature, pressure, and the bulk composition of
306 the melt (Montel 1986; Rapp and Watson 1986; Rapp et al. 1987; Skora and Blundy 2012;
307 Stepanov et al. 2012; Duc-Tin and Keppler 2015). The solubility of monazite increases with
308 higher H₂O concentrations in melt (Stepanov et al. 2012) and with decreasing phosphorus

309 concentrations in the melt (Duc-Tin and Keppler 2015). Studies that couple experimentally-
310 determined solubility equations of monazite with phase equilibria modelling show that monazite
311 dissolution increases during prograde metamorphism above the solidus and that for UHT
312 metamorphism most monazite will be completely consumed along the prograde path (Kelsey et
313 al. 2008; Yakymchuk and Brown 2014b). Based on these theoretical models of monazite
314 behaviour, no prograde monazite growth is expected above the solidus. This contrasts with
315 studies of natural rocks that record prograde suprasolidus monazite growth (e.g., Hermann and
316 Rubatto 2003; Hacker et al. 2015). Johnson et al. (2015) suggest that apatite dissolution during
317 prograde metamorphism may have contributed to LREE saturation of the melt and resulted in
318 prograde suprasolidus monazite crystallization. However, the role of apatite breakdown and
319 monazite growth in suprasolidus metamorphic rocks is still poorly understood.

320 Finally, similar to zircon, Ostwald ripening could produce prograde monazite growth in
321 suprasolidus metamorphic rocks (e.g., Nemchin and Bodorkos 2000). However, this has not been
322 extensively studied and may only apply to monazite growth just above the solidus where the
323 modal proportion of anatectic melt is relatively low.

324

325 **Xenotime**

326 Xenotime is a common phosphate in low- to high-grade metamorphic rocks (e.g., Franz
327 et al. 1996; Bea and Montero 1999; Pyle and Spear 1999; Spear and Pyle 2002). It can be
328 directly dated using U–Pb geochronology (e.g., Parrish 1990; Rasmussen et al. 2005, 2007,
329 Sheppard et al. 2007; Janots et al. 2009; Crowley et al. 2009) and can be used as a thermometer
330 when in equilibrium with monazite and garnet (Pyle et al. 2001). Some of the proposed
331 mechanisms of metamorphic xenotime growth are: (1) growth due to dissolution/reprecipitation

332 of detrital or diagenetic zircon (Dawson et al. 2003; Rasmussen et al. 2011), (2) growth during to
333 the breakdown of detrital zircon (Franz et al. 2015), (3) growth during the breakdown of allanite
334 and/or monazite (Janots et al. 2008), (4) crystallization from anatectic melt (Pyle and Spear
335 1999; Crowley et al. 2009), and (5) growth during the breakdown of Y-rich garnet (e.g., Pyle and
336 Spear 1999).

337 There are three important controls on the stability of xenotime in metamorphic rocks.
338 First, higher bulk rock Yttrium concentrations allow xenotime to persist to higher pressures and
339 temperatures (Spear and Pyle 2010). Second, similar to monazite and zircon, partial melting will
340 result in xenotime consumption in order to contribute to REE and P saturation of the melt (e.g.,
341 Wolf and London 1995; Duc-Tin and Keppler 2015). Third, xenotime will be consumed during
342 the growth of garnet, which partitions Y and HREE (Pyle and Spear 1999, 2000; Spear and Pyle
343 2002). Consequently, xenotime in the matrix of high-grade garnet-bearing rocks is relatively rare
344 although xenotime inclusions may be present in garnet (e.g., Pyle and Spear 1999). By contrast,
345 xenotime may be present in the matrix at various metamorphic grades in garnet-absent rocks.

346

347

PHASE EQUILIBRIA MODELLING

348 Phase assemblages in metamorphic rocks generally change through continuous or
349 multivariant reactions rather than discontinuous or univariant reactions (e.g., Stüwe and Powell
350 1995; Kelsey and Hand 2015). Petrogenetic grids display discontinuous reactions and may yield
351 the false impression that reactions occur over narrow intervals along a P - T path (e.g., Vernon
352 1996). In reality, mineral (and melt) modes and compositions are continuously changing along a
353 P - T evolution and these variations represent the reaction sequence of the rock. The particular
354 reaction sequence experienced by a metamorphic rock is dependent on bulk composition, the P -

355 *T* path and whether the system is open or closed (e.g., White and Powell 2002; Brown and
356 Korhonen 2009; Yakymchuk and Brown 2014a). The phase assemblages expected for a
357 particular bulk rock composition over a range of *P–T* conditions are commonly depicted using
358 *P–T* pseudosections, phase diagrams constructed for a fixed bulk composition (e.g., Hensen
359 1971; Worley et al. 1998; Spear et al. 2016). More detailed information about the reaction
360 sequences can be determined by using the intersection of a *P–T* path with isopleths (or contours)
361 of different variables (e.g., mineral modes and compositions) on a pseudosection or by
362 constructing mode–temperature or mode–pressure diagrams (e.g., White et al. 2011).

363 Here, we present pseudosections and mode-box diagrams (*P/T*–mode plots) for different
364 bulk chemical compositions to investigate the reaction sequences along common *P–T* paths and
365 we discuss the consequences for interpreting the ages obtained from accessory mineral
366 geochronology. The effects of fractionation of growing porphyroblasts and changes in bulk
367 composition are discussed later. Because we use phase equilibria modelling, it is assumed that
368 there are no kinetic controls (e.g., nucleation barriers or sluggish diffusion) that impact the
369 growth or dissolution of major and accessory minerals. Although this assumption may not be
370 strictly valid in some circumstances (Watt and Harley 1993; Pattison and Tinkham 2009; Gaides
371 et al. 2011; Pattison and Debuhr 2015), the modelling here provides a general framework for
372 investigating reaction sequences.

373

374 **Bulk compositions**

375 Three bulk chemical compositions are modelled that represent the most common
376 protoliths investigated during most metamorphic studies. For the subsolidus and suprasolidus *P–*
377 *T* paths, this includes an average amphibolite-facies metapelite (Ague 1991) and an average

378 passive margin greywacke (Yakymchuk and Brown 2014a). These two compositions were
379 chosen because they are expected to dominate passive margin turbidite sequences that are
380 involved in orogenesis at convergent plate boundaries. A mid-ocean ridge basalt (MORB) from
381 Sun and McDonough (1989) was also used to investigate a suprasolidus reaction sequence for a
382 typical mafic protolith. The modelled bulk compositions are summarized in Table 1. Note that
383 we do not consider melt loss nor fractionation of minerals away from the reacting volume in the
384 modelling here, both of which can modify the composition of the equilibrium volume during a
385 metamorphic evolution (e.g., White and Powell 2002; Evans 2004; Guevara and Caddick 2016;
386 Mayne et al. 2016).

387

388 **Computational Methods**

389 Forward phase equilibria modelling is used to evaluate the changes to metamorphic
390 mineral assemblages along several schematic P – T paths. There are two main approaches to phase
391 equilibria modelling. The first approach is to use Gibbs free energy minimization to determine
392 the stable mineral assemblage at a given P – T condition and there are two commonly used
393 software packages available for this approach, including `Perple_X` (Connolly and Petrinì 2002)
394 and `Theriak–Domino` (de Capitani and Brown 1987; de Capitani and Petrakakis 2010). A second
395 approach is to determine the solution of simultaneous non-linear equations to build up an array
396 of points and lines that make up a metamorphic phase diagram (`THERMOCALC`: Powell and
397 Holland 1988; Powell et al. 1998; Powell and Holland 2008). Both of these approaches require
398 an internally consistent thermodynamic database derived from the results of experimental
399 studies. There are several commonly-used databases available: Berman (1988), Holland and
400 Powell (1998) and most recently Holland and Powell (2011). Finally, the calculations require

401 activity–composition models that relate end-member proportions to end-member activities for
402 the solid-solution minerals as well as for complex fluids and melt.

403 Here, calculations were performed using THERMOCALC v.3.40 (Powell and Holland 1988)
404 using the internally consistent dataset of Holland and Powell (2011). For metapelite and
405 greywacke compositions, modelling was undertaken in the MnO–Na₂O–CaO–K₂O–FeO–MgO–
406 Al₂O₃–SiO₂–H₂O–TiO₂–Fe₂O₃ (MnNCKFMASHTO) chemical system using the activity–
407 composition relations in White et al. (2014a,b). An average MORB composition was
408 investigated in the NCKFMASHTO chemical system using the activity-composition models
409 from Green et al. (2016). Phases modelled as pure end-members are quartz, rutile, titanite,
410 aqueous fluid (H₂O), kyanite and sillimanite. Mineral abbreviations are from Holland and Powell
411 (2011) with the exception of titanite (ttn).

412 Phase equilibria modelling of subsolidus and suprasolidus systems requires different
413 approaches to approximate the bulk H₂O content of the system. For subsolidus phase equilibria
414 modelling, the amount of H₂O in each bulk composition was set to be in excess, such that $a_{\text{H}_2\text{O}}$
415 =1. By contrast, suprasolidus rocks are generally not considered to have excess H₂O as the very
416 small amount of free H₂O at the solidus is partitioned into anatectic melt (Huang and Wyllie
417 1973; Thompson 1982; Clemens and Vielzeuf 1987; White and Powell 2002; White et al. 2005).
418 Along the solidus, the solubility of H₂O in anatectic melt increases with pressure. Therefore, for
419 phase equilibria modelling of suprasolidus rocks, the amount of H₂O in the bulk composition
420 must be set, and was adjusted so that the melt is just saturated with H₂O at solidus at 8 kbar. If
421 the modelled prograde path crossed the solidus at lower or higher pressures, the quantity of melt
422 produced will be slightly overestimated and underestimated, respectively.

423

424 **Modelled P – T paths**

425 Several simplified typical P – T paths for subsolidus and suprasolidus systems are
426 modelled to investigate the reaction sequences, changes in mineral modes, amount of melt
427 generated and/or consumed, variations in bulk rock a_{TiO_2} and the consequences for interpreting
428 the results of accessory mineral geochronology. Two subsolidus and two suprasolidus P – T paths
429 were investigated for the metapelite and greywacke compositions that represent different tectonic
430 evolutions. We also model one suprasolidus P – T path for a MORB composition applicable to
431 high-pressure mafic granulites.

432 For the subsolidus systems, the first P – T path has a clockwise trajectory and is
433 representative of collisional orogenesis (e.g., England and Thompson 1984; Thompson and
434 England 1984). An important feature of this P – T path is that peak P occurs before peak T , which
435 is consistent with a relatively long residence time in the core of an orogenic belt. The model path
436 ('clockwise' in Figs 3, 5) contains four segments: (1) heating and burial up to 10 kbar and
437 550°C, (2) isobaric heating up to 650°C, (3) decompression and minor heating to 8 kbar and
438 660°C, and (4) decompression and cooling to 6.2 kbar and 610°C.

439 The second subsolidus P – T path has a 'hairpin' trajectory where peak P and peak T
440 nearly coincide (e.g., 'hairpin' in Figs 3, 5). This style of P – T path is also common in collisional
441 orogenesis (e.g., Brown 1998; Kohn 2008) and reflects relatively short residence times at depth
442 and can represent crustal thickening in an accretionary zone of a propagating orogen (Jamieson
443 et al. 2004). The model hairpin P – T path contains three segments: (1) heating and burial up to 7
444 kbar and 610°C, (2) heating and decompression to 6 kbar and 650°C, and (3) decompression and
445 cooling to 4 kbar and 600°C. Peak metamorphism is at the same temperature for both subsolidus
446 P – T paths, but the pressures at peak T are different.

447 Two P – T paths for suprasolidus rocks that are typical of granulite-facies metamorphism
448 are investigated. First, a clockwise P – T path is modelled (e.g., ‘clockwise’ in Figs 7, 9) that is
449 associated with crustal thickening and heating and is typical of many granulite terranes (e.g.,
450 Clark et al. 2011). This P – T path contains four segments: (1) increase in pressure and heating
451 across the solidus up to 9 kbar at 850°C, (2) isobaric heating up to 900°C, (3) isothermal
452 decompression to 7 kbar, and (4) cooling and decompression to 5 kbar at 750°C.

453 Second, a suprasolidus counterclockwise (or anticlockwise) P – T path is considered (e.g.,
454 ‘counterclockwise’ in Figs 7, 9) and is common in high temperature–low pressure terranes (e.g.,
455 Clarke et al. 1987; Collins and Vernon 1991) and some ultrahigh temperature terrains (e.g.,
456 Korhonen et al. 2013a, 2014). This P – T path reflects heating prior to thickening (usually at peak
457 T) followed by near-isobaric cooling. In some cases, this has been attributed to the inversion of a
458 hot back-arc basin during collisional orogenesis (Clark et al. 2014). The modelled
459 counterclockwise P – T path contains four segments: (1) isobaric heating up to 830°C and 5 kbar,
460 (2) heating and an increase in pressure up to 860°C and 6 kbar, (3) cooling and an increase in
461 pressure to 850°C and 7 kbar, and (4) isobaric cooling to 750°C.

462 A high-pressure clockwise path was also chosen to model the reaction sequences for
463 high-pressure mafic granulites (e.g., O’Brien and Rötzler 2003) and to investigate the growth and
464 breakdown of garnet for these rocks. This P – T path contains four segments: (1) heating and an
465 increase in pressure up to 12 kbar and 850°C, (2) isobaric heating up to 950°C, (3) isothermal
466 decompression to 8 kbar, and (4) cooling and decompression to 800°C and 6 kbar. The
467 decompression segments of this P – T path are also compatible with the decompression segments
468 of some ultrahigh-pressure metamorphic rocks (Hacker et al. 2010; Chen et al 2013; Xu et al
469 2013).

470

471 **Modelling suprasolidus zircon and monazite dissolution**

472 For the suprasolidus P – T paths, the growth and dissolution of zircon and monazite are
473 modelled using the method of Kelsey et al. (2008). Note that the bulk composition used in this
474 modelling is fixed without fractionation of elements into growing porphyroblasts (e.g., zirconium
475 into garnet) or loss of melt. These factors are discussed later. First, the saturation concentrations
476 of the melt in ppm are calculated as follows. The major element composition of the anatectic
477 melt is calculated at a specified P – T . This composition of the melt is combined with the
478 solubility equations of Boehnke et al. (2013) for zircon and Stepanov et al. (2012) for monazite,
479 and the stoichiometric concentrations of Zr in zircon (497,664 ppm Zr) and LREE in monazite
480 (566,794 ppm LREE) to determine the saturation concentrations of Zr and LREE in ppm (Kelsey
481 et al. 2008). These initial calculations do not account for the proportion of anatectic melt or the
482 bulk-rock concentrations of Zr and LREE. Second, the saturation concentrations of Zr or LREE
483 (ppm) are multiplied by the proportion of anatectic melt in the system (in normalized mol%
484 which is approximately equivalent to vol%) at P – T to arrive at concentrations in ppm that are
485 required to saturate the melt in the equilibration volume of the rock. Finally, these values are
486 divided by the assumed bulk-rock chemical concentrations of Zr and LREE. The Zr and LREE in
487 pelites and the greywackes are generally similar to each other and to the values of
488 metasedimentary migmatites; a value of 150 ppm was chosen here (e.g., Yakymchuk and Brown
489 2014b). For MORB, the Zr concentration was set at 103 ppm, which is an average of global
490 MORB compositions (White and Klein 2014). The sensitivity of these calculations to bulk rock
491 concentrations of Zr and LREE and the assumptions and limitations of this methodology are
492 discussed by Kelsey et al. (2008) and Yakymchuk and Brown (2014b). The result of the

493 calculations is the proportion of zircon or monazite dissolution required to saturate the anatectic
494 melt in Zr and LREE in the equilibration volume of the rock. This value is subtracted from 100%
495 and the results are reported as the percent of zircon and monazite remaining relative to the
496 amounts existing at the fluid-present solidus for each P - T path.

497 One important limitation of the monazite saturation equation used here is that it does not
498 account for phosphorus in anatectic melt, which can affect monazite stability (e.g., Duc-Tin and
499 Keppler 2015). The main repository of phosphorus in most metamorphic rocks is apatite, which
500 is expected to break down during partial melting (Wolf and London 1994). Phosphorus
501 saturation in melt is a function of temperature, SiO_2 concentration (Harrison and Watson 1984)
502 and the aluminum saturation index of the melt ($\text{ASI} = \text{molar } [\text{Al}_2\text{O}_3] / [\text{K}_2\text{O} + \text{Na}_2\text{O} + \text{CaO}]$,
503 Wolf and London 1994). The solubility of apatite in melt increases with rising ASI, which is
504 common during prograde partial melting in migmatites (e.g., Johnson et al. 2015). For example,
505 an increase in the ASI from 1.1 to 1.2 increases the solubility of apatite by an order of magnitude
506 (Wolf and London 1994). The breakdown of LREE-rich apatite during prograde metamorphism
507 may contribute to LREE saturation of anatectic melt and monazite crystallization (e.g., Johnson
508 et al. 2015; Rocha et al. 2016). Here, we use melt ASI as a qualitative tool to investigate the
509 behaviour of apatite during partial melting and the consequences for prograde monazite growth
510 in suprasolidus metamorphic rocks.

511

512 **SUBSOLIDUS PHASE RELATIONS AND REACTION SEQUENCES**

513 Phase relations and reaction sequences for two subsolidus P - T paths for two bulk compositions
514 are discussed below for a closed system where bulk composition does not change. Fractionation
515 of porphyroblasts away from the reacting volume is not considered in the calculations here.

516

517 **Metapelite**

518 *Phase relations.* The pseudosection for the amphibolite-facies metapelite is shown in
519 Figure 3. Biotite is stable at $T > 450\text{--}460^\circ\text{C}$ with increasing pressure. Garnet is stable across
520 most of the diagram except at low P and low T and at $P < 5$ kbar just below the solidus. The
521 staurolite-in field boundary extends from 550°C at 4 kbar up to 650°C at 10 kbar. The stability
522 field of the aluminosilicate minerals (kyanite and sillimanite) ranges from $T > 580^\circ\text{C}$ at 4 kbar to
523 $T > 660^\circ\text{C}$ at 10 kbar. Rutile is stable at low P at low temperatures and at $P > 10.5$ at $T > 550^\circ\text{C}$.

524

525 *Reaction sequence for the clockwise $P\text{--}T$ path.* A mode-box for the reaction sequence
526 for the clockwise $P\text{--}T$ path is shown in Figure 4a. During the prograde segment of the clockwise
527 $P\text{--}T$ path, chlorite is progressively consumed to produce biotite (starting at 455°C) and then
528 garnet (starting at 460°C). Chlorite is exhausted by 580°C . Rutile is completely consumed by
529 $\sim 530^\circ\text{C}$. Epidote breaks down along the prograde path and the epidote-out field boundary is
530 encountered at 620°C . The complete breakdown of epidote at 620°C may result in minor zircon
531 crystallization or more likely the growth of new rims on pre-existing zircon.

532 The staurolite stability field is encountered during the heating and decompression
533 segment at 9 kbar and 650°C . In natural examples, prograde monazite growth has been
534 associated with garnet, staurolite and kyanite/sillimanite growth in metapelites (Catlos et al.
535 2001; Smith and Barriero 1990; Wing et al. 2003; Kohn and Malloy 2004; Fitzsimons et al.
536 2005; Corrie and Kohn 2008), although kyanite/sillimanite is not encountered along the $P\text{--}T$ path
537 modelled here. In general, monazite could grow at different points along the prograde $P\text{--}T$ path.
538 Because garnet growth is predicted to be continuous up to the staurolite-in field boundary, new

539 monazite grown during this segment of the P – T path is expected to have low concentrations of
540 HREE and Y; these elements are expected to partition into garnet. Staurolite growth is at the
541 expense of garnet and occurs during the decompression portion of the prograde path (e.g.,
542 Florence and Spear 1993). Monazite formed during this segment is expected to have elevated
543 HREE and Y concentrations due to the breakdown of garnet and the age would record the initial
544 stages of decompression from peak P . By contrast, zircon is expected to undergo only minor
545 changes to its mode and will be slightly consumed during subsolidus prograde metamorphism
546 (e.g., Kohn et al. 2015). Two possible exceptions are minor zircon growth during the breakdown
547 of Zr-rich epidote and/or rutile. However, at low temperatures, rutile is not expected to be
548 zirconium rich.

549 The stable mineral assemblage at the metamorphic peak of 660°C and 8 kbar includes
550 garnet, staurolite, biotite, plagioclase, ilmenite, muscovite and quartz. In general, the mode of
551 monazite in metapelites is expected to reach a maximum at peak P – T and monazite is not
552 expected to grow during cooling (e.g., Spear and Pyle 2010). An additional factor is that
553 monazite is susceptible to dissolution/reprecipitation in the presence of fluids (e.g., Williams et
554 al. 2011). While fluid-mediated dissolution/reprecipitation should not increase the net mode of
555 monazite, it can occur at almost any point along the P – T evolution. Therefore, it is possible that
556 monazite in a subsolidus metapelite can record a range of ages that vary from monazite-in up to
557 the metamorphic peak and dissolution/precipitation may result in ages that record retrogression
558 (Harlov et al. 2005; Williams et al. 2011). On the other hand, zircon is expected to be relatively
559 unreactive and may not record prograde to peak metamorphism in subsolidus metamorphic
560 rocks.

561 After the exhaustion of rutile during the heating and burial segment the value of a_{TiO_2}
562 decreases to 0.95 initially and remains nearly constant until the decompression and heating
563 segment, where a_{TiO_2} drops to 0.8. During the subsequent decompression and cooling segment
564 the a_{TiO_2} value remains stable at 0.8. After rutile is exhausted, any growth of new zircon or
565 modification of existing zircon will occur when $a_{\text{TiO}_2} < 1.0$; this needs to be considered when
566 applying the Ti-in-zircon thermometer.

567

568 ***Reaction sequence for the hairpin P–T path.*** A mode-box for the reaction sequence for
569 the hairpin P – T path is shown in Figure 4b. The prograde reaction sequence includes garnet-in at
570 520°C and 4.5 kbar, staurolite-in at 590°C and 6.5 kbar and sillimanite-in at 640°C and 6 kbar.
571 Garnet both grows and is consumed during the prograde portion of the reaction sequence. The
572 value of a_{TiO_2} decreases from 0.88 at 500°C to 0.79 at peak T . For the decompression and cooling
573 segment, a small amount of biotite grows at the expense of garnet; the modes of the other
574 minerals are little changed and the a_{TiO_2} value remains close to 0.79.

575 As with the clockwise P – T path, monazite growth may occur at various times during the
576 prograde segment of the hairpin P – T path. Because the mode of garnet increases and decreases
577 multiple times during the prograde reaction sequence, linking Y and HREE in monazite would
578 be challenging in the absence of microstructural context, such as monazite in coronae
579 surrounding resorbed garnet. For example, HREE and Y-rich monazite could be generated
580 during: (1) the heating and burial segment during garnet breakdown to staurolite, (2) the heating
581 and decompression segment during garnet breakdown to biotite near the metamorphic peak, or
582 (3) garnet breakdown during the retrograde segment of the P – T path. Furthermore, xenotime
583 breakdown during prograde metamorphism would also contribute to the growth of Y and HREE-

584 rich monazite with or without a contribution from garnet (e.g., Spear and Pyle 2010). Therefore,
585 for the hairpin P – T path modelled here, HREE- or Y-rich monazite can record burial and/or
586 exhumation.

587

588 **Greywacke**

589 ***Phase relations.*** The pseudosection for the greywacke is shown in Figure 5. Garnet is
590 stable across the entire diagram. Titanite is stable at $T < 460$ – 470°C with increasing pressure.
591 Biotite is stable across the diagram except at the high- P –low- T corner. Chlorite is unstable above
592 600°C at 7 kbar and unstable above 530°C at 12 kbar. Albite is stable below 565°C at 4 kbar and
593 below 620°C at 11 kbar. White mica (muscovite and/or paragonite) is stable up to the solidus at
594 $P > 10$ kbar and at $T < 570^\circ\text{C}$ at 4 kbar. There are two important sets of quasi-linear field
595 boundaries with positive slopes in P – T space. First is the set of boundaries that range from
596 480°C at 4 kbar up to 640°C at 12 kbar represent the breakdown of epidote to produce
597 plagioclase. A second set of fields extends from 570°C at 4 kbar to the solidus with increasing P
598 that represent the consumption of paragonite to produce sillimanite or kyanite.

599

600 ***Reaction sequence for the clockwise P – T path.*** A mode-box for the assemblage
601 sequence for the clockwise P – T path is shown in Figure 6a. For the clockwise P – T path the
602 important changes along the prograde reaction sequence include: (1) the growth of rutile
603 commencing at 460°C resulting in a_{TiO_2} increasing from 0.9 at 450°C to 1.0 at 460°C , (2) the
604 complete consumption of titanite by 475°C , (3) the growth of biotite and garnet at the expense of
605 chlorite and muscovite from 450°C to 550°C , which results in the complete consumption of
606 chlorite by 550°C , (4) the breakdown of epidote to produce 2 mol% plagioclase at $\sim 605^\circ\text{C}$, (5)

607 the consumption of ~0.5 mol% garnet to produce plagioclase and hematite from 605°C to 620°C,
608 and (6) the growth of 0.5 mol% garnet from 620°C to 650°C. Similar to the metapelite, the
609 breakdown of Zr-rich epidote may result in minor zircon growth.

610 Decompression and heating from 10 kbar at 650°C to 8 kbar at 660°C results in: (1) the
611 disappearance of rutile at 9.8 kbar and a subsequent drop in a_{TiO_2} , and (2) the breakdown of
612 paragonite and garnet to produce kyanite, biotite and plagioclase at 8.8 kbar. The peak
613 metamorphic assemblage at 660°C and 8 kbar includes garnet, kyanite, muscovite, biotite,
614 plagioclase and ilmenite. Further decompression and cooling results in the growth of biotite at
615 the expense of garnet and a gradual decrease in a_{TiO_2} .

616 Like the metapelite, garnet breakdown occurs during a portion of the heating and burial
617 segment of the P - T path (at ~600°C) and during the heating and decompression segment
618 immediately before the metamorphic peak. Therefore, Y- and HREE-rich monazite associated
619 with garnet breakdown could record the timing of burial and/or exhumation.

620

621 *Reaction sequence for the hairpin P - T path.* A mode-box for the reaction sequence for
622 the hairpin P - T path is shown in Figure 6b. The hairpin P - T path has three notable differences in
623 its reaction sequence compared with the clockwise path for the greywacke. First, the P - T path
624 does not intersect the epidote stability field. Second, the value of a_{TiO_2} is always less than one
625 and decreases from 0.95 to 0.80 along the P - T path. And third, white mica is completely
626 consumed by the end of the P - T path. In contrast to the multiple garnet growth/consumption
627 segments along the clockwise P - T path, the mode of garnet increases during burial and decreases
628 during decompression for the hairpin path. Therefore, Y- and HREE-depleted monazite is

629 predicted to record burial and monazite enriched in these elements would document garnet-
630 breakdown during decompression and/or cooling and may also record xenotime breakdown.

631

632 SUPRASOLIDUS PHASE RELATIONS AND REACTION SEQUENCES

633 Phase relations and reaction sequences for three suprasolidus scenarios are discussed below for a
634 closed system where bulk composition does not change and the water content is fixed to just
635 saturate the rock in H₂O at 8 kbar at the solidus. This approach provides important first-order
636 constraints on the reaction sequences for the modelled compositions; it does not take into
637 account melt loss and the associated effects on rock fertility, solidus temperature and zircon and
638 monazite stability. The consequences of open system behaviour on the reaction sequence and
639 accessory mineral stability are discussed later.

640

641 Metapelite

642 *Phase Relations.* The pseudosection for the metapelite is shown in Figure 7. Ilmenite and
643 plagioclase are stable across the entire diagram. Garnet is stable at high pressures across the
644 diagram except at $T < 700^{\circ}\text{C}$ at $P < 5.5$ kbar. Cordierite is stable at high- T -low- P conditions.
645 Orthopyroxene is not stable in this diagram. Rutile is restricted to $P > 10$ – 11.5 kbar across the
646 modelled temperature range. The three important partial melting reactions for the metapelite
647 include: (1) the consumption of any free aqueous fluid to produce melt at the wet solidus, which
648 ranges from 660°C at low pressures to 710°C at high pressure, (2) the incongruent breakdown of
649 muscovite to produce K-feldspar, which is represented by a narrow low-variance field that
650 extends from the wet solidus at low P to $T > 780^{\circ}\text{C}$ at $P > 12$ kbar, and (3) the progressive
651 breakdown of biotite at temperatures above the muscovite stability field to produce either garnet

652 at higher pressure or cordierite ± garnet at lower pressure. Biotite is exhausted by 850°C at $P > 7$
653 kbar and melting progresses via the consumption of quartz and feldspar.

654

655 ***Reaction sequence for the clockwise P – T path.*** The mode-box and titania activity for the
656 clockwise assemblage sequence are shown in Figure 8a. The predicted amount of monazite and
657 zircon remaining as well as melt ASI are shown in Figure 8b. During the prograde segment of
658 the clockwise P – T path, the metapelite begins to melt at the wet solidus at ~680°C through the
659 consumption of any free H₂O as well as quartz and plagioclase. The amount of aqueous fluid at
660 the solidus is expected to be small given the limited porosity of high-grade metamorphic rocks
661 (e.g., Yardley and Valley 1997). Therefore, the amount of melt produced at the wet solidus is
662 expected to be limited. For the modelled metapelite, the amount of melt produced is 4 mol%,
663 although this is mostly an artefact of the modelling which assumes H₂O saturation of the solidus
664 at 8 kbar.

665 After any free H₂O is consumed, partial melting continues with increasing temperature
666 through the consumption of muscovite up until 728°C. This produces an additional 3 mol% melt
667 with a generally constant melt ASI. At 728°C and 7.5 kbar, melting proceeds via the incongruent
668 breakdown of muscovite to generate K-feldspar over a narrow (~2°C) temperature range. This
669 narrow field produces an additional 3 mol% melt. After muscovite is exhausted, the rock
670 contains ~9 mol% melt and approximately 8% of the zircon and 12% of the monazite that was
671 present at the solidus is predicted to have been consumed. A minor amount of apatite breakdown
672 is expected in order to saturate the anatectic melt in phosphorus.

673 After muscovite is completely consumed, partial melting continues through the
674 consumption of biotite, plagioclase and quartz to produce melt, garnet, K-feldspar and ilmenite.

675 This produces an additional 32 mol% melt. Because biotite is an important host of accessory
676 mineral inclusions (e.g., Watson et al. 1989), biotite breakdown may release inclusions of
677 monazite and zircon into the reaction volume of the rock. The liberation of these accessory
678 minerals may contribute to LREE and Zr saturation of the anatectic melt. On the other hand,
679 some zircon and monazite may be included in growing garnet and will be sequestered away from
680 the reaction volume. These minerals will be shielded from dissolution and are more likely to
681 preserve inherited (or detrital) ages as well as any subsolidus to early suprasolidus prograde
682 metamorphic ages. The inclusion of accessory minerals in major minerals effectively reduces the
683 Zr and LREE available to the system and proportionally more zircon and monazite dissolution
684 will be required to maintain melt saturation in these elements (e.g., Yakymchuk and Brown
685 2014b). Therefore, the sequestration of zircon and monazite in stable peritectic minerals such as
686 garnet will promote the dissolution of accessory minerals along grain boundaries in the matrix of
687 the rock with increasing T .

688 Titania activity is predicted to reach its highest value of 0.92 at 850°C. At this point, all
689 of the monazite and 85% of the zircon are expected to be consumed. However, during this
690 interval, the ASI of the melt increases from 1.10 to 1.15, which may result in enhanced apatite
691 dissolution. For example, Pichavant et al. (1992) estimate that a similar increase in ASI at 800°C
692 and 5 kbar would change the P_2O_5 concentration of melt from 0.50 to 0.75 wt%. If apatite is
693 LREE rich, the dissolution of apatite may delay the complete dissolution of monazite to higher
694 temperatures or, in extreme cases, may even promote prograde monazite crystallization (e.g.,
695 Johnson et al. 2015).

696 After the complete consumption of biotite, the hydrous minerals have been exhausted and
697 the residuum is essentially composed of anhydrous minerals. Melting continues through the

698 continued consumption of quartz and K-feldspar up to the modelled peak temperature of 900°C.
699 The ASI of the melt reaches 1.18 at peak T and significant apatite dissolution is likely. At this
700 point, the metapelite contains 43 mol% melt and both monazite and zircon are absent. After
701 zircon and monazite are completely consumed, further anatexis is expected to generate melt that
702 is undersaturated in Zr and LREE. Therefore, even with the breakdown of LREE-rich apatite,
703 prograde monazite crystallization is not expected above ~850°C.

704 Isothermal decompression from 9 to 7 kbar produces an additional 11 mol% melt at the
705 expense of quartz and K-feldspar. At this point the rock contains the maximum amount of melt
706 of 54 mol%. During this decompression segment, melt ASI increases from 1.18 to 1.23 and a_{TiO_2}
707 decreases from 0.86 to 0.74. Approximately 1 mol% garnet is consumed during decompression,
708 which would liberate some HREE and Y into the reaction volume. If significant apatite
709 dissolution occurs and promotes monazite crystallization during decompression, this may be
710 reflected as elevated HREE concentrations in monazite. However, the melt is undersaturated in
711 LREE given the complete exhaustion of monazite at 850°C so new monazite growth is unlikely
712 unless the apatite is very enriched in LREE or the modal proportion of apatite is high.

713 Decompression and cooling from peak T results in: (1) melt crystallization, (2) the
714 growth of K-feldspar and quartz until the cordierite stability field is reached at ~880°C and 6.7
715 kbar, (3) garnet and cordierite consumption to produce biotite, (4) new zircon and monazite
716 growth, (5) a decrease in melt ASI and a_{TiO_2} . Melt crystallization and cordierite growth are
717 concomitant with the consumption of garnet and sillimanite; this is a common reaction sequence
718 for high-temperature decompression in migmatites. Zircon or monazite produced over this
719 reaction interval is expected to be enriched in HREE and Y (e.g., Yakymchuk et al. 2015).
720 Protracted monazite and zircon growth will occur from peak T to the solidus. U–Pb zircon and

721 monazite ages that spread down Concordia have been interpreted to record protracted growth
722 during cooling and melt crystallization in various migmatite terranes with clockwise P - T
723 evolutions (Korhonen et al. 2012; Reno et al. 2012; Morrisey et al. 2014; Walsh et al. 2015).

724

725 ***Reaction sequence for the counterclockwise P - T path.*** The mode-box and titania
726 activity for the counterclockwise assemblage sequence are shown in Figure 8c. The amount of
727 monazite and zircon remaining as well as melt ASI are shown in Figure 8d. During isobaric
728 heating at 5 kbar, partial melting begins at the wet solidus at $\sim 670^\circ\text{C}$ and continues through the
729 breakdown of muscovite to produce a total 8 mol% melt by 685°C . Melting continues via the
730 progressive breakdown of biotite to produce peritectic garnet and K-feldspar. By 765°C , 17
731 mol% melt is present and roughly 80% and 60% of the initial amount of zircon and monazite,
732 respectively, remains. From the solidus up to 765°C , the a_{TiO_2} value of the system has decreased
733 from 0.9 to 0.6 and the ASI of the melt has increased from 1.13 to 1.18. An increase in melt ASI
734 increases the solubility of apatite; this may liberate some LREE and P that could contribute to
735 minor monazite crystallization because the melt is predicted to be saturated with respect to the
736 LREE. Similar to the clockwise P - T path, the breakdown of biotite may liberate zircon and/or
737 monazite that was sequestered away from the reacting volume.

738 Cordierite enters the phase assemblage at $\sim 770^\circ\text{C}$ and melting continues through the
739 breakdown of biotite, sillimanite and plagioclase to produce K-feldspar, garnet and cordierite.
740 Sillimanite is completely consumed by 780°C and biotite is exhausted by 800°C . During the
741 interval from 770°C to 800°C , approximately 15 mol% melt is generated and melt ASI decreases
742 slightly from 1.18 to 1.16. Although the change in ASI would decrease the solubility of apatite,

743 this may be counteracted by the additional melt generation during biotite breakdown and apatite
744 growth is not expected.

745 In the absence of biotite, melting continues through the breakdown of quartz, plagioclase
746 and K-feldspar, which results in a progressively drier melt. Monazite and zircon are predicted to
747 be completely consumed by 820°C and 850°C, respectively. The increase in pressure near the
748 metamorphic peak results in minor cordierite consumption to produce garnet. At the
749 metamorphic peak, the metapelite has generated ~44 mol% melt.

750 Isobaric cooling at 6.5 kbar from 860°C to 750°C results in: (1) melt crystallization; (2)
751 new zircon and monazite growth commencing at 855°C and 825°C, respectively; (3) a decrease
752 in melt ASI, which would decrease the solubility of apatite and contribute to apatite
753 crystallization; (4) the retrogression of cordierite and garnet to biotite; (5) the consumption of K-
754 feldspar and the growth of plagioclase; and (6) a decrease in a_{TiO_2} . The concurrent breakdown of
755 garnet and growth of zircon, monazite and plagioclase may result in elevated HREE and Y
756 concentrations and more pronounced negative Eu anomalies in newly crystallized accessory
757 minerals. New zircon growth occurs when the a_{TiO_2} value of the system ranges from 0.6 to 0.7.
758 Applying the Ti-in-zircon thermometer assuming an a_{TiO_2} value of 1.0 would underestimate the
759 true temperature by ~50°C (Fig. 1b). Protracted retrograde monazite and zircon growth is
760 expected during melt crystallization along the isobaric cooling segment of the counterclockwise
761 P - T path (e.g., Korhonen et al. 2013b)

762

763 **Greywacke**

764 ***Phase relations.*** The pseudosection for the greywacke is shown in Figure 9. When
765 compared with the metapelite, the greywacke composition is less fertile (e.g., Clemens and

766 Vielzeuf 1987; Thompson 1996; Vielzeuf and Schmidt 2001; Johnson et al. 2008; Yakymchuk
767 and Brown 2014a), contains different mineral assemblages, and yields different reaction
768 sequences. For the greywacke, quartz, ilmenite, plagioclase and garnet are stable across the
769 entire diagram. Cordierite becomes stable at pressures of 5–7 kbar with increasing temperature.
770 Orthopyroxene is stable at $P < 6$ kbar and $T > 800^{\circ}\text{C}$. Rutile is stable at $P > 9$ –11 kbar.
771 Muscovite is stable at $P > 7$ –12 kbar with increasing temperature. Similar to the metapelite,
772 partial melting of the greywacke begins at the wet solidus at temperatures of ~ 670 – 700°C . A
773 narrow low-variance field representing the breakdown of muscovite to produce K-feldspar is
774 restricted to $P > 10.8$ kbar; this contrasts with a similar field in the metapelite pseudosection that
775 extends from < 4 to > 12 kbar (Fig. 7). For the greywacke composition, biotite breakdown
776 generates garnet at high pressure, cordierite at low pressure, and orthopyroxene at low pressure
777 and high temperature.

778

779 ***Reaction sequence for the clockwise P – T path.*** The mode-box and titania activity for the
780 clockwise assemblage sequence are shown in Figure 10a. The predicted amount of monazite and
781 zircon remaining along with the ASI of melt are shown in Figure 10b. Along the clockwise P – T
782 path, partial melting begins at $\sim 675^{\circ}\text{C}$ at the wet solidus and muscovite is not stable. Therefore,
783 further melting proceeds through the breakdown of biotite and sillimanite to produce garnet.
784 Similar to the metapelite, growing garnet has the potential to capture inherited or prograde
785 monazite and zircon allowing their preservation to higher temperatures. K-feldspar becomes
786 stable at 800°C and 8.2 kbar and biotite is completely consumed by 850°C . Up to this point, the
787 greywacke has produced 14 mol% melt, which is less than the 34 mol% melt generated by the
788 pelite for the same P – T path. Up to 850°C , melt is produced gradually for the greywacke

789 composition whereas melting of the metapelite occurs as a pulse in the narrow muscovite–K-
790 feldspar field followed by more gradual melting due to biotite breakdown.

791 Beyond the loss of biotite from the stable assemblage, melting continues through the
792 breakdown of K-feldspar, plagioclase and quartz and an additional 4 mol% melt is produced by
793 900°C. Although monazite dissolution is modelled to continue from 850 to 900°C, some
794 prograde monazite crystallization may be possible in this temperature range for three reasons.
795 First, the melt is saturated with respect to LREE due to progressive monazite dissolution.
796 Second, after the exhaustion of biotite the melt becomes progressively drier, which decreases the
797 solubility of monazite. Third, melt ASI increases, which increases the dissolution of apatite and
798 may liberate enough LREE to support new monazite growth. However, unless this monazite is
799 sequestered away in a growing peritectic mineral (such as garnet) it is expected to be consumed
800 during further heating and partial melting.

801 All of the monazite and ~65% of the zircon are consumed by 900°C. Isothermal
802 decompression from 9 kbar to 7 kbar produces an additional 4 mol% melt and results in minor
803 (<1 mol%) garnet consumption. Melt ASI increases from 1.18 to 1.23 during decompression,
804 which enhances the solubility of apatite. However, the melt is undersaturated with respect to
805 LREE and new monazite growth is not expected during decompression. Approximately 10% of
806 the initial amount of zircon present at the solidus is consumed during decompression and titania
807 activity decreases from 0.95 to 0.85.

808 Cooling and decompression from peak T to 750°C results in: (1) melt crystallization, (2)
809 the consumption of garnet and sillimanite to produce cordierite commencing at 890°C and biotite
810 starting at 840°C, (3) a decrease in a_{TiO_2} from 0.85 to 0.65, (4) a decrease in melt ASI from 1.23
811 to 1.16, (5) the growth of new zircon (likely as overgrowths on existing zircon), and (6) the

812 crystallization of monazite starting at $\sim 885^{\circ}\text{C}$. Zircon and monazite growth occurs during garnet
813 breakdown and these accessory minerals are expected to have elevated Y and HREE
814 concentrations. The application of Ti-in-zircon thermometry should use an a_{TiO_2} value ranging
815 from 0.85 to 0.65. Assuming a a_{TiO_2} value of 1.0 would underestimate temperatures by up to
816 50°C (Fig. 1b).

817

818 ***Reaction sequence for the counterclockwise P–T path.*** The mode-box and titania
819 activity for the counterclockwise assemblage sequence are shown in Figure 10c. The amount of
820 monazite and zircon remaining as well as melt ASI are shown in Figure 10d. Similar to the
821 clockwise P–T path for the greywacke, the prograde segment of the counterclockwise P–T path
822 generates melt gradually in contrast to the more pulsed melting in the metapelite. Melting
823 commences at the wet solidus and proceeds via the breakdown of biotite to produce cordierite
824 and garnet followed by K-feldspar at 760°C . Orthopyroxene enters the phase assemblage at
825 825°C and biotite is completely consumed by 830°C . The value of a_{TiO_2} decreases from ~ 0.80 to
826 0.60 up to the orthopyroxene-in field boundary and then increases for the remainder of the
827 prograde path. The increase in pressure near peak T results in the breakdown of orthopyroxene to
828 produce garnet (e.g., White et al. 2008). By the end of the heating segment, $\sim 50\%$ of the zircon
829 and $\sim 70\%$ of the monazite that was present at the solidus has been consumed. Melt ASI varies
830 between 1.16 and 1.11 during the prograde path. As with the clockwise P–T path, melting above
831 the biotite-stability field (in this case from $830\text{--}860^{\circ}\text{C}$) has the potential to generate some
832 prograde monazite if LREE-rich apatite is consumed. A total of 18 mol% melt is predicted to be
833 generated during heating, which is significantly less than the 44 mol% produced along the same
834 P–T path for the metapelite.

835 Consistent with the metapelite for the same P - T path, isobaric cooling results in melt
836 crystallization and monazite and zircon growth. The mode of garnet decreases by 7 mol% and K-
837 feldspar breaks down to sillimanite in the presence of melt at 815°C. Zircon and monazite
838 crystallization is predicted to have elevated Y and HREE due to the breakdown of garnet. Zircon
839 growth occurs when the a_{TiO_2} value of the system is ~ 0.7 .

840

841 **Average Mid Ocean Ridge Basalt**

842 *Phase relations.* The P - T pseudosection for an average MORB composition is shown in
843 Figure 11. The wet solidus has a negative slope from 4 to 11.5 kbar and a positive slope at T
844 > 11.5 kbar. The temperature of the wet solidus ranges from 620 to 700°C over the modelled P - T
845 range. Garnet is stable from 10–14 kbar with decreasing temperature. Orthopyroxene enters the
846 assemblage at temperatures of 800°C to 900°C with increasing P and is not stable above 10 kbar
847 at high T . Garnet and orthopyroxene are only stable together at pressures of 9.5–10 kbar and at T
848 $> 900^\circ\text{C}$. Rutile is stable at $T > 790^\circ\text{C}$ at $P > 7$ kbar.

849

850 *Reaction sequence for the clockwise P - T path.* The mode-box and titania activity for the
851 reaction sequence as well as the amount of zircon remaining and melt ASI are shown in Figure
852 12. Melting starts at the wet solidus at 630°C and 9.7 kbar and generates ~ 2 mol% melt and ~ 2
853 mol% clinopyroxene at the expense of epidote and biotite. A minor amount (~ 2 mol%) of zircon
854 is expected to be consumed in order to saturate the melt in Zr. Epidote can be an important
855 source of Zr (e.g., Kohn et al. 2015) and the breakdown of Zr-rich epidote may result in Zr
856 saturation and minor zircon crystallization at this stage. After the exhaustion of biotite at
857 $\sim 650^\circ\text{C}$, melting continues via the breakdown of hornblende, titanite and quartz to produce an

858 additional 4 mol% melt as well as peritectic clinopyroxene by 800°C and 11.4 kbar. During this
859 portion of the prograde path, a_{TiO_2} increases from 0.6 at the wet solidus up to 0.9 at 800°C. The
860 amount of zircon remaining is ~80 mol% of the amount present at the solidus. Melt ASI
861 increases slightly from 0.99 to 1.00.

862 Garnet becomes stable at ~800°C and melting proceeds through the breakdown of
863 hornblende and titanite. Garnet growth may include zircon grains that would be isolated from the
864 reacting volume of the rock and zircon could be preserved to higher T . Rutile becomes stable at
865 818°C and titanite is completely consumed by 821°C. The peak pressure of 12 kbar is reached at
866 850°C and at this point the system contains ~13 mol% melt, 10 mol% garnet and 65% of the
867 zircon has been consumed. Although zircon is expected to be consumed during partial melting,
868 the breakdown of Zr-rich amphibole (e.g., Sláma et al. 2007) may yield enough Zr to
869 oversaturate the melt and grow new zircon; this zircon may be relatively depleted in HREE in
870 response to the presence of garnet in the rock.

871 The isobaric heating segment of the P - T path at peak P produces garnet at the expense of
872 hornblende and quartz. Quartz is exhausted at 905°C. This has implications for applying the Ti-
873 in-zircon thermometer, which uses a_{SiO_2} as a variable. Zircon is completely consumed by 880°C
874 and the melt produced at higher T is expected to be undersaturated in Zr. A consequence of this
875 is that prograde zircon growth at $T > 880^\circ\text{C}$ is unlikely because any excess Zr due to the
876 breakdown of other minerals (e.g., amphibole) will be incorporated into the Zr-undersaturated
877 melt. At the metamorphic peak of 950°C, the system contains ~28 mol% melt, ~20 mol% garnet
878 and ~14% hornblende.

879 Isothermal decompression from 12 to 10 kbar results in significant garnet consumption
880 (from 20 to 9 mol%), hornblende growth (from 14 to 24 mol%) and minor melt consumption

881 (from 28 to 27 mol%). Zirconium liberated from garnet breakdown is expected to be partitioned
882 between the Zr-undersaturated melt and hornblende. Consequently, zircon crystallization is
883 unlikely. Orthopyroxene enters the assemblage at 10.1 kbar and grows at the expense of garnet,
884 which is exhausted by 9.7 kbar. If garnet is completely consumed, any liberated zircon may
885 become available to the reacting volume and will likely be consumed into the Zr-undersaturated
886 melt. Rutile is completely consumed by 9.1 kbar and a_{TiO_2} decreases with further decompression.
887 At the end of the isothermal decompression segment of the P - T path the amount of
888 orthopyroxene reaches 5 mol%. No new zircon growth is expected during the isothermal
889 decompression segment because: (1) there is a <1 mol% melt change during isothermal
890 decompression and the melt is already significantly undersaturated in Zr, (2) the M value of the
891 melt (cation ratio of $[\text{Na} + \text{K} + 2\text{Ca}] / [\text{Al} \times \text{Si}]$) increases from 1.6 to 1.7, which results in an
892 increase in the Zr required to saturate the melt (Harrison and Watson 1983; Boehnke et al. 2013),
893 (3) the mode of hornblende increases from 14 to 23 mol%, likely accommodating significant
894 amounts of Zr at high temperature (e.g., Kohn et al. 2015).

895 Cooling and decompression from 950°C at 8 kbar to 800°C at 6 kbar results in the
896 consumption of 17 mol% melt and the complete breakdown of orthopyroxene by ~830°C.
897 Zirconium saturation of the melt is reached at ~870°C and zircon begins to crystallize. At
898 ~850°C, approximately 20% of the amount of zircon originally present at the solidus has grown
899 back and this zircon is expected to be enriched in HREE because there is no garnet present in the
900 system. Zircon growth at this stage occurs in a system with an a_{TiO_2} value of 0.9 and in the
901 absence of quartz, which is important for the application of Ti-in-zircon thermometry.

902

903 **Summary of reaction sequence modelling**

904 Linking the trace element concentrations of accessory minerals to the key major minerals
905 (e.g., garnet and plagioclase) requires an understanding of the reaction sequences for a particular
906 bulk composition. Key minerals like garnet and plagioclase can grow or be consumed multiple
907 times along a P – T path during heating and cooling or burial and exhumation (Figs 3–10). For
908 example, garnet is predicted to grow in the subsolidus greywacke composition during heating
909 and burial and garnet breakdown occurs during cooling and exhumation for the hairpin P – T path
910 (Fig. 6b). By contrast, garnet growth and consumption occurs multiple times along the heating
911 and increasing pressure segment of the P – T paths for the subsolidus metapelite composition (Fig.
912 4). Therefore, high-Y and high-HREE zones in monazite that can be linked to garnet breakdown
913 may reflect the burial and/or exhumation portions of a P – T path.

914 Titania activity also varies along the modelled P – T paths and this needs to be taken into
915 account when applying Ti-in-zircon or Ti-in-quartz thermometers. In general, new zircon growth
916 in subsolidus rocks is expected to be limited because major minerals such as hornblende and
917 garnet can accommodate more Zr with increasing temperature. One exception to this may be
918 minor zircon growth through the breakdown of Zr-rich epidote. For the suprasolidus P – T paths
919 for the metapelite and greywacke, new zircon growth is expected during cooling and melt
920 crystallization; this growth is predicted to occur when bulk rock a_{TiO_2} is less than one for all of
921 the modelled P – T paths. If the Ti-in-zircon thermometer is applied with an a_{TiO_2} value of one,
922 then the result will be an underestimation in peak metamorphic temperatures by up to $\sim 40^\circ\text{C}$. For
923 the MORB composition, new zircon growth occurs at $a_{\text{TiO}_2} < 1.0$ as cooling and melt
924 crystallization occurs outside the stability field of rutile (Fig. 12).

925 In suprasolidus metamorphic rocks, zircon and monazite are expected to be consumed
926 along the prograde path and new growth is generally predicted to occur along the cooling path.

927 Zircon and monazite dissolution is non-linear and the rate increases with increasing temperature.
928 For the clockwise P - T path for the metapelite and greywacke, an increase in the melt ASI leads
929 to more apatite dissolution. Because the melt is saturated in LREE with respect to monazite up to
930 820°C, apatite breakdown may liberate enough LREE to promote new monazite growth.
931 However, after the exhaustion of monazite, the melt is expected to be undersaturated in LREE
932 and prograde monazite growth at the expense of LREE-rich apatite is unlikely. For the
933 counterclockwise P - T path for the metapelite and greywacke, melt ASI increases and decreases
934 during the P - T evolution and monazite growth from apatite breakdown will be more complex to
935 interpret.

936 The metapelite is expected to lose most of the inherited or subsolidus prograde monazite
937 and zircon during heating above the solidus (unless these mineral are sequestered away from the
938 reaction volume) whereas a larger proportion of these minerals remains in the greywacke
939 composition except for the clockwise P - T path where monazite is completely consumed for the
940 greywacke. The difference reflects the fertility of the two rocks; pelites generate more melt than
941 the greywacke and require more Zr and LREE to saturate this melt. Therefore, less fertile
942 compositions, such as greywackes, are more likely to preserve subsolidus zircon and monazite.

943

944

COMPLICATING FACTORS

945 **Changes in effective bulk composition**

946 Phase equilibria modelling requires an assessment of the effective bulk composition of a
947 system that is used to model the P - T phase relations for a rock (e.g., Stüwe 1997). The effective
948 bulk composition approximates the composition ‘available’ to the rock in which equilibrated
949 mineral assemblages and reaction sequence develop. In natural systems, the effective bulk

950 composition may change along a P – T path (e.g., Guevara and Caddick 2016), which has
951 subsequent implications for modelling reaction sequences in both subsolidus and suprasolidus
952 metamorphic rocks. There are two main mechanisms that can change the effective bulk
953 composition of a metamorphic system: fractionation of elements into growing porphyroblasts
954 and melt loss.

955 Growing porphyroblasts can fractionate certain elements into their cores that become
956 unavailable to the reacting system in the remainder of the rock. For example, chemical zoning in
957 garnet is commonly preserved in metamorphic rocks because cation diffusion in garnet is
958 relatively slow (e.g., Chakraborty and Ganguly 1992). The preferential partitioning of elements
959 into garnet cores reduces their effective composition in the reactive volume of the rock (e.g.,
960 Lanari and Engi, 2017). Some of the consequences for using phase diagrams to infer
961 metamorphic conditions considering garnet fractionation include reduced stability fields for
962 garnet (Gaides et al. 2008) and other minerals (Zuluaga et al. 2005; Moynihan and Pattison
963 2013) as well as changes in mineral compositional isopleths for garnet (Evans 2004; Gaides et al.
964 2006) and plagioclase (Moynihan and Pattison 2013). While crystal fractionation needs to be
965 assessed on a case-by-case basis, it is generally most important to consider for greenschist- and
966 amphibolite-facies metamorphic assemblages where mineral chemistries are particularly useful
967 for determining P – T conditions. At higher grades, the use of mineral composition isopleths is
968 generally less effective due to retrograde exchange reactions (e.g., Spear and Florence 1992;
969 Kohn and Spear 2000; Pattison et al. 2003). For zircon, the fractionation of Zr into growing
970 garnet (e.g., Kohn et al. 2015) may reduce the Zr budget available for zircon growth.

971 In high-grade metamorphic rocks that underwent anatexis the preservation of peritectic
972 minerals and lack of extensive retrogression supports melt drainage during prograde

973 metamorphism (Fyfe 1973; Powell 1983; White and Powell 2002; Guernina and Sawyer 2003;
974 Reno et al. 2012). Melt loss produces progressively more refractory bulk compositions, which
975 results in elevated solidus temperatures in the residual rocks. In migmatites that have undergone
976 melt loss, suprasolidus zircon and monazite growth is generally expected to occur during cooling
977 from peak T to the elevated solidus (e.g., Kelsey et al. 2008; Spear and Pyle 2010; Yakymchuk
978 and Brown 2014b). Therefore, rocks that have experienced identical P – T histories but variable
979 amounts of melt loss and have different solidus temperatures should record a range of ages (e.g.,
980 Korhonen et al. 2013b).

981

982 **Bulk composition and the suprasolidus behaviour of zircon and monazite**

983 For the modelled suprasolidus reaction sequences, the preservation of subsolidus (e.g.,
984 prograde or inherited) zircon and monazite is mainly related to the fertility of the rocks. The
985 metapelite generates more melt along the same P – T path than the greywacke. Consequently,
986 zircon and monazite are completely consumed for the metapelite composition along both P – T
987 paths (Figs 8a, b) whereas some zircon and/or monazite can survive in the greywacke. An
988 additional factor is the bulk rock content of Zr for zircon and LREE and phosphorus for monazite
989 (e.g., Kelsey et al. 2008; Yakymchuk and Brown 2014b). Both of these factors are explored
990 together in Figure 13.

991 The temperature–composition diagrams in Figure 13 illustrate the variation in melt mode
992 and the stability of zircon and monazite for compositions ranging linearly from the metapelite
993 (left side) to that of the greywacke (right side). The diagrams are isobaric and were calculated at
994 7 kbar because this pressure intersects the main melt producing reactions for both bulk
995 compositions. The common reaction to both compositions is the breakdown of biotite to produce

996 cordierite at ~840–845°C (Fig. 13a). The amount of melt in the metapelite composition is
997 roughly twice the amount in the greywacke composition over the range of modelled temperatures
998 (Fig. 13b). For a bulk composition of 150 ppm Zr, zircon is completely consumed in the
999 metapelite composition by 925°C whereas only 30% of the zircon has been consumed in the
1000 greywacke composition at the same temperature (Fig. 13c). For a bulk composition of 150 ppm
1001 LREE, monazite is completely consumed by 830°C for the metapelite and 880°C for the
1002 greywacke (Fig. 13d). The slopes of the dissolution contours are steeper for zircon than for
1003 monazite in Figures 13c and 13d. This indicates that zircon dissolution is more sensitive to bulk
1004 composition (metapelite vs. greywacke) than monazite.

1005 The sensitivity of zircon and monazite dissolution to bulk composition of Zr and LREE is
1006 explored in Figures 13e and 13f. The contours represent the complete dissolution of zircon and
1007 monazite for various bulk rock Zr and LREE contents. For low-Zr pelites with concentrations of
1008 50 ppm, zircon is completely consumed by 860°C, which is 65°C lower than for a closer-to-
1009 average Zr concentration of 150 ppm. Greywackes with low concentrations of Zr (e.g., 50 ppm)
1010 can still preserve zircon up to UHT conditions due to their lower fertility. Monazite dissolution
1011 contours in Figure 13f have shallower slopes than those for zircon (Fig. 13e), which again
1012 suggests that monazite dissolution is less sensitive to bulk composition. A metapelite with low
1013 concentrations of LREE (e.g., 50 ppm) is predicted to lose monazite by 780°C and monazite is
1014 expected to be completely consumed in the greywacke composition by 820°C. Preserving
1015 subsolidus monazite to UHT conditions requires very LREE-rich compositions of >400 ppm for
1016 the metapelite and >180 ppm for the greywacke. Therefore, a strategy for finding subsolidus
1017 prograde zircon and monazite in migmatites is to choose samples with the highest concentrations
1018 of Zr and LREE.

1019 The breakdown of LREE-rich apatite is a potential mechanism to promote prograde
1020 suprasolidus monazite growth (e.g., Johnson et al. 2015). This is most likely to occur when the
1021 anatectic melt is saturated with respect to monazite in LREE. After the exhaustion of monazite,
1022 the melt is expected to be undersaturated and LREE liberated from the breakdown of apatite is
1023 not predicted to generate new prograde monazite growth. In principle, the higher the bulk rock
1024 LREE the longer monazite will persist during a suprasolidus heating path and the longer the melt
1025 will remain saturated with respect to LREE (Fig. 13f). A further consideration is that bulk
1026 compositions with low phosphorus concentrations may lose apatite during the prograde path and
1027 would result in melt that is undersaturated in P. This may promote further monazite dissolution
1028 instead of monazite growth. Therefore, LREE and P-rich bulk compositions should be targeted
1029 for accessory mineral geochronology to constrain the timing of suprasolidus prograde monazite
1030 growth.

1031

1032 **Effects of open system behaviour on accessory minerals**

1033 Dissolution/re-precipitation of accessory minerals in metamorphic rocks due to the
1034 infiltration of an externally derived fluid has been documented in experiments and studies of
1035 natural samples (e.g., Tomaschek et al. 2003; Crowley et al. 2008; Harlov and Hetherington
1036 2010; Blereau et al. 2016). The careful integration of petrography with the chemistries of these
1037 minerals can be used to provide information on the timing of fluid infiltration (e.g., Williams et
1038 al. 2011) and on fluid chemistry (e.g., Harlov et al. 2011; Taylor et al. 2014; Shazia et al. 2015).
1039 However, the timing of fluid ingress relative to the metamorphic history and the chemistry of
1040 these fluids are highly variable and should to be assessed on a case-by-case basis.

1041 For migmatites that have lost melt, the extraction of melt saturated in Zr or LREE will
1042 also change the effective composition of the residuum (e.g., Rapp et al. 1987), which has
1043 subsequent consequences for zircon and monazite dissolution. For example, consider the
1044 muscovite to K-feldspar melting reaction for the metapelite along the clockwise P - T path at
1045 $\sim 730^\circ\text{C}$ and 7.5 kbar and consider bulk rock values of 150 ppm for LREE and Zr. This reaction
1046 involves a large positive volume change (e.g., Powell et al. 2005), which may be accommodated
1047 by melt extraction from the rock. When muscovite is exhausted the rock contains ~ 9 mol% melt.
1048 The saturation concentrations of LREE and Zr in the melt at this point are 195 ppm and 137
1049 ppm, respectively. Assuming 8 vol% melt (approximately equivalent to 8 mol%) is extracted
1050 (leaving 1 vol% in the rock along grain boundaries), mass balance can be used to determine the
1051 amount of Zr and LREE left in the system. For this example, the effective concentration of
1052 LREE in the rock decreases to 146 ppm and the effective concentration of Zr increases to 151
1053 ppm. Although the changes to the bulk composition are minor in this example, monazite is
1054 predicted to be completely consumed at lower T and zircon may persist to higher T than for
1055 closed system (Fig. 13). For rocks with very low concentrations of Zr and LREE, melt extraction
1056 can have a more significant impact on the effective bulk concentrations of these elements and the
1057 stability of zircon and monazite (e.g., Yakymchuk and Brown 2014b).

1058

1059 **Inclusion/host relationships**

1060 An important consideration for accessory mineral reactivity is their inclusion in the major
1061 rock-forming minerals (Watson et al. 1989; Bea et al. 2006). Watson et al. (1989) showed that
1062 for a Himalayan migmatite sample roughly 78% of the zircon mass is located along grain
1063 boundaries and the remaining 28% is included in major minerals (predominately biotite and

1064 garnet), though how representative this is of typical migmatitic gneiss is unknown.
1065 Consequently, the breakdown of major minerals may liberate accessory minerals into the
1066 reacting volume of the rock that would otherwise be sequestered away.

1067 Inclusions of zircon and monazite that are isolated from the reaction volume may also
1068 reduce the effective bulk rock concentration of Zr and LREE (Yakymchuk and Brown 2014b).
1069 For example, consider a bulk rock composition of LREE with half of the monazite sequestered
1070 away from the reaction volume as inclusions. For a metapelite with a bulk rock LREE
1071 composition of 300 ppm and considering that half of this is unavailable, the effective
1072 concentration of LREE is 150 ppm. The complete dissolution of monazite is modelled to occur at
1073 830°C in contrast to 870°C for the scenario where all monazite is available for reaction (Fig.
1074 13f). Therefore, applying the models in Figure 13 to natural examples requires an estimate of the
1075 amount of zircon or monazite sequestered away from the reaction volume as inclusions as well
1076 as an estimate of the amount of Zr and LREE locked away in the major minerals.

1077 The heterogeneous distribution of melt and minerals in high-grade metamorphic rocks
1078 also has implications for the dissolution and preservation of accessory minerals. Even for an
1079 initially homogenous protolith, *in situ* melt may be spatially associated with peritectic minerals
1080 in isolated patches; this produces a heterogeneous melt framework (e.g., White et al. 2004).
1081 Zircon and monazite proximal to the zones of incipient melting and in chemical communication
1082 with this melt are more likely to contribute to Zr and LREE saturation of the melt whereas more
1083 distal grains may not. Consequently, detrital, inherited or prograde (subsolidus or early
1084 suprasolidus) zircon and monazite are more likely to be preserved in domains away from
1085 incipient melting whereas post-peak and retrograde zircon and monazite may be spatially
1086 associated with *in situ* leucosome.

1087

1088

CONCLUDING REMARKS

1089

One important facet of petrochronology is to link the ages of accessory mineral

1090

chronometers to the P - T information obtained from major rock-forming minerals in

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metamorphic rocks. The growth and consumption of major minerals is important because these

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minerals: (1) may contain the necessary essential structural constituents to promote accessory

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mineral growth directly from their breakdown, (2) are repositories of the trace elements used to

1094

link accessory mineral chronometers to P - T conditions (e.g., Sr and Eu related to stability of

1095

plagioclase as well as Y and HREE reflecting the growth/consumption of garnet), (3) are

1096

important hosts for accessory mineral inclusions, and (4) play a role in controlling the

1097

component activities (e.g., a_{TiO_2}) along a P - T evolution. The main controls on accessory mineral

1098

behaviour differ between subsolidus and suprasolidus metamorphic conditions.

1099

For subsolidus metamorphism, zircon is generally unreactive and monazite can grow

1100

during the prograde and retrograde segments. Linking ages from these accessory mineral to a

1101

metamorphic history requires an understanding of the major mineral reaction sequence as well as

1102

the behaviour of accessory minerals like xenotime, apatite and allanite. Major minerals such as

1103

garnet or plagioclase may experience growth and breakdown stages at any point along a P - T

1104

path: linking their behaviour to the trace element chemistries of accessory minerals requires an

1105

assessment of the reaction sequence for a particular rock along a well-constrained P - T path.

1106

For suprasolidus metamorphism, phase equilibria modelling predicts that both zircon and

1107

monazite will be consumed during prograde metamorphism and grow during cooling and melt

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crystallization. However, this contrasts with some studies that have convincingly showed

1109

evidence for suprasolidus prograde zircon and monazite growth. For monazite, apatite

1110 dissolution may have contributed to minor prograde monazite growth if the anatectic melt is
1111 saturated in LREE. For zircon, solid-state breakdown of major minerals that contain appreciable
1112 quantities of Zr may facilitate prograde zircon growth. Ostwald ripening may also play a role in
1113 the prograde growth of both zircon and monazite, but this mechanism is still incompletely
1114 understood.

1115

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1121

1122 **References**

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1697 **Figure Captions**

1698 **Figure 1.** Underestimation of metamorphic temperature when the bulk rock value of a_{TiO_2} is less
1699 than 1.0 (the true value is shown by the contours), but a value of 1.0 is assumed in: (a) the Ti-in-
1700 quartz thermometer of Wark and Watson (2006), and (b) the Ti-in-zircon thermometer of Ferry
1701 and Watson (2007).

1702

1703 **Figure 2.** Compilation of the range of allanite to monazite reaction temperatures for variable
1704 bulk rock pelite compositions. Curves are from Spear (2010).

1705

1706 **Figure 3.** P – T pseudosection for the subsolidus metapelite. The solidus is shown by the heavy
1707 dashed line.

1708

1709 **Figure 4.** (a) Mode-box diagram and a_{TiO_2} for the subsolidus clockwise P – T path for the
1710 metapelite. (b) Mode-box diagram and a_{TiO_2} for the subsolidus hairpin P – T path.

1711

1712 **Figure 5.** P – T pseudosection for the subsolidus greywacke. The solidus is shown by the heavy
1713 dashed line.

1714

1715 **Figure 6.** (a) Mode-box diagram and a_{TiO_2} for the subsolidus clockwise P – T path for the
1716 greywacke composition. (b) Mode-box diagram and a_{TiO_2} for the subsolidus hairpin P – T path.

1717

1718 **Figure 7.** P – T pseudosection for suprasolidus metapelite. The solidus is shown by the heavy
1719 dashed line.

1720

1721 **Figure 8.** (a) Mode-box diagram and a_{TiO_2} for the suprasolidus clockwise P – T path for the
1722 metapelite. (b) Amount of zircon and monazite remaining relative to the amount at the solidus
1723 and the ASI value of melt. (c) Mode-box diagram and a_{TiO_2} for the counterclockwise P – T path
1724 for the metapelite. (d) Amount of zircon and monazite remaining relative to the amount at the
1725 solidus for the counterclockwise P – T path.

1726

1727 **Figure 9.** P – T pseudosection for the suprasolidus greywacke. The solidus is shown by the heavy
1728 dashed line.

1729

1730 **Figure 10.** (a) Mode-box diagram and a_{TiO_2} for the suprasolidus clockwise P – T path for the
1731 greywacke. (b) Amount of zircon and monazite remaining relative to the amount at the solidus
1732 and the ASI value of melt. (c) Mode-box diagram and a_{TiO_2} for the counterclockwise P – T path
1733 for the greywacke. (d) Amount of zircon and monazite remaining relative to the amount at the
1734 solidus for the counterclockwise P – T path.

1735

1736 **Figure 11.** P – T pseudosection for the suprasolidus MORB. The solidus is shown by the heavy
1737 dashed line.

1738

1739 **Figure 12.** (a) Mode-box diagram and a_{TiO_2} for the suprasolidus clockwise P – T path for the
 1740 MORB. (b) Amount of zircon remaining relative to the amount at the solidus and the ASI value
 1741 of melt.

1742

1743 **Figure 13.** (a) T – X pseudosection calculated at 7 kbar showing the change in mineral
 1744 assemblages for compositions ranging from the metapelite (left side) to the greywacke (right
 1745 side). (b) T – X diagram with melt isopleths (mol%). (c) T – X diagram with the calculated amount
 1746 of zircon dissolution assuming a bulk Zr concentration of 150 ppm. (d) Calculated amount of
 1747 monazite dissolution assuming a bulk LREE concentration of 150 ppm. (e) T – X diagram with
 1748 contours for the complete dissolution of zircon for a range of bulk rock Zr concentrations. (f) T –
 1749 X diagram with contours for the complete dissolution of monazite for a range of bulk rock LREE
 1750 concentrations.

1751

1752

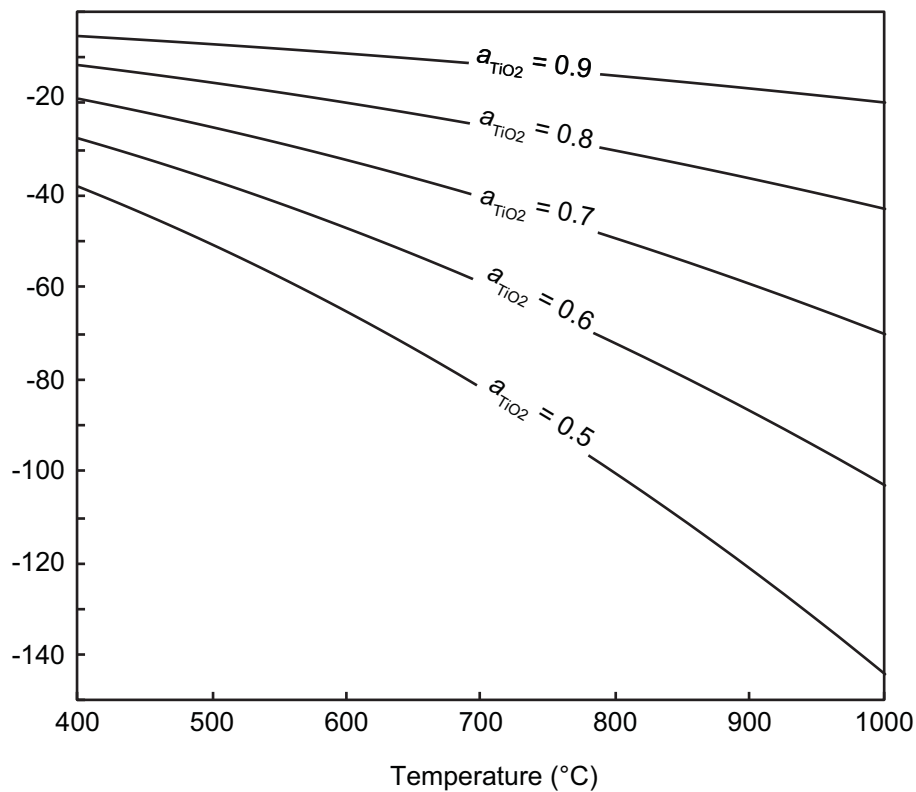
Table 1. Bulk compositions used for phase equilibria modelling (mol%)

	Figures	H ₂ O	SiO ₂	Al ₂ O ₃	CaO	MgO	FeO	K ₂ O	Na ₂ O	TiO ₂	MnO	O
<i>subsolidus</i>												
metapelite	3,4	+	64.58	13.65	1.59	5.53	8.03	2.94	2.00	0.91	0.17	0.60
greywacke	5,6	+	77.62	8.20	1.15	3.85	4.21	1.26	2.85	0.52	0.35	0.32
<i>suprasolidus</i>												
metapelite	7,8,13	6.24	60.55	12.80	1.49	5.18	7.52	2.76	1.88	0.85	0.16	0.60
greywacke	9,10,13	2.61	75.35	7.96	1.12	3.74	4.09	1.22	2.77	0.51	0.34	0.31
MORB	11,12	6.19	42.58	10.45	12.27	14.76	9.68	0.22	2.62	0.66	–	0.58

1753

+ H₂O in excess, – not considered

(a) Ti-in-quartz thermometer



(b) Ti-in-zircon thermometer

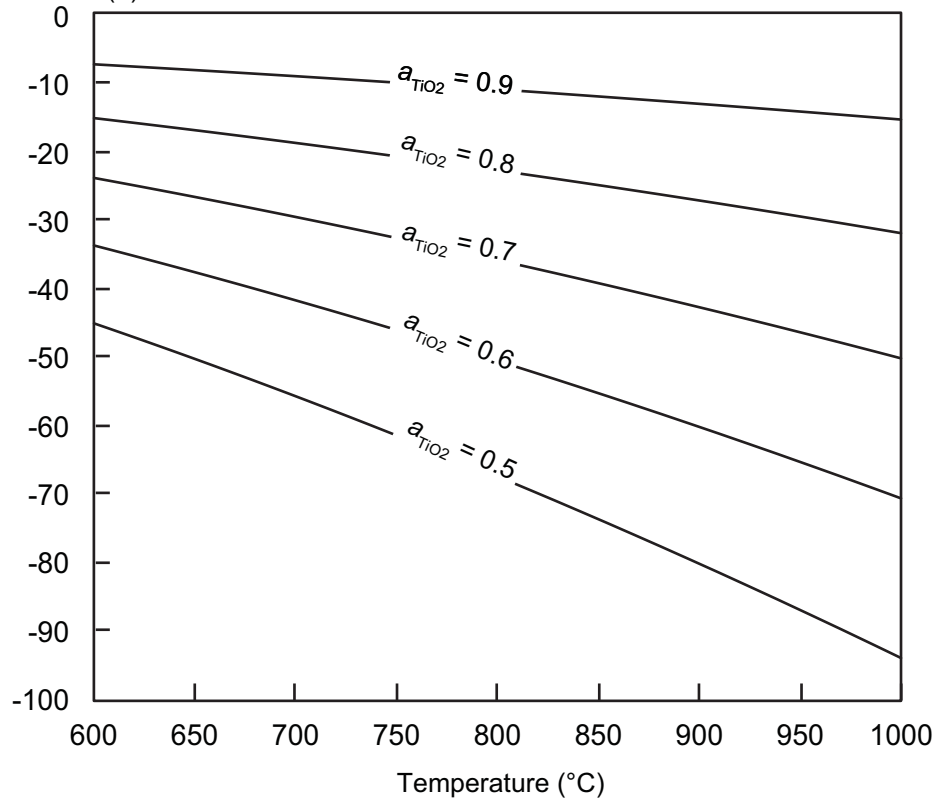


Figure 1

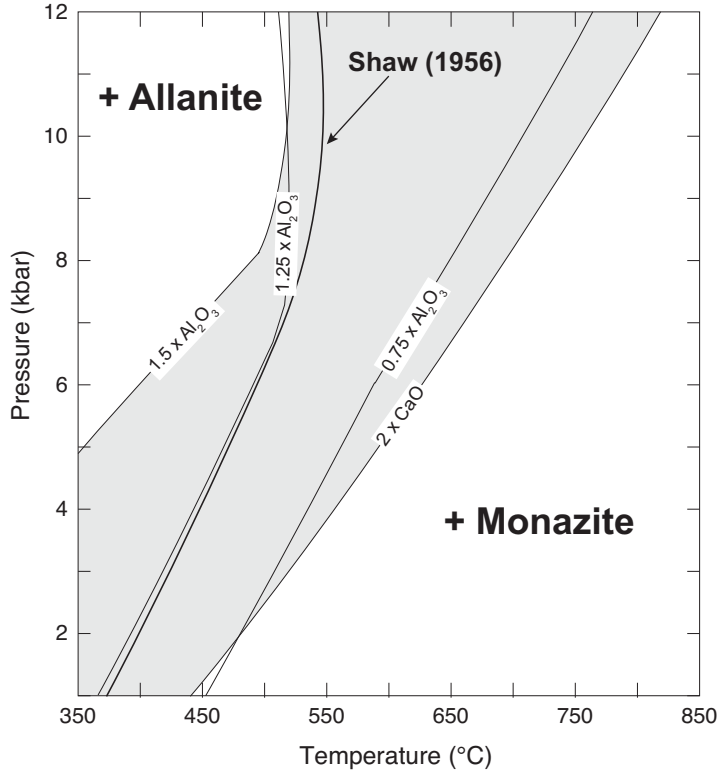
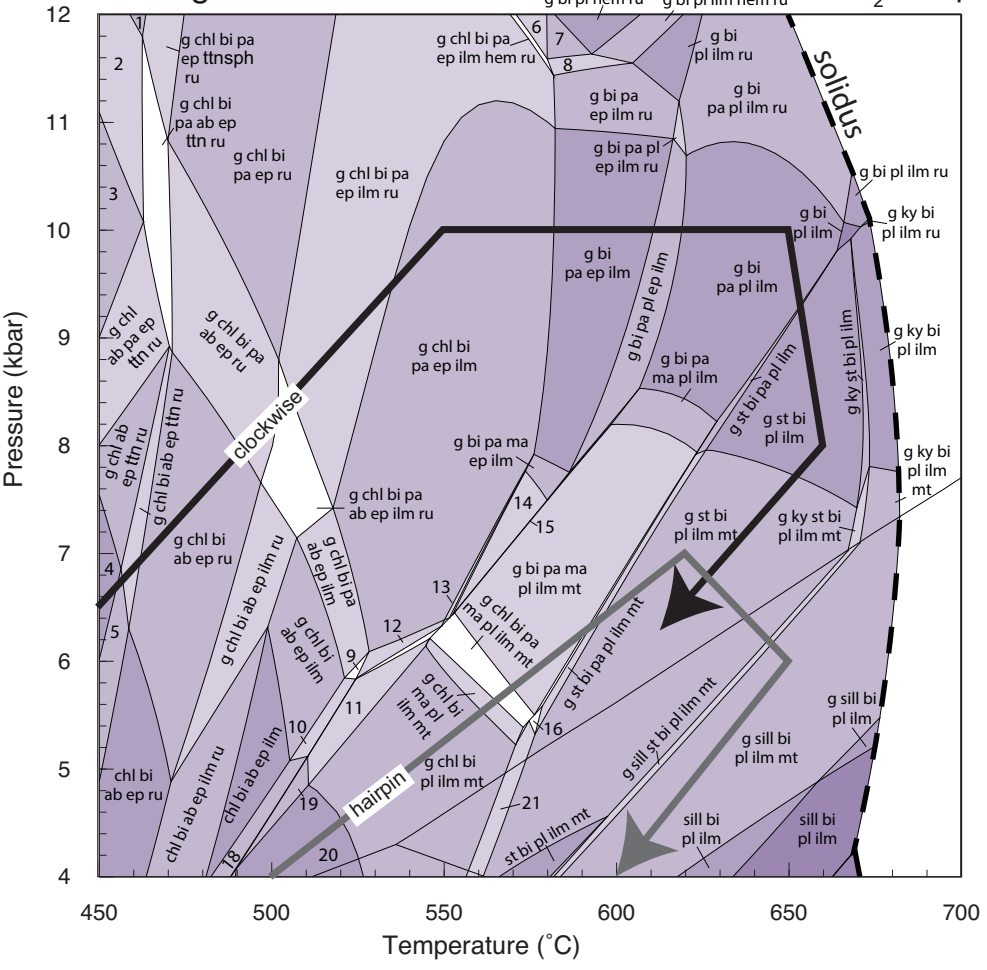


Figure 2

Average Pelite

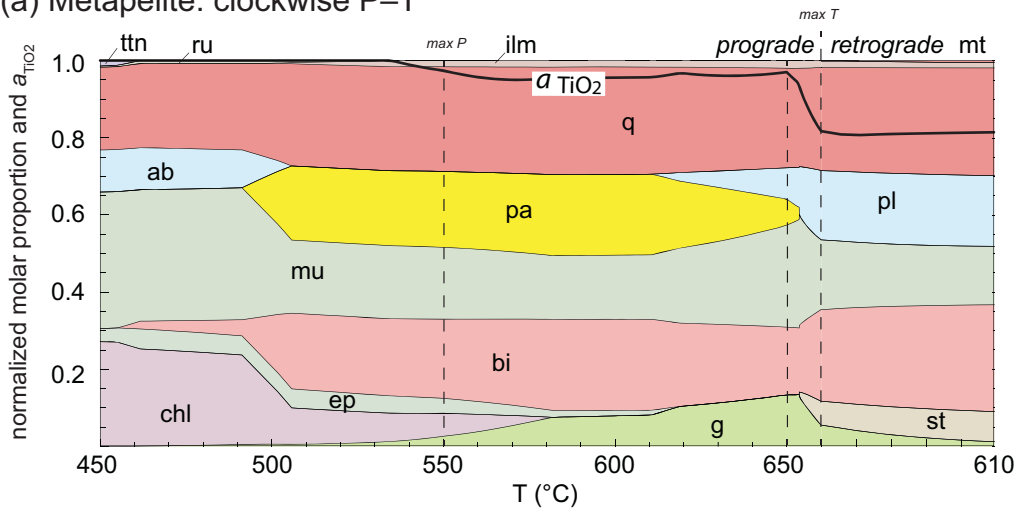
+ H₂O + q



- | | | | |
|--------------------------|-----------------------------|-----------------------------|------------------------------|
| 1) g chl bi pa ep ttn | 6) g chl bi pa ep hem ru | 11) g chl bi pl ep ilm mt | 16) g st chl bi pa pl ilm mt |
| 2) g chl bi pa ab ep ttn | 7) g bi pa ep hem ru | 12) g chl bi pa pl ep ilm | 17) chl bi pl ab ep ilm ru |
| 3) g chl pa ab ep ttn | 8) g bi pa ep ilm hem ru | 13) g chl bi pa ma ep ilm | 18) chl bi pl ab ep ilm |
| 4) chl ab ep ttn ru | 9) g chl bi pa pl ab ep ilm | 14) g bi pa ma ep ilm mt | 19) chl bi pl ep ilm mt |
| 5) chl bi ab ep ttn ru | 10) g chl bi pl ab ep ilm | 15) g bi pa ma pl ep ilm mt | 20) chl bi pl ilm mt |
| | | | 21) g st chl bi pl ilm mt |

Figure 3

(a) Metapelite: clockwise P-T



(b) Metapelite: hairpin P-T

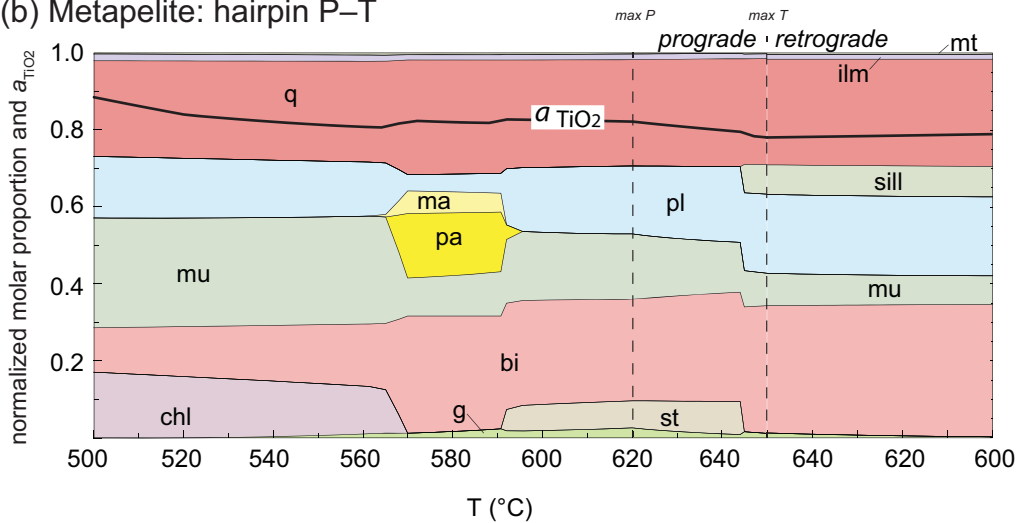
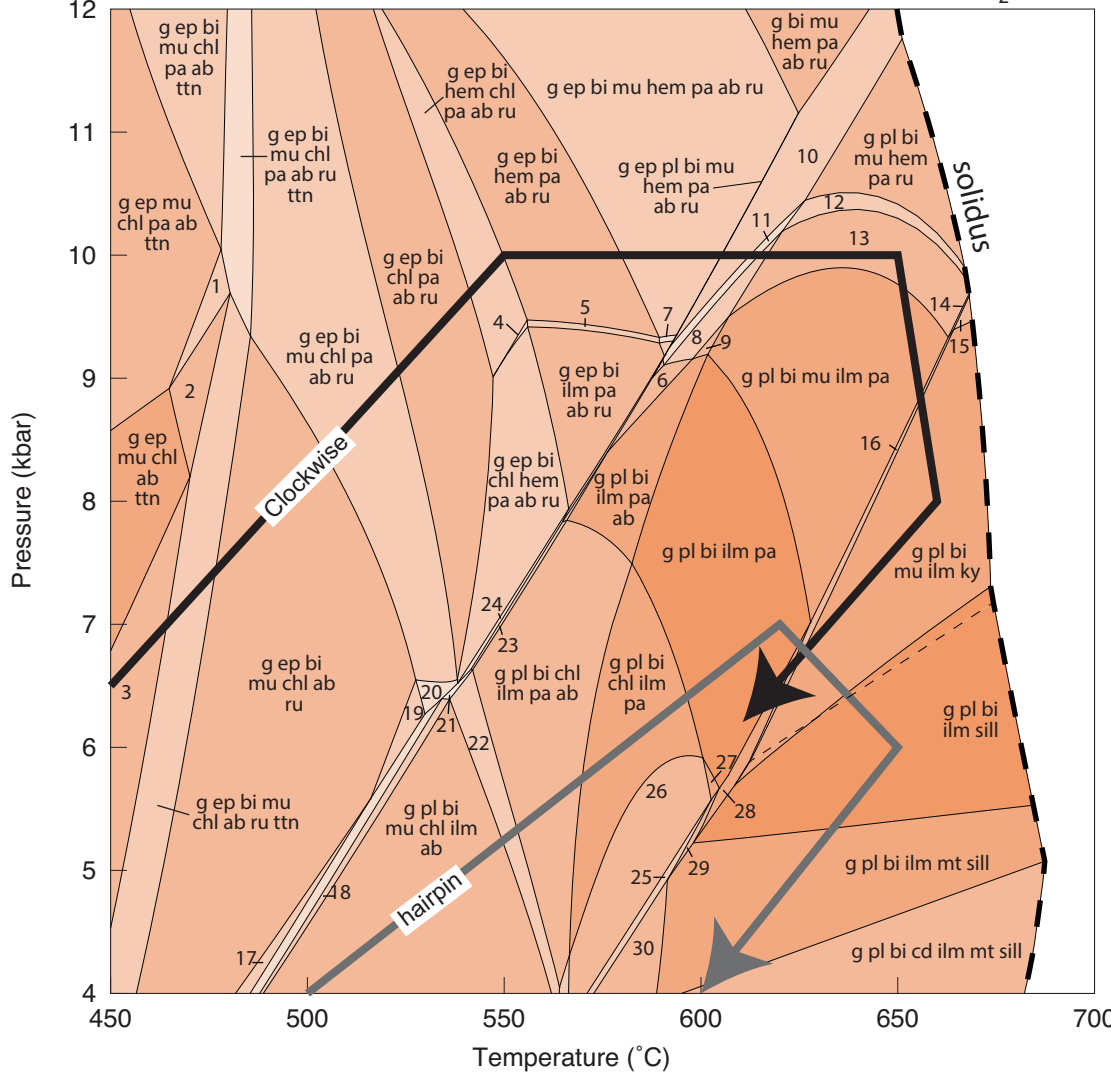


Figure 4.

Average passive margin greywacke

+ H₂O + q



- | | | |
|-----------------------------------|---------------------------------|------------------------------------|
| 1. g ep mu chl pa ab ru ttn | 11. g pl bi mu ilm hem pa ab ru | 21. g pl ep bi mu chl ilm pa ab ru |
| 2. g ep mu chl ab ru ttn | 12. g pl bi mu ilm hem pa ru | 22. g pl bi mu chl ilm pa ab |
| 3. g ep bi mu chl ab ttn | 13. g pl bi mu ilm pa ru | 23. g pl ep bi chl ilm pa ab |
| 4. g ep bi chl ilm hem pa ab ru | 14. g pl bi mu ilm pa ru ky | 24. g pl ep bi chl ilm pa ab ru |
| 5. g ep bi ilm hem pa ab ru | 15. g pl bi mu ilm ru ky | 25. g pl bi chl ilm pa mt sill |
| 6. g pl bi ilm pa ab ru | 16. g pl bi mu ilm pa ky | 26. g pl bi chl ilm pa mt |
| 7. g ep pl bi mu ilm hem pa ab ru | 17. g pl ep bi mu chl ab ru | 27. g pl bi ilm pa mt |
| 8. g pl bi mu ilm pa ab ru | 18. g pl ep bi mu chl ilm ab | 28. g pl bi ilm pa sill |
| 9. g pl bi mu ilm pa ab | 19. g ep bi mu chl ilm ab ru | 29. g pl bi ilm pa mt sill |
| 10. g pl bi mu hem pa ab ru | 20. g ep bi mu chl ilm pa ab ru | 30. g pl bi chl ilm mt sill |

Figure 5

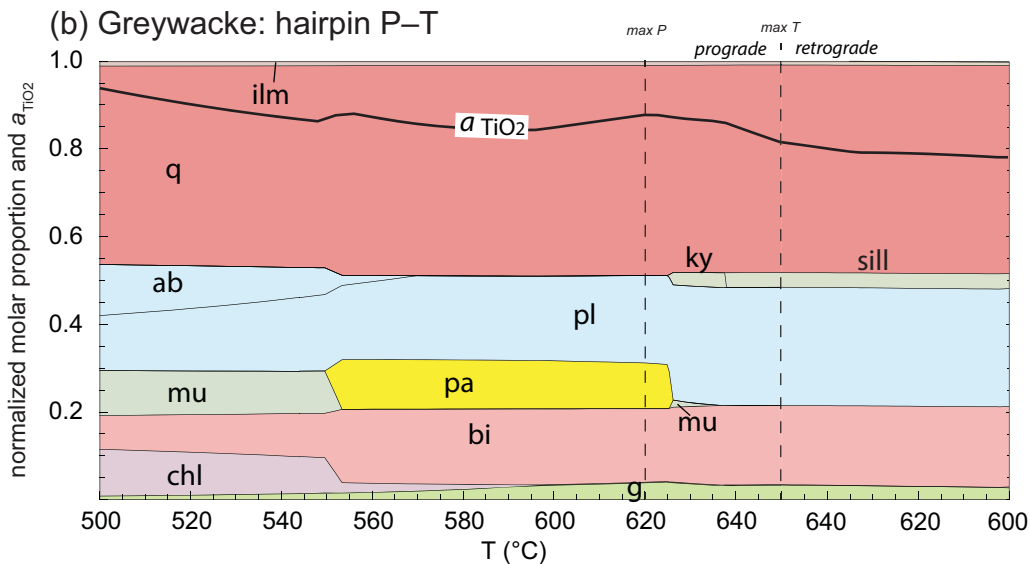
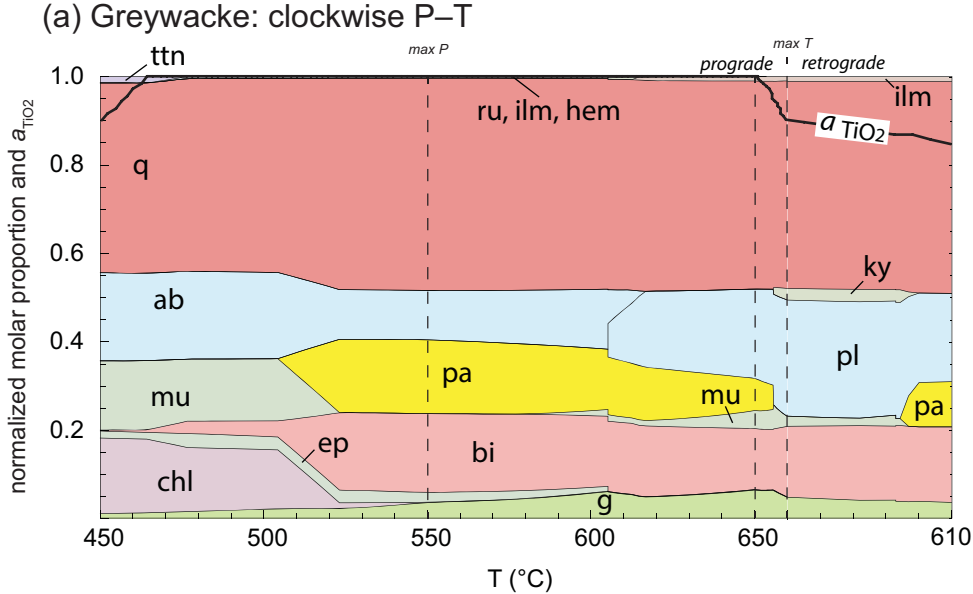
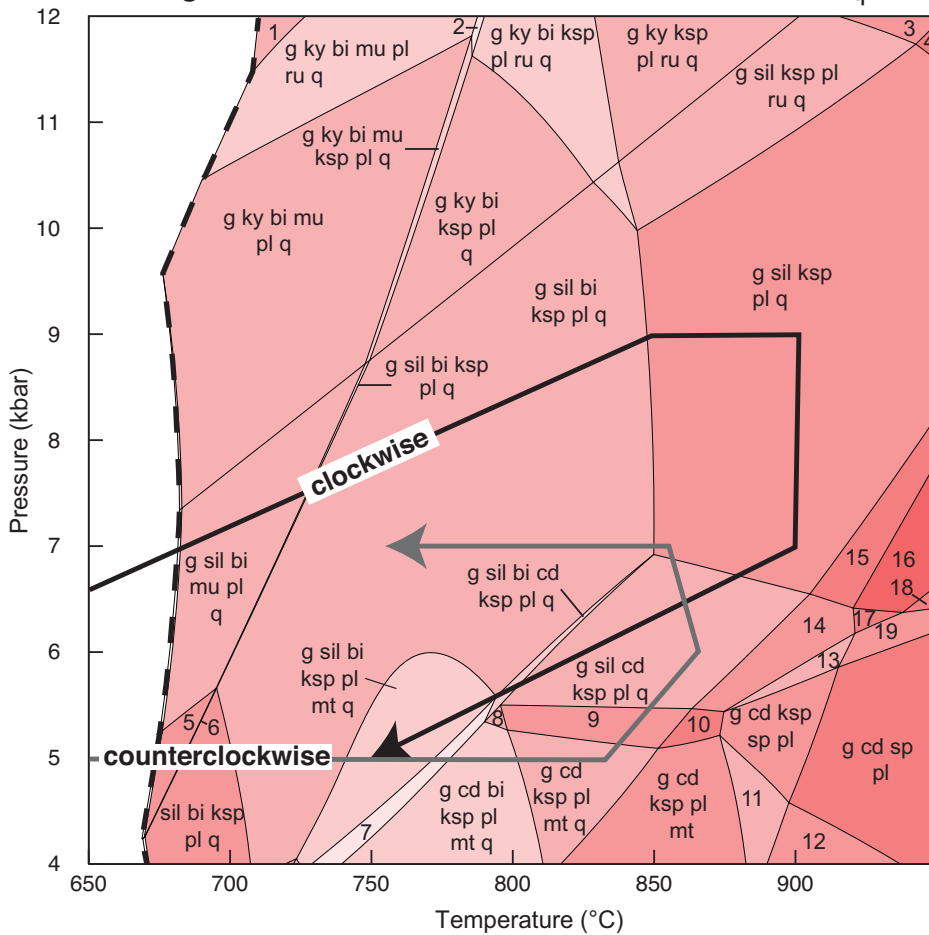


Figure 6

Average Pelite

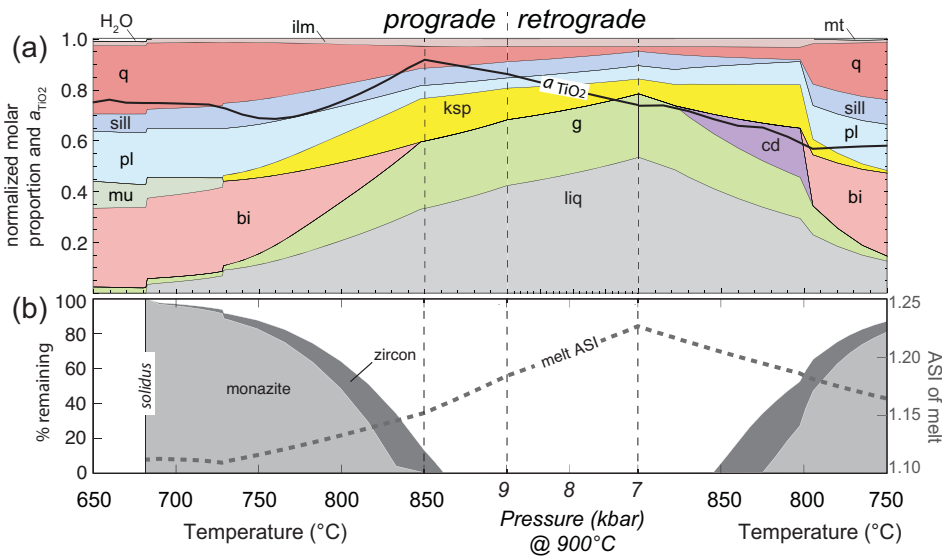
+ liq + ilm



- | | | |
|----------------------------|------------------------|--------------------|
| 1: g bi bi mu pl ru q | 8: g cd bi ksp pl q | 15: g sil ksp pl |
| 2: g ky bi mu ksp pl ru q | 9: g cd ksp pl ru q | 16: g sil pl |
| 3: g sil ksp ru q | 10: g cd ksp pl ru | 17: g sil cd pl |
| 4: g sil ksp q | 11: g cd ksp sp pl mt | 18: g sil sp pl |
| 5: sil bi mu pl q | 12: g cd sp pl mt | 19: g sil cd sp pl |
| 6: sil ksp bi mu pl q | 13: g sil cd ksp sp pl | |
| 7: g sil cd bi ksp pl mt q | 14: g sil cd ksp pl | |

Figure 7

Metapelite: clockwise P-T



Metapelite: counterclockwise P-T

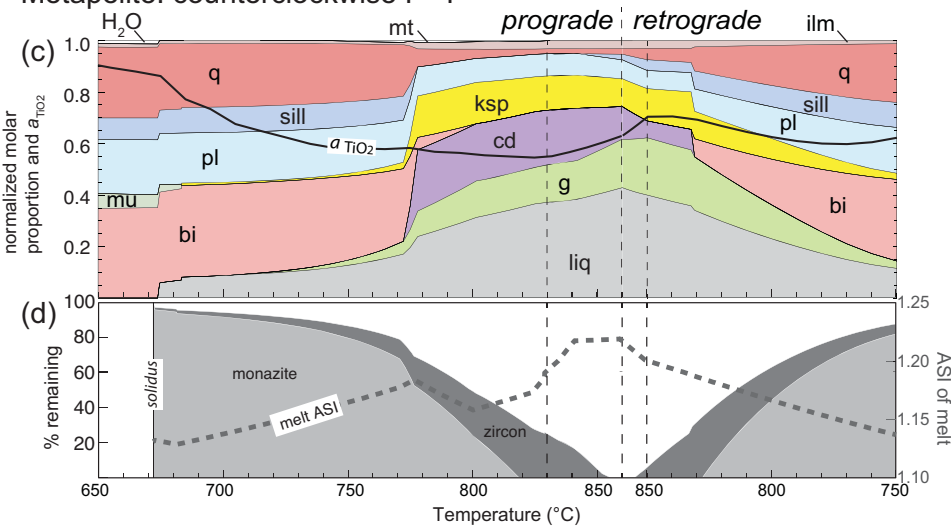


Figure 8

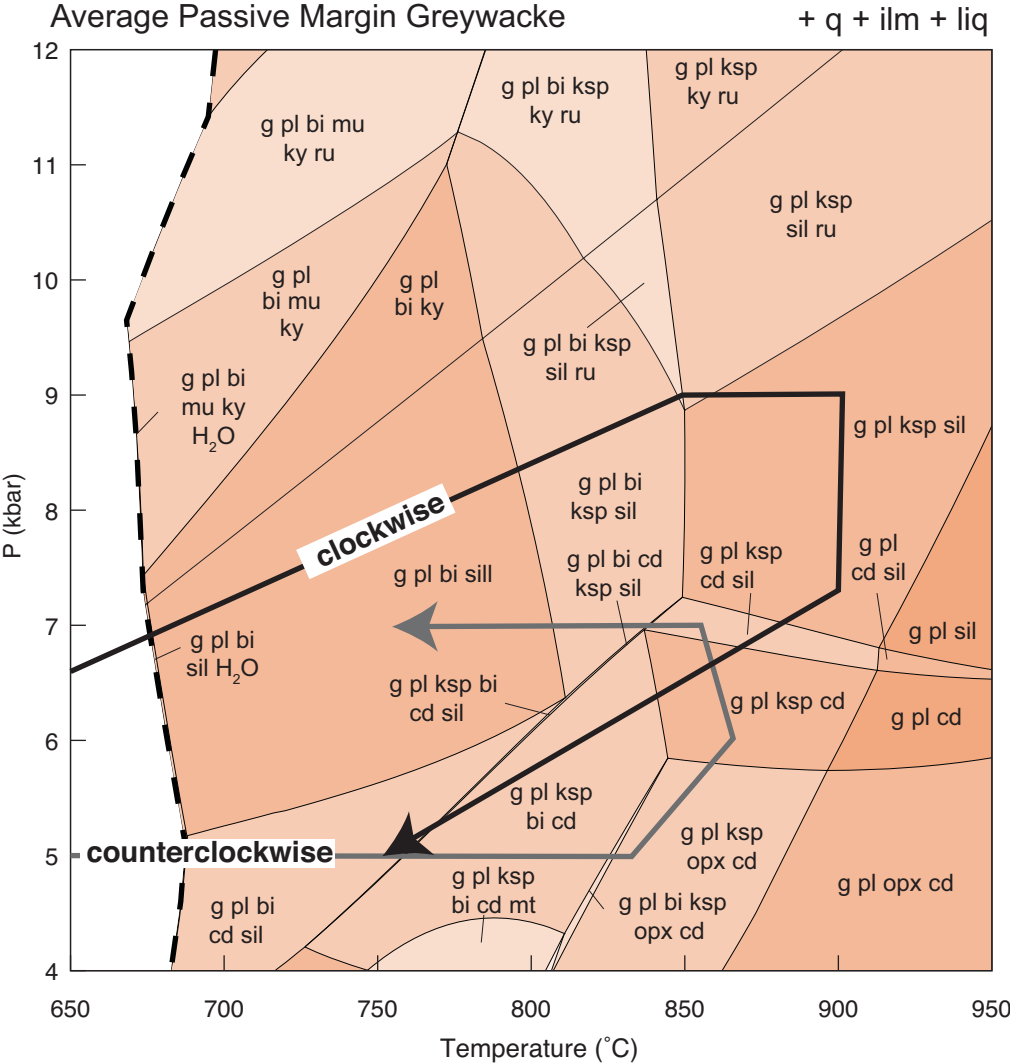
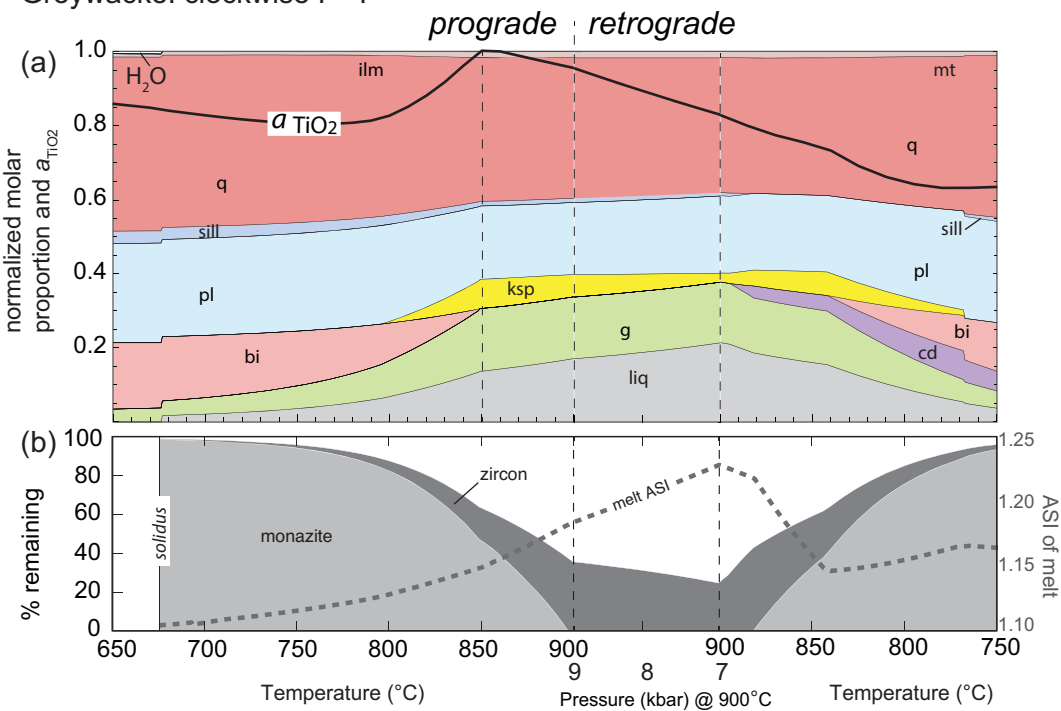


Figure 9

Greywacke: clockwise P-T



Greywacke: counterclockwise P-T

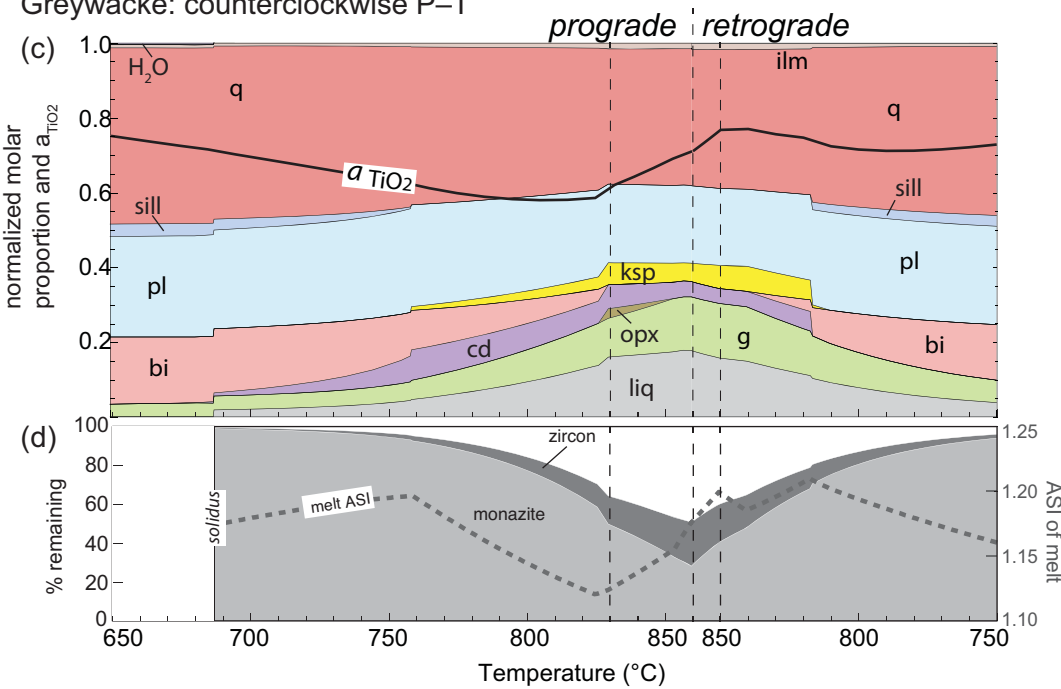


Figure 10

Average MORB

+ liq

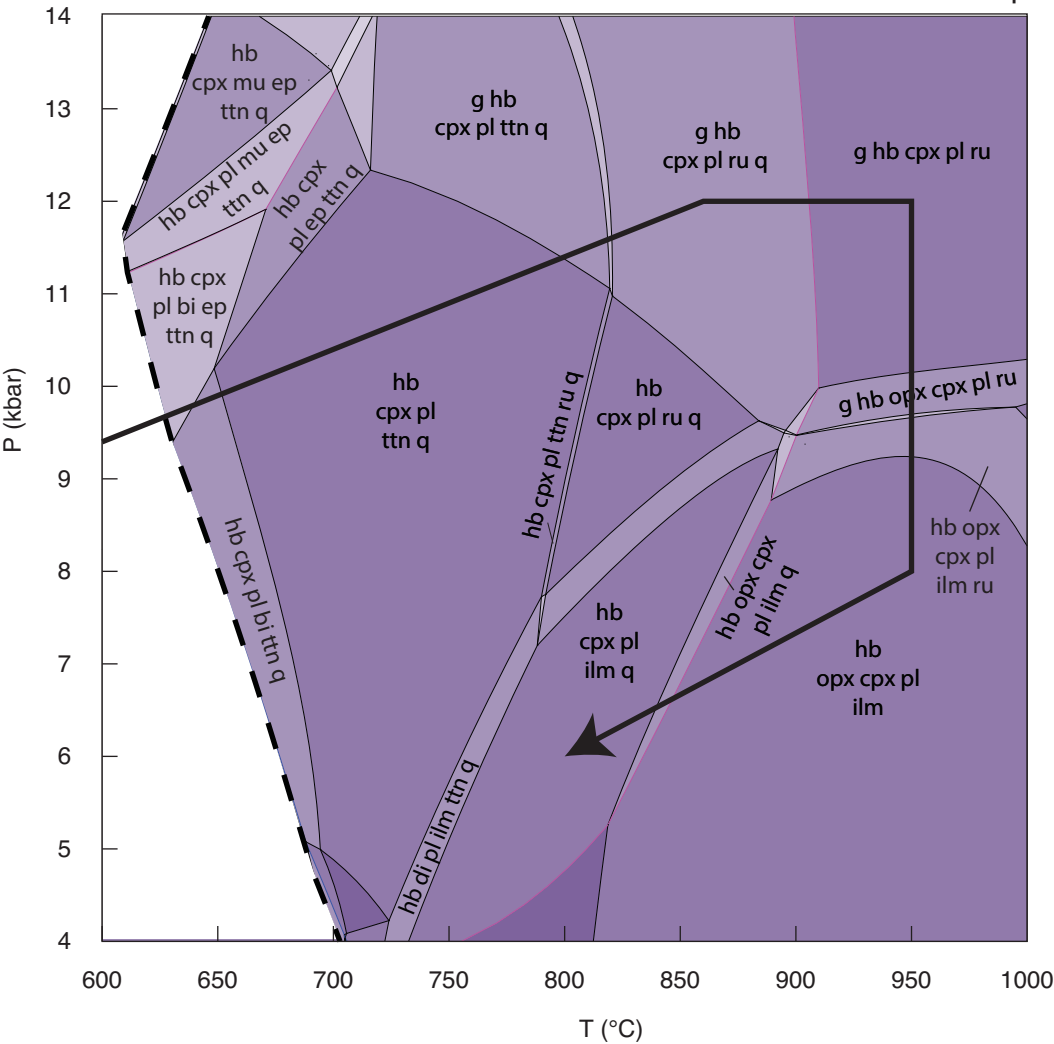


Figure 11

MORB: clockwise P-T

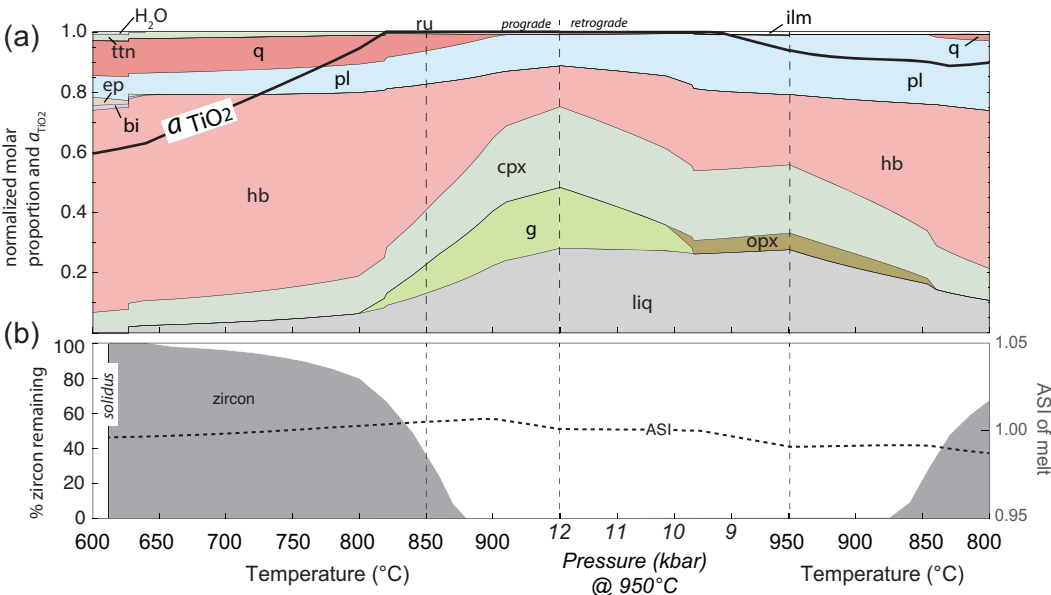


Figure 12

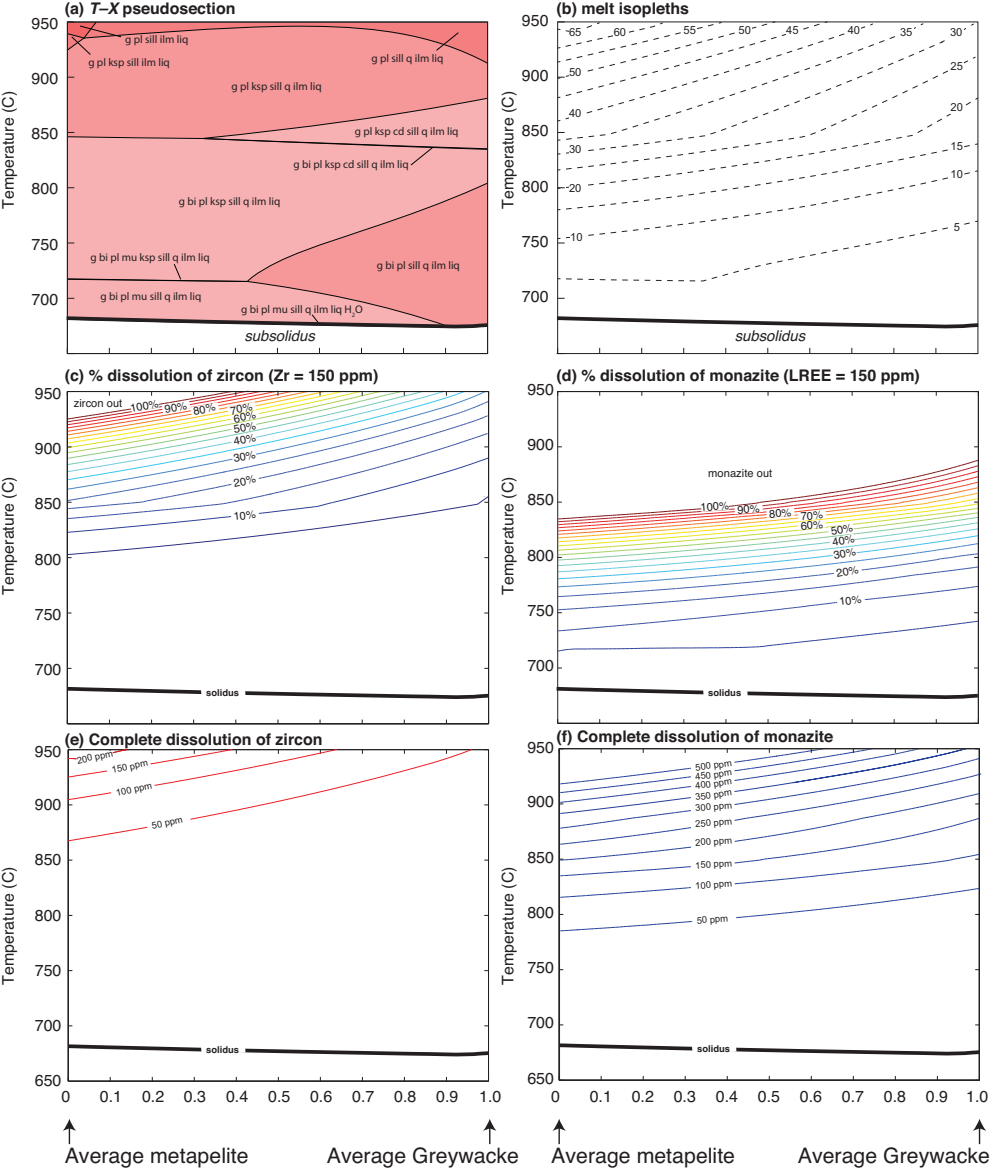


Figure 13