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Author/s:

Webb, G;Powell, R;Mclaren, S

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**Phase equilibria constraints on the melt fertility of crustal
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1 **Phase equilibria constraints on the melt fertility of**
2 **crustal source rocks: The effect of sub-solidus water**
3 **loss**

4 G. Webb, R. Powell and S. McLaren

5 School Earth Sciences, The University of Melbourne, Vic 3010, Australia

6 *Corresponding author: gwebb@student.unimelb.edu.au

7 Short title: Prograde water loss and crustal melt fertility

8 **ABSTRACT**


9 During prograde metamorphism, free H₂O generated by ongoing dehydration reactions is
10 likely to be continuously lost from a rock primarily driven by compaction. Classical melting
11 experiments cannot easily simulate this phenomena because all run products are ideally
12 conserved within the experimental charge even though significant equilibration and
13 H₂O generation may occur during heating. Phase equilibria modelling is used to model the
14 effect of subsolidus water loss on subsequent melting relationships of felsic lithologies
15 (including metapelite, metagreywacke and metatonalite) in the suprasolidus. Subsolidus
16 water loss drives the bulk composition towards the minimum saturation point within the
17 wet melting field and results in significantly reduced subsequent melt generation when
18 compared to melting experiments involving conservation of H₂O in an experimental charge
19 from some temperature well below the solidus. This effect is most obvious at *P-T*
20 conditions just above the solidus. The reduction in melt fertility due to subsolidus water
21 loss causes rheologically critical thresholds to be shifted to higher temperatures(e.g. melt
22 connectivity threshold, melt escape threshold and the solid-to-liquid transition). Subsolidus
23 water loss is also likely to impact on crystallising melts as they cross the solidus. As a
24 result, the H₂O content of granites may not reflect the H₂O contents of the magma from
25 which they were derived. Instead they record the H₂O contents in the vicinity of the
26 minimum saturation point at the *P-T* conditions at which they crossed the solidus.

27

28 keywords: prograde metamorphism, melting, water, anatexis, THERMOCALC

29 INTRODUCTION

30 Understanding crustal melt fertility is important for developing robust models for
31 migmatite formation, granite petrogenesis and the differentiation and evolution of the
32 continental crust (Brown & Fyfe, 1970; Huppert & Sparks, 1988; Clemens, 1990;
33 Thompson, 1996; Sawyer, 1998; Annen *et al.*, 2008). Hydrous felsic lithologies melt
34 incongruently at mid- to lower-crustal depths (10–60 km) at temperatures attainable during
35 high-grade metamorphism (650–1000 °C; Thompson & Connolly, 1995; Thompson, 1999;
36 White *et al.*, 2001). The amount of melt produced and its composition is determined by the
37 composition of the source rock, the pressure and temperature conditions and any open
38 system behaviour including the loss or gain of fluid, melt or solid phases. Of these,
39 H₂O content is one of the key compositional variables affecting melting behaviour in the
40 crust (Tuttle & Bowen, 1958; Holtz *et al.*, 1995; Johannes & Holtz, 1996).

41 Two dominant modes of melting have been identified to explain the observed
42 characteristics of natural granites, migmatites and granulites. ‘Fluid-present melting’ or
43 ‘wet melting’ describes the situation where a system is saturated with respect to a
44 H₂O-dominated fluid phase as it crosses the wet solidus with increasing T and/or P . In
45 fluid-present melting the activity of H₂O is effectively unity ($a_{\text{H}_2\text{O}} \approx 1$), neglecting the
46 proportions of other constituents in the fluid that are likely to be small, at least for most
47 crustal conditions. The resulting melt phase is saturated with respect to H₂O. Fluid-present
48 melting may occur in some migmatite terrains (especially at low P ; e.g. Sawyer, 2010).
49 However it is not generally considered to be the dominant mode of melting in the
50 continental crust (Clemens, 1984). By contrast, ‘fluid-absent’, ‘vapour-absent’ or
51  **‘dehydration’ melting** proceeds via reactions involving the breakdown of hydrous minerals

52 (mainly muscovite, biotite and hornblende) in the absence of a free fluid ($a_{\text{H}_2\text{O}} < 1$) at
 53 higher temperatures than the wet solidus. It has been argued that fluid-absent melting is
 54 the dominant mechanism for the formation of most granite magmas, which are generally
 55 thought to have formed at high T and are undersaturated with respect H_2O (Clemens,
 56 1984; Pickering & Johnston, 1998).

57 Estimates of the amount of melt generated during partial melting have largely come
 58 from melting experiments on a variety of starting compositions including metapelites
 59 (Vielzeuf & Holloway, 1988; Carrington & Harley, 1995; Gardien *et al.*, 1995; Pickering &
 60 Johnston, 1998), metagreywackes (Patino-Douce & Beard, 1996; Vielzeuf & Montel, 1994;
 61 Montel & Vielzeuf, 1997), tonalites (Huang & Wyllie, 1986; Rutter & Wyllie, 1988; Skjerlie
 62 & Johnston, 1993; Patino-Douce, 1997; 2004) and amphibolites (Rushmer, 1991; Wolf &
 63 Wyllie, 1994). The majority of recent experimental studies have focussed on fluid-absent
 64 melting because of its perceived importance in magma genesis and crustal differentiation.
 65 The results of these fluid-absent melting studies indicate that different crustal lithologies
 66 display markedly different melting behaviour at similar P - T conditions. **Furthermore,**
 67 **different melt fractions have been reported for compositionally-similar starting materials**
 68 **(e.g. Gardien *et al.*, 1995).**



69 **Fluid-absent melting experiments are often described as having ‘no added water’ or**
 70 **being ‘vapour-absent’. This is generally achieved by pre-heating the starting material above**
 71 **$\sim 150^\circ\text{C}$ for a sustained period (ca. 24 hours), causing any adsorbed H_2O to escape, prior to**
 72 sealing the experimental charge (e.g. Patino-Douce, 2004). It is then assumed that any
 73 H_2O retained within the starting material is structurally bound within hydrous minerals
 74 (e.g. muscovite, biotite and amphibole). However, hydrous minerals that are stable up to
 75 150°C at ambient pressures may begin to break down, via subsolidus dehydration reactions,

76 as the experimental charge is heated (and pressurised) to the desired conditions of the
 77 experimental run. Because of the intentional ‘closed system’ nature of most high P melting
 78 experiments, any H_2O generated via subsolidus dehydration will be effectively trapped
 79 within the experimental charge. In this way a ‘vapour absent’ starting material can evolve,
 80 becoming ‘fluid-present’ at the experimental P – T conditions. Additionally, London *et al.*
 81 (2012) suggested that powdered samples loaded in air may absorb between 1 and 4 weight
 82 percent H_2O before the experimental charge is sealed. It has also been suggested that
 83 water is often lost from the experimental charge via diffusion of molecular H_2O through the
 84 metal casing at temperatures above ~ 950 °C as suggested by decreasing melt modes and
 85 increasing modal proportions of solid phases (e.g. plagioclase and garnet) with increasing T
 86 during experimental runs (Patino-Douce & Beard, 1994). The combined effects of
 87 absorption during handling and diffusive loss at high temperature work in opposite
 88 directions, thereby increasing the uncertainty in the actual H_2O content inside the
 89 experimental charge during high temperature melting experiments. Furthermore, if H^+ is
 90 lost through the capsule rather than molecular H_2O , as has been suggested by some
 91 authors, this may affect the oxidation state during a melting experiment (Patino-Douce &
 92 Beard, 1994; White *et al.*, 2011).

93 An important constraint on the amount of H_2O available for melting reactions during
 94 regional metamorphism can be obtained by considering the evidence for the nature of the
 95 fluid regime prior to the onset of melting. During prograde metamorphism of rocks with an
 96 initially high H_2O content (e.g. metapelites and metagreywackes), free H_2O is likely to be
 97 continuously generated via devolatilisation reactions up to the point at which the wet
 98 solidus is crossed with increasing T and/or P . Above this point, H_2O is consumed via wet
 99 melting reactions and decreases until it is no longer stable as a free phase at progressively

100 higher T. Several authors have argued that any H₂O generated during subsolidus
101 dehydration has a strong tendency to escape the source rock due to the combined effects of
102 compaction and increasing fluid pressure (Thompson & Connolly, 1990; Connolly, 1997;
103 Yardley, 2009). Decreasing H₂O content with increasing metamorphic grade is observed in
104 most metamorphic terrains providing evidence that much of the H₂O generated during
105 subsolidus dehydration escapes the system prior to the onset of melting (Miyashiro, 1961).
106 Indeed, the loss of fluid from metamorphic terrains may be a requirement for preserving
107 metamorphic isograds during retrograde cooling and/or decompression (Thompson, 1983;
108 Guiraud *et al.*, 2001). Subsolidus H₂O loss (SWL) may be especially significant at the
109 pressure conditions of the middle to lower crust where rock porosity is thought to be small
110 (<0.1–0.5 %), and positive excursions in hydrostatic pressure above the prevailing
111 lithostatic pressure cannot be maintained for significant time periods (Norton & Knapp,
112 1977; Connolly, 1997; 2010).

113 Importantly, there are few existing datasets to constrain sub-solidus open system
114 processes involving H₂O, and experimentally determined phase relationships do not
115 adequately constrain melting relationships, particularly with respect to H₂O. Quantitative
116 thermodynamic calculations using internally-consistent datasets can be readily employed in
117 multicomponent systems that approach the compositional complexity of natural rocks (e.g.
118 White & Powell, 2002; Powell *et al.*, 2005; White *et al.*, 2005; White & Powell, 2010). *P-X*
119 or *T-X* pseudosections can be used to visualise the effect of changes in bulk composition, as
120 a result of the loss of a fluid and/or solid component from the system. This kind of
121 thermodynamic modelling has the potential to complement experimental studies to provide
122 a clearer picture of open-system processes that are thought to occur in the continental
123 crust. Here we use pseudosections to investigate the effect of SWL on melting relationships



124 for a variety of bulk compositions representing common crustal rock types (metapelites,
 125 metagreywackes, tonalites, granodiorites). The impact of SWL on melt fertility is modelled
 126 at temperatures appropriate to partial melting during regional metamorphism at mid- to
 127 lower-crustal pressures. The modelling results are compared to the results derived from
 128 melting experiments to provide improved constraints on the nature of melting during
 129 crustal anatexis.

130 CALCULATED PHASE RELATIONS

131 Mineral equilibria calculations were undertaken using THERMOCALC 3.37i (Powell &
 132 Holland, 1988; updated 2013) with an updated version of the internally consistent data set
 133 of Holland & Powell (1998; data set `tcds55`, file created 22 November 2003). The
 134 calculations were undertaken in the chemical system
 135 $\text{Na}_2\text{O}-\text{CaO}-\text{K}_2\text{O}-\text{FeO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{SiO}_2-\text{H}_2\text{O}-\text{TiO}_2-\text{O}$ (NCKFMASHTO) of White *et al.*
 136 (2007). The phases considered in the calculations, and references to the
 137 activity-composition models used are: garnet, biotite and silicate melt (White *et al.*, 2007),
 138 cordierite and epidote (Holland & Powell, 1998), orthopyroxene and magnetite (White &
 139 Powell, 2002), muscovite (Coggon & Holland, 2002), plagioclase - K-feldspar (Holland &
 140 Powell, 2003), ilmenite (White *et al.*, 2000), and hornblende (Diener *et al.*, 2007).
 141 Sillimanite, kyanite and quartz are pure end-member phases. Mineral abbreviations used in
 142 diagrams and in the text are: opx – orthopyroxene; g – garnet; cd – cordierite; hb –
 143 hornblende; bi – biotite; mu – muscovite; ky – kyanite; sill – sillimanite; ksp – K-feldspar; pl
 144 – plagioclase; ilm – ilmenite; mt – magnetite; q – quartz; liq – silicate liquid/melt.

145 Bulk compositions for phase equilibria modelling were derived from starting

146 compositions of published melting experiments for which melt fraction (Φ) estimates are
 147 available (Table 1). The four starting compositions were chosen to represent common felsic
 148 rock types and have compositions that are comparable to the average compositions of the
 149 middle and upper crust (Fig. 1). Two metapelitic bulk compositions are from Vielzeuf &
 150 Holloway (1988) and Gardien *et al.* (1995) and denoted VH88P and G95P, respectively.
 151 VH88P is modelled on a natural metapelite from the Cabo Ortegal metamorphic complex
 152 (Galicia, NW Spain) composed of quartz (39 wt%), plagioclase (19 wt%), kyanite (7 wt%),
 153 muscovite (9 wt%), biotite(21 wt%), garnet (2 wt%), staurolite (1 wt%), secondary chlorite
 154 (1 wt%) and accessory minerals (1 wt%). VH88P is strongly peraluminous and has
 155 comparatively high molar K_2O/Na_2O and high FeO contents. G95P is derived from a model
 156 two-mica metapelite generated from a mixture of quartz (39 wt%), plagioclase (30 wt%),
 157 muscovite (15 wt%) and biotite (15 wt%). It is weakly peraluminous and has comparatively
 158 low K_2O/Na_2O , high SiO_2 and low CaO. A metagreywacke bulk composition (denoted
 159 VM94G) is derived from melting experiments on a natural metagreywacke from the vallée
 160 de la Beaume (Ardèche, France) composed of quartz (40 wt%), plagioclase (32 wt%), and
 161 biotite (25 wt%), and accessory minerals (3 wt%; Vielzeuf & Montel, 1994; Montel &
 162 Vielzeuf, 1997). This composition is weakly peraluminous with high K_2O/Na_2O , SiO_2 and
 163 low FeO. A tonalitic bulk composition (denoted PD97T) was derived from melting
 164 experiments on a natural tonalite starting material from the Sierra Nevada Batholith (Lee
 165 Vining Canyon area; Patino-Douce, 1997). The sample was reported to contain of quartz
 166 (20 wt%), plagioclase (45 wt%), K-feldspar (7 wt%), hornblende (13 wt%), biotite (13 wt%)
 167 and accessory minerals (2 wt%). PD97T is metaluminous with low K_2O/Na_2O , SiO_2 and
 168 high CaO and Na_2O .

169 Some of the bulk compositions contain low concentrations of F, P, Mn and CO_2 . Small

170 amounts P_2O_5 , CO_2 and MnO are unlikely to have a measurable effect on the melting
171 behaviour of felsic rocks, whereas F contents in the range 0.01–4 wt% may have the effect of
172 lowering the solidus of granitic melts (Manning, 1981) and stabilising biotite to higher
173 temperatures (Peterson *et al.*, 1991). Fluorine is not currently incorporated into the
174 activity-composition models for biotite or melt. As a result, the models presented here may
175 differ slightly in terms of melting behaviour from the more complex starting compositions of
176 the corresponding experiments. Furthermore, it should be noted that the melt model used
177 here does not incorporate TiO_2 or ferric iron. TiO_2 is readily incorporated into biotite and
178 oxide minerals (e.g. ilmenite and magnetite) and is important for stabilising biotite at high
179 temperature (White *et al.*, 2007). Hayden & Watson (2007) investigated the incorporation
180 of TiO_2 into haplogranitic melt. Their results indicated that even for rutile-saturated bulk
181 compositions the amount of TiO_2 incorporated into the melt phase was small and as such,
182 is unlikely to have a first order effect on melt fertility. The ferric iron content of all of the
183 bulk compositions in this study are fixed at low values such that the only stable iron oxide
184 minerals are ilmenite \pm magnetite.

185 For each starting composition the H_2O content was varied to produce a low- H_2O and a
186 high- H_2O end-member (Table 2). The high and low values were chosen to encompass the
187 transition from H_2O undersaturated to H_2O saturated conditions at the wet solidus. Each
188 end-member pair were then used to construct T - M_{H_2O} pseudosections (Guiraud *et al.*,
189 2001) for each model starting composition. Diagrams were constructed at 10 kbar for the
190 starting compositions VH88P, G95P and VM94G and at 8 kbar for composition PD97T to
191 facilitate comparison with the reported experimental results. Models were also generated at
192 5 kbar for VH88P, G95P and VM94G and at 4 kbar for PD97T to investigate melting
193 equilibria at low pressure.

194 **Pseudosections**

195 T - $M_{\text{H}_2\text{O}}$ pseudosections calculated for compositions VH88P, G95P, VM94G at 10 kbar, and
 196 PD97T at 8 kbar over the temperature range 450 to 1100 °C are shown in Fig. 2. The
 197 pseudosections for all four compositions have a sub horizontal H_2O -saturated solidus
 198 (wet-solidus). VH88P has the highest T wet solidus at ~ 680 °C. The wet solidi for G95P,
 199 VM94G and PD97T are all similar, occurring at ~ 645 °C. In each case the wet solidus
 200 extends beyond the range of the high H_2O end-member but is limited at low H_2O contents
 201 by the stability of free H_2O . The point at which the solidus is minimally saturated with
 202 respect to H_2O is referred to here as the minimum saturation point (MSP; Fig. 2). The
 203 narrow fields immediately above the wet-solidus in all four T - $M_{\text{H}_2\text{O}}$ pseudosections are low
 204 variance fields related to the underlying NCKASH univariant reaction,



205 The H_2O -out boundaries of these ‘wet-melting’ fields are intersected within 10–40 °C
 206 above the wet-solidus. The upper boundaries increase in T only slightly as they trend from
 207 the minimum saturation point towards the high H_2O end-member for each diagram.

208 At H_2O contents less than the value of the minimum saturation point the temperature
 209 of the solidus increases steeply as the equilibrium composition becomes increasingly
 210 undersaturated with respect to H_2O . Inflections in the solidus curves at low H_2O contents
 211 on each of the T - $M_{\text{H}_2\text{O}}$ pseudosections correspond to changes in the modal mineralogy of
 212 low variance fields emanating from ‘vapour absent’ univariant reactions in the underlying
 213 NCKASH and NCKFMASH systems including,



214 For all compositions, the subsolidus H₂O-out boundary trends to lower bulk
 215 H₂O contents as it approaches the solidus (at the minimum saturation point) with
 216 increasing T (Fig. 2). The H₂O-out boundary for VH88P rises steeply from <450 °C up to
 217 the plagioclase-in boundary (~615 °C) and then migrates towards a lower bulk H₂O value as
 218 it approaches the solidus. For compositions G95P and VM94G the H₂O-out boundary is
 219 steep and approximately linear from <450 °C up to the wet-solidus. The H₂O-out boundary
 220 for PD97T trends gradually towards lower bulk H₂O, with distinct inflections corresponding
 221 to muscovite-out, K-feldspar-in and hornblende-in boundaries, as it approaches the
 222 minimum saturation point at the wet solidus with increasing temperature.

223 The pseudosections for VH88P and G95P both contain sillimanite and kyanite. For
 224 VH88P, sillimanite is stable from the sill-ky phase transition (~800 °C) up to >1000 °C at
 225 high H₂O contents and is stable above 1100 °C at low H₂O contents (Fig. 2 a & b). Kyanite
 226 is stable from 800 °C down to below 450 °C over the entire range of modelled H₂O contents
 227 (0.11–14.02 mol%). Sillimanite is stable for G95P from the sill-ky phase transition to above
 228 1100 °C near the low-H₂O end-member, but its upper limit of stability decreases to ~900 °C
 229 at the high-H₂O end-member. Kyanite is stable from 800 °C down to 450 °C at
 230 H₂O contents below ~3 mol%. Above this value the stability is limited by the kyanite-out
 231 boundary along the low T sides of the quadrivariant bi-mu-ksp-pl-liq-ilm-q-ky and the
 232 quinivariant bi-mu-pl-liq-ilm-q-ky fields. Neither sillimanite nor kyanite are stable for

233 VM94G and PD97T over the range modelled (Fig. 2 c & d).

234 K-feldspar is unstable at temperatures above the solidus at high bulk H₂O contents for
 235 all four modelled compositions. At bulk H₂O contents close to the minimum saturation
 236 point, K-feldspar is stable over the ranges ~790–1000 °C, ~700–980 °C, ~810–990 °C and
 237 <450–990 °C for compositions VH88P, G95P, VM94G and PD97T, respectively.

238 It is important to note that even at high T (>1000 °C) none of the bulk compositions
 239 reach their liquidus and form pure melts (including the metatonalite - PD97T; Fig. 2).
 240 Even for the high H₂O end-members the high T equilibrium assemblages comprise liq-g-ilm,
 241 liq-opx-ilm, liq-opx-ilm-q and liq-hb-pl-ilm for VH88P, G95P, VM94G and PD97T,
 242 respectively. Lower variance high T assemblages become increasingly stable at lower bulk
 243 H₂O contents in each pseudosection.

244 Ilmenite is stable for all for starting compositions over the entire modelled ranges of
 245 temperature and H₂O contents. The pseudosection for composition G95P contains rutile at
 246 low temperature and magnetite is stable for intermediate H₂O contents between
 247 800–880 °C. Staurolite is stable from just below to just above the solidus for composition
 248 VM94G. While these phases are involved in the NCKFMASHTO equilibria, their modal
 249 abundances (1-oxide molar basis) are always low (<0.01) and are, therefore, unlikely to
 250 exert a significant influence on the melting behaviour of the system modelled.

251 **Prograde metamorphism and sub-solidus water loss**

252 The contours of melt and free H₂O corresponding to the pseudosections in Fig. 2 are shown
 253 in Figure 3. While each T - $M_{\text{H}_2\text{O}}$ pseudosection displays distinct phase relationships in
 254 detail, the relationships between the free H₂O and melt bearing regions in each case are

255 broadly similar. The proportion of free H₂O present along a particular prograde heating
256 path in the subsolidus (vertical line) can be described in terms of df/dT (representing the
257 change in free H₂O productivity with increasing T). Similarly, the amount of melt produced
258 along a path in the suprasolidus can be described in terms of dm/dT (i.e. change in melt
259 productivity with increasing T ; Johnson *et al.*, 2008).

260 At sub-solidus conditions df/dT is positive from $<450^\circ\text{C}$ up to the solidus at bulk
261 H₂O values above the H₂O-out boundary for each of the four compositions (grey regions in
262 Fig. 3). As a consequence, all four modelled compositions will produce free H₂O along a
263 prograde path, with fixed H₂O contents, that traverses the ‘free H₂O-stable’ region (i.e.
264 bulk H₂O contents greater than the value of the minimum saturation point). At
265 supra-solidus conditions, H₂O & melt are both stable within the narrow wet-melting fields.
266 Within this region, df/dT becomes negative and the amount of free H₂O decreases rapidly
267 towards the H₂O-out boundary with increasing T . This region also corresponds to an initial
268 increase in melt production with melt contours that are perpendicular to the T -axis
269 ($dm/dT > 10 \text{ mol\%/}10^\circ\text{C}$). Above the H₂O+melt region the melt contours have a slope
270 influenced by the shape of the H₂O-undersaturated solidus such that increases in T and/or
271 bulk H₂O correspond to increased melt production.

272 An idealised melting experiment involves conserving all of the starting components
273 (including the initial H₂O contents) throughout the experimental run. This behaviour is
274 represented by the conservation of subsolidus water (CSW) scenario, shown by the vertical
275 grey arrows in Fig. 3. The water contents for the CSW scenario for VH88P, G95P and
276 VM94G are derived from the reported water contents for the bulk starting material of the
277 original experiments (2.15 w%, 1.9 w% and 1.44 w%, respectively; Table 2). The
278 H₂O contents of PD97T was not reported. The CSW path for this composition was chosen

279 so as to intersect the observed melt fraction in the experiments (i.e. $\Phi = 0.30$ melting at
280 950°C ; Patino-Douce, 1997).

281 If subsolidus water loss occurs, free H_2O will be incrementally lost along the prograde
282 heating path (Guiraud *et al.*, 2001), continuously driving the bulk composition towards the
283 H_2O -out boundary. To reduce the bulk H_2O contents of the source rock below the value of
284 the H_2O -out boundary would require the liberation of structurally bound H_2O from hydrous
285 minerals. As there is no viable mechanism to achieve this, the subsolidus H_2O -out
286 boundary effectively limits the minimum H_2O -contents of the source rock. As the rock
287 crosses the solidus with increasing temperature any remaining free H_2O is rapidly consumed
288 within the wet-melting fields and the newly generated melt becomes increasingly
289 undersaturated with respect to H_2O . Because such melts are effectively ‘water-starved’ as
290 soon as they cross the solidus, the H_2O contents of the system become fixed at the value of
291 the minimum saturation point, if H_2O loss is efficient and the porosity is small, as generally
292 suggested. As a result an isobaric prograde heating path for a rock composition affected by
293 SWL will involve:

- 294 1. A sub-solidus path that traverses the free H_2O region at bulk H_2O contents just
295 greater than the H_2O -out boundary as it approaches the minimum saturation point;
- 296 2. A supra-solidus path with the bulk H_2O fixed at the value of the minimum saturation
297 point up to the maximum temperature of the system.

298 In natural systems, H_2O may additionally be lost via incorporation into escaping melt
299 or as part of hydrous peritectic minerals that are themselves entrained into escaping melt
300 (Clemens & Stevens, 2012). However, both of these mechanisms will lead to other changes
301 in the bulk composition that cannot be determined by varying any single compositional

302 variable. The appropriate diagrams for modelling such phenomena are more complex and
303 involve varying the equilibrium bulk composition at each increment of melt- and/or
304 mineral-loss.

305 The mineral mode changes along the subsolidus sections of the SWL paths were
306 calculated along the H₂O-out boundary so that the bulk H₂O contents continuously changes
307 along this segment. The supra-solidus portion of the SWL path was modelled at a fixed
308 H₂O value (corresponding to the minimum saturation point; Table 2). Changes in the
309 modal abundances of the equilibrium assemblages along the modelled CSW and SWL paths
310 for the four starting compositions are shown in Fig. 4.

311 For composition VH88P the SWL heating path results in significantly less melt
312 generation when compared with the CSW path (especially at low T; Fig. 4a & 4b). The
313 reduction in melt fraction along the SWL path corresponds to increases in the modal
314 proportions of quartz, plagioclase and biotite. K-feldspar is stabilised at temperatures
315 above ~790 °C, whereas K-feldspar is completely absent in the CSW scenario. Very high
316 dm/dT melting (20 mol%/10 °C) along the CSW path ($\Phi < 0.01$ to ~ 0.11) corresponds to
317 the H₂O-out boundary just above the solidus, at ~680 °C. This feature is not observed
318 along the SWL path. Moderate dm/dT melting (~3 mol%/10 °C) over the interval
319 790–880 °C on the SWL path corresponds to breakdown of biotite and increased growth of
320 garnet. Melting at higher temperatures continues to increase in a broadly monotonic
321 fashion reaching melt fractions of ~0.65 and ~0.44 at 1000 °C for the CSW and SWL paths,
322 respectively ($dm/dT \sim 1\text{--}2$ mol%/10 °C). Interestingly, muscovite begins to breakdown
323 below the solidus (at ~615 °C, corresponding to the first appearance of plagioclase) and
324 completely disappears just above the solidus, at ~720 °C, for both the SWL and CSW
325 paths; however, there is no significant increase in the melt fraction corresponding to the

326 disappearance of muscovite from the equilibrium assemblage along either path.

327 The SWL path for G95P corresponds to reduced melt fertility and slight increases in
 328 the modal proportions of quartz and plagioclase at temperatures just above the solidus
 329 ($\sim 640^\circ\text{C}$) and an increase in the proportion of K-feldspar at temperatures above 760°C
 330 when compared to the CSW path (Fig. 4c & 4d). Two steep increases in melting occur
 331 along the CSW path at low melt fraction. The first corresponds to the H_2O -out boundary
 332 just above the solidus ($dm/dT \sim 6\text{--}8 \text{ mol}\%/10^\circ\text{C}$). The second step corresponds to the
 333 abrupt disappearance of muscovite and the first appearance of kyanite and K-feldspar (at
 334 $\sim 760^\circ\text{C}$; $dm/dT \sim 10\text{--}20 \text{ mol}\%/10^\circ\text{C}$). Due to the negligible amount of H_2O at the solidus
 335 in the SWL scenario the melt fraction remains low ($\Phi < 0.02$) until muscovite disappears at
 336 $\sim 760^\circ\text{C}$ at which point it increases abruptly to ($\Phi \sim 0.09$). Melting at higher temperatures
 337 continues to increase steeply with melt fractions reaching ~ 0.69 and ~ 0.51 at 1000°C for
 338 the CSW and SWL paths, respectively ($dm/dT \sim 3\text{--}5 \text{ mol}\%/10^\circ\text{C}$).

339 For composition VM94G, the SWL scenario results in significantly reduced melt
 340 fertility, moderate increases in the modal proportions of quartz and plagioclase at
 341 temperatures just above the solidus (Fig. 4e & 4f). K-feldspar increases in modal
 342 abundance and becomes stabilised over a broader temperature range ($\sim 805\text{--}1000^\circ\text{C}$) when
 343 compared to the CSW path ($\sim 905\text{--}930^\circ\text{C}$). A sharp increase in melting along the CSW
 344 path corresponds to the H_2O -out boundary just above the solidus (at $\sim 650^\circ\text{C}$; dm/dT
 345 $\sim 8\text{--}10 \text{ mol}\%/10^\circ\text{C}$). Again, this feature is absent from the SWL path. Immediately above
 346 the solidus the melt fraction remains low ($\Phi < 0.01$) along the SWL path with an initial,
 347 gradual increase in melt fraction, beginning at $\sim 755^\circ\text{C}$, corresponding to the first
 348 appearance of garnet and the initial breakdown of biotite (dm/dT increasing from < 0.1 up
 349 to $2 \text{ mol}\%/10^\circ\text{C}$). A steep increase in melting, from $\Phi \approx 0.09$ to $\Phi \approx 0.19$, corresponds to

350 the final disappearance of biotite and the first appearance of orthopyroxene across the
351 temperature interval $\sim 895\text{--}910\text{ }^{\circ}\text{C}$ for the SWL path ($dm/dT \sim 7\text{ mol\%/10 }^{\circ}\text{C}$). A similar
352 step occurs along the CSW path over the same temperature interval with melting abruptly
353 increasing from from $\Phi \approx 0.28$ to $\Phi \approx 0.36$. Above the biotite-out boundary, melt fractions
354 climb steeply to values of ~ 0.56 and ~ 0.34 at $1000\text{ }^{\circ}\text{C}$ for the CSW and SWL paths,
355 respectively ($dm/dT \sim 3\text{--}4\text{ mol\%/10 }^{\circ}\text{C}$).

356 The SWL scenario for composition PD97T results in decreased melt fertility and slight
357 increases in the modal proportions of quartz, plagioclase and K-feldspar when compared to
358 the CSW path (Fig. 4g & 4h). A small, moderate increase in melt fraction, from 0.0 to
359 ~ 0.02 , occurs just above the solidus, at $\sim 640\text{ }^{\circ}\text{C}$, along the CSW path (dm/dT
360 $\sim 5\text{ mol\%/10 }^{\circ}\text{C}$). Again, this initial step in melt fraction is absent from the SWL path. The
361 melt fraction along the CSW path increases gently to 0.09 over the interval $\sim 645\text{--}840\text{ }^{\circ}\text{C}$
362 ($dm/dT \sim 0.1\text{--}0.2\text{ mol\%/10 }^{\circ}\text{C}$) and then abruptly increases to 0.17 between $\sim 840\text{--}850\text{ }^{\circ}\text{C}$
363 ($dm/dT \sim 10\text{ mol\%/10 }^{\circ}\text{C}$). A similar sharp increase in melt fraction occurs along the SWL
364 path (from $\Phi \sim 0.03$ to 0.10) over the same temperature interval. For both paths this step
365 corresponds to the first appearance of orthopyroxene, the breakdown and disappearance of
366 biotite and an increase in the modal abundance of K-feldspar. Above the biotite-out
367 boundary melt fractions increase moderately steeply to reach ~ 0.45 and ~ 0.36 at $1000\text{ }^{\circ}\text{C}$
368 for the CSW and SWL paths, respectively ($dm/dT \sim 10\text{ mol\%/10 }^{\circ}\text{C}$). Interestingly,
369 hornblende modes do not noticeably decrease for either path over the calculated melting
370 interval ($\sim 640\text{--}1100\text{ }^{\circ}\text{C}$).

371 Melting at low pressure

372 The minimum saturation point for each composition, and the corresponding SWL melting
 373 path, increases as a function of pressure. Therefore, it is not possible to create a single P - T
 374 pseudosection for a bulk composition affected by SWL. Rather it is necessary to determine
 375 T - $M_{\text{H}_2\text{O}}$ phase relationships and H_2O value of the minimum saturation point at each
 376 pressure of interest. As a result this method is only applicable to the consideration of
 377 isobaric melting paths. To examine the effect of pressure on the phase equilibria,
 378 temperature-mode diagrams were generated for the SWL scenario along prograde isobaric
 379 heating paths for VH88P, G95P and VM94G at 5 kbar and PD97T at 4 kbar (Fig. 5).

380 At 5 kbar, VH88P (SWL) produces very little melt ($\Phi < 0.03$) between the solidus, at
 381 $\sim 690^\circ\text{C}$, and $\sim 775^\circ\text{C}$ ($dm/dT \ll 0.1 \text{ mol\%/}10^\circ\text{C}$). The melt fraction then rises sharply to
 382 $\Phi \approx 0.07$ at $\sim 780^\circ\text{C}$ ($dm/dT \sim 10\text{--}20 \text{ mol\%/}10^\circ\text{C}$). This step corresponds to the
 383 disappearance of sillimanite from the equilibrium assemblage, a sharp decrease in the modal
 384 abundance of biotite, an increase in the modal abundance of cordierite and the first
 385 appearance of garnet. This feature is absent from the SWL scenario for composition VH88P
 386 at 10 kbar. Above $\sim 780^\circ\text{C}$ the melt fraction increases moderately steeply reaching Φ
 387 ≈ 0.67 at $\sim 1000^\circ\text{C}$ ($dm/dT \sim 2\text{--}3 \text{ mol\%/}10^\circ\text{C}$; *c.f.* 0.44 for VH88P at 10 kbar).

388 The melt fraction for composition G95P (SWL) at 5 kbar remains low (< 0.01) between
 389 the solidus ($\sim 655^\circ\text{C}$) and $\sim 675^\circ\text{C}$ ($dm/dT \ll 0.1 \text{ mol\%/}10^\circ\text{C}$). Between $675\text{--}680^\circ\text{C}$ the
 390 melt fraction jumps abruptly to ~ 0.07 ($dm/dT \sim 10\text{--}20 \text{ mol\%/}10^\circ\text{C}$). This step
 391 corresponds to the disappearance of muscovite and the first appearance of sillimanite in the
 392 equilibrium assemblage. A similar high dm/dT step is observed at $\sim 760^\circ\text{C}$ in the SWL
 393 melting curve for G95P at 10 kbar. From $680\text{--}835^\circ\text{C}$ the melt fraction increases gradually

394 to 0.19 ($dm/dT \sim 0.5\text{--}0.7 \text{ mol\%/10 }^\circ\text{C}$). From 835 °C the melting curve begins to increase
395 steeply corresponding to the breakdown and disappearance of biotite and the first
396 appearance of orthopyroxene in the equilibrium assemblage ($dm/dT \sim 8 \text{ mol\%/10 }^\circ\text{C}$).
397 Melting then proceeds with a steep slope reaching 0.70 at ~ 1000 °C (dm/dT
398 $\sim 4\text{--}5 \text{ mol\%/10 }^\circ\text{C}$; *c.f.* 0.51 for G95P at 10 kbar).

399 At 5 kbar, the melting curve for composition VM94G (SWL) increases slightly, from
400 $\Phi = 0.0$ to 0.02, between the solidus (675 °C) and the first appearance of garnet in the
401 equilibrium assemblage at 785 °C ($dm/dT \ll 0.1 \text{ mol\%/10 }^\circ\text{C}$). Between 785–840 °C the melt
402 fraction increases sharply to 0.22 corresponding to the first appearance of orthopyroxene
403 and the breakdown and disappearance of biotite from the equilibrium assemblage (dm/dT
404 $\sim 8\text{--}10 \text{ mol\%/10 }^\circ\text{C}$). Above 840 °C the melting curve rises steeply to reach 0.60 at ~ 1000 °C
405 ($dm/dT \sim 4 \text{ mol\%/10 }^\circ\text{C}$; *c.f.* 0.34 for G95P at 10 kbar).

406 At 4 kbar PC97T (SWL) produces very little melt (< 0.02) between the solidus, at
407 ~ 655 °C, and 795 °C corresponding to the first appearance of orthopyroxene (dm/dT
408 $\ll 0.1 \text{ mol\%/10 }^\circ\text{C}$). Between 795–805 °C the melt fraction increases sharply to 0.11 (dm/dT
409 $\sim 10\text{--}15 \text{ mol\%/10 }^\circ\text{C}$). This interval corresponds to the rapid breakdown and disappearance
410 of biotite and an increase in the abundance of K-feldspar and orthopyroxene in the
411 equilibrium assemblage. Above 805 °C, the melt fraction increases moderately steeply to
412 reach 0.47 at ~ 1000 °C (*c.f.* 0.36 for G95P at 10 kbar).

413 CRUSTAL MELT FERTILITY

414 Comparisons with melting experiments

415 The experimentally derived melt fractions corresponding to VH88P, G95P, VM94G and
 416 PD97T are shown in Figs. 3 & 6. As previously noted, the starting H₂O contents for the
 417 melting experiments reported by Patino-Douce (1997) are not known, so it is not possible to
 418 make a comparison with the modelled CSW melting path for PD97T. Nevertheless, it is still
 419 useful to compare an inferred CSW path from matching the reported experimental melt
 420 fraction with the modelled SWL scenario.

421 Vielzeuf & Holloway (1988) infer a very steep increase in melt fraction, from $\Phi = 0.10$
 422 to 0.60 over the temperature range 850–900 °C corresponding to a biotite-out “reaction”
 423 (i.e. $bi + sill/ky + pl + q \rightarrow liq + g + ksp$; Fig. 6a). Above 900 °C, these authors describe a
 424 melting ‘plateau’ corresponding to the gradual breakdown of sillimanite and garnet. Neither
 425 of these features are observed for the CSW path for VH88P which displays broadly
 426 monotonic melting behaviour above 800 °C. At temperatures higher than ~950 °C, they
 427 report lower melt fractions than the VH88P() melting curve. A similar misfit is observed
 428 between the the experimental results of Vielzeuf & Montel (1994) and the modelled CSW
 429 path for VM94 at 1000 °C (Fig. 6c). Experiments above 950 °C were not reported in
 430 Gardien *et al.* (1995) or Patino-Douce (1997), so any high T effect cannot be assessed for
 431 these studies. Nevertheless, the generation of lower than predicted melt fractions at high T
 432 is consistent with observations from other experiments that melt modes often decrease at
 433 temperatures above 950 °C (possibly due to diffusive H₂O-loss through the wall of the
 434 experimental capsule; Patino-Douce & Beard, 1994).

435 The experimental melt fraction estimates reported by Gardien *et al.* (1995) show a

436 general trend of increasing melting with increasing T that broadly corresponds to the CSW
437 melting curve for composition G95P (Fig. 6b). Interestingly, both the CSW and SWL
438 melting curves for composition G95P record a steep increase in melt fraction at $\sim 760^\circ\text{C}$,
439 corresponding to the mu-out boundary with increasing T , while the reported experimental
440 melt fraction actually decreases by 5% between 750°C and 800°C before increasing at
441 higher T . The melting estimates of Vielzeuf & Montel (1994) are slightly above the
442 corresponding VM94G() melting curve at temperatures below $\sim 900^\circ\text{C}$ (Fig. 6b).

443 Importantly, the experimentally-derived melt fractions for all starting compositions are
444 higher than the corresponding SWL melting curves as a function of T . This general result is
445 consistent with the assumption that SWL constrained melting represents a minimum
446 melting end-member for a particular bulk composition.


447 The general advantages and limitations of melting experiments and phase equilibria
448 modelling have been discussed in detail elsewhere (e.g. Johnson *et al.*, 2008; White *et al.*,
449 2011), so the following discussion focusses on the issues that are specific to estimating
450 crustal melt fertility. The differences between the experimental estimates and the modelled
451 CSW melting curves can be understood in terms of the different limitations inherent for
452 each approach.

453 A key assumption of most melting experiments is that the analysed composition of the
454 starting material is representative of the equilibrating bulk composition at experimental run
455 conditions. As previously discussed above, the combined effects of H_2O absorption into
456 powders during handling as well as diffusive H_2O loss through the experimental casing at
457 high temperature may result in significant changes in the actual H_2O content of a given
458 melting experiment. The CSW scenarios reported here represent idealised melting

459 experiments that are unaffected by these undesirable experimental phenomena.

460 Detailed melting experiments are labour and time intensive with individual experiments
461 often taking several weeks to approach equilibration (e.g. Patino-Douce, 2004). As a result,
462 the total number of data points generated is usually low and performing repeat experiments
463 at the same P - T conditions is uncommon making it difficult to assess the statistical
464 significance or the reproducibility of any single melting estimate. By contrast, phase
465 equilibria modelling allows the determination of melt modes over a range of P - T - X
466 conditions for which the constituent phase models are calibrated. This allows for the
467 calculation of continuous melting curves along a particular P - T path of interest.

468 Melt fractions for each experimental charge are typically estimated either by mass
469 balance calculations or by graphical assessment of the proportion of melt in
470 two-dimensional back-scattered electron scans. Both methods have high analytical
471 uncertainties that may be difficult to quantify (e.g. Vielzeuf & Montel, 1994). Quantitative
472 determination of melt fractions from experimental data is particularly problematic at low
473 melt fractions due to the small size of melt/glass volumes that may be observed and
474 analysed within the matrix of the experimental charge. As a result, experimental studies
475 can often only approximate the degree of melting occurring at temperatures near the
476 solidus. This constraint does not apply to phase equilibria modelling where the degree of
477 melting can be calculated in detail from the solidus up to high temperatures.

478 The NCKFMASH melt model used in this study is constrained by well calibrated
479 experimental data on unary, binary and ternary subsystems (Holland & Powell, 2001). It
480 can, therefore,  expected to satisfactorily describe the topology of the melt bearing fields in
481 NCKFMASHTO, if not, for example, the precise PT position of fields. As previously


482 discussed, the effects of TiO_2 and Fe^{3+} on the modelled melt fertility are likely to be small
483 and certainly less than the uncertainties inherent in the determination of experimental
484 estimates on the corresponding natural rock compositions. The influence of other minor
485 components (e.g. Mn, F, Cl, P, B) on melting behaviour cannot be assessed with the
486 models used here and in any case it is unlikely that any of these components (with the
487 possible exception of F) will significantly affect melting behaviour at the low concentrations
488 reported for the experimental starting compositions (shown in Table 1). We, therefore,
489 argue that the modelled melting curves presented here represent reliable estimates of the
490 melt fertility of the starting compositions under conditions of CSW and SWL.

491 **Estimates of crustal melt fertility**

492 It is clear that the overall ‘shapes’ of the melting curves for the CSW and SWL scenarios
493 for each starting composition are similar and reflect similar underlying melt-producing
494 equilibria (Fig. 6). Inflections in the melting curves occur at similar T indicating that they
495 are largely independent of bulk H_2O content over the ranges modelled. The key difference
496 between the CSW and SWL curves, in each case, is that the CSW melting curve contains
497 an initial steep increase, just above the solidus, corresponding to the disappearance of free
498 H_2O from the equilibrium assemblage. This initial melting step effectively increases the
499 baseline for subsequent melting but does not significantly affect the shape of the curve at
500 higher T . The most obvious observation that can be made from comparing these curves is
501 that CSW paths that incorporate excess free H_2O will yield more melt than the
502 corresponding SWL path.

503 The SWL melting curves for each starting composition at high and low pressure are

504 shown in Figure 7. With decreasing pressure the solidus migrates to higher T (shown by
 505 the grey and black arrows in Fig. 7), while at the same time the H_2O solubility of the melt
 506 phase increases. At low pressure, SWL constrained heating paths intersect key melting
 507 equilibria at lower T compared to the corresponding high P paths (cf. SWL versus CSW
 508 paths at constant P ; Fig. 6). This is because the underlying fluid-absent melting reactions
 509 have a positive slope in P - T space. The resulting melting curves show increased melt
 510 generation in the lower pressure scenarios for all starting compositions. This is likely to be a
 511 general result that is applicable to most felsic rock compositions.



512 As noted above there is **no obvious geological mechanism that would cause the bulk**
 513 **composition to contain less H_2O than the value corresponding to the minimum saturation**
 514 **point.** As a result the SWL path results in the minimum melt fertility scenario for a given
 515 source rock. Increasing melt fertility above the minimum defined by efficient SWL may be
 516 achieved via the retention of sub-solidus H_2O or the addition of H_2O to the system at
 517 temperatures above the solidus (compare paths A, B & C in Fig. 8).

518 H_2O may be released from a source rock with a higher T solidus into an adjacent rock
 519 with a lower T solidus (White *et al.*, 2005). For example, at 10 kbar composition VH88P
 520 intersects its wet-solidus at $\sim 680^\circ\text{C}$ whereas the solidus for composition G95P occurs at
 521 $\sim 640^\circ\text{C}$. If these two compositions represent hypothetical adjacent source regions subjected
 522 to the same heating path, VH88P could provide up to 3% H_2O to G95P over the
 523 temperature interval between their respective solidi. assuming the source rocks were in
 524 equal proportions, this would result in a significant increase in the melt fertility of the G95P
 525 source rock ($\sim 10\%$ more melt at 800°C , $\sim 15\%$ more melt at 1000°C). Water may also be
 526 added to a particular source rock via dewatering of migrating melts. During cooling and
 527 crystallisation water is released as the melt composition intersects the H_2O saturated fields

528 near the solidus (e.g. paths D & E in Fig. 8). H₂O saturation in melt is strongly dependent
529 on pressure (Tuttle & Bowen, 1958; Johannes & Holtz, 1996). Therefore, the volume of
530 water released by a crystallising melt will depend on the initial H₂O content of the melt and
531 the pressure at which the melt intersects the H₂O-in boundary. These three mechanisms
532 may result in greater melt fertility than is defined by a SWL-moderated minimum melting
533 curve. However, under most geological conditions SWL paths and melt fertility are likely to
534 apply.

535 The modelling presented here does not take into consideration the effect of melt loss on
536 melt fertility of felsic rocks as this process involves modification of several compositional
537 vectors the cannot be represented on a simple T - M_{H_2O} diagram. The loss of melt will cause
538 the bulk composition to become more residual with the largest compositional change likely
539 to be a decrease in water content due to the hydrous nature of the melt derived from
540 melting of felsic rocks (White & Powell, 2002; 2010).

541 **DISCUSSION and CONCLUSIONS**

542 **General conclusions**

543 The water content of the rock comprises the sum of free H₂O and structurally-bound
544 H₂O residing in hydrous minerals (e.g. biotite, muscovite and hornblende). While free H₂O
545 can be lost from a source rock (due to compaction, deformation, migrating porosity, etc.),
546 structurally-bound H₂O is only made mobile when the hydrous host mineral becomes
547 unstable due to changes in the P - T - X conditions at equilibrium. This may occur at sub
548 solidus and/or suprasolidus conditions. Above the solidus, any H₂O liberated by the
549 breakdown of hydrous phases is incorporated into the melt phase (until it is H₂O

550 saturated). The H₂O contents of the systems described here cannot evolve to be less than
551 the value of the minimum saturation point without also losing melt and/or solid phases
552 (Fig. 8). In this way subsolidus water loss can be viewed as defining the minimum
553 H₂O condition for a particular bulk composition that has undergone prograde heating. A
554 corollary of this is that the subsolidus water loss path defines the minimum melt fertility of
555 a rock, at least prior to melt loss.

556 Most melting experiments are likely to conserve any H₂O generated by devolatilisation
557 of hydrous minerals as the experimental charge is heated and pressurised from ambient
558 temperatures up to the conditions of the experimental run. Given the likelihood that
559 subsolidus water loss is a significant phenomena affecting prograde metamorphic rocks,
560 most experimentally derived melting curves will tend to overestimate the melt fertility of
561 the starting material at the onset of melting within the continental crust. The modelled
562 curves presented here allow subsolidus water loss to be taken into account and, therefore,
563 allow determination of the minimum melting relationships for compositions of interest.

564 **Implications for melt connectivity, melt loss and the rheology of the crust** 565 **during anatexis**

566 An important consequence of the reduced melt fertility implied by the subsolidus water loss
567 model is that critical melt thresholds will be intersected at significantly higher temperatures
568 (Fig. 6). At low-melt fractions melt will initially form in isolated domains along grain edges
569 and corners between reactant phases (Mehnert *et al.*, 1973). As the melt fraction increases
570 these isolated melt volumes begin to coalesce eventually forming an interconnected network
571 through the rock volume (Vigneresse *et al.*, 1996). Rosenberg & Handy (2005) identified a

572 melt connectivity threshold which occurs at $\Phi \approx 0.07$ and corresponds to $\sim 80\%$ of grain
573 boundaries containing melt. Experimental data suggests that rocks experience significant
574 loss of strength between the onset of melting and the melt connectivity threshold (from
575 ~ 800 MPa down to ~ 200 MPa; Rosenberg & Handy, 2005). It follows that rocks having
576 experienced subsolidus water loss will remain comparatively stronger at higher
577 temperatures than rocks that have not. For the bulk compositions modelled here the
578 differences in the temperature of the melt connectivity threshold range from $<10^\circ\text{C}$ to
579 $>150^\circ\text{C}$. Such a relationship between degree of melting, temperature and strength is likely
580 to significantly influence the rheological behaviour of crust undergoing regional
581 metamorphism and anatexis.

582 Higher degrees of melting are associated with critical thresholds governing melt
583 segregation and the transition from a solid to a liquid rheology. The melt escape threshold
584 refers to the melt fraction at which melt is able to segregate from the source (Vigneresse
585 *et al.*, 1996). The melt escape threshold is likely to depend on a number of factors including
586 viscosity and strain rate but is likely to occur at $\Phi \sim 0.2\text{--}0.25$ (Sawyer, 1994; Vigneresse
587 *et al.*, 1996). The ‘solid-to-liquid transition’ is the melt content at the transition from a
588 solid-supported to a liquid-supported structure (Rosenberg & Handy, 2005). The
589 solid-to-liquid transition occurs at $\Phi \sim 0.4\text{--}0.6$. Both the melt escape threshold and the
590 solid-to-liquid transition will be shifted to higher temperatures for a rock that has
591 experienced subsolidus water loss.

592 Subsolidus water loss during magma crystallisation



593 The discussion so far has concerned the consequences of subsolidus water loss during
 594 prograde heating, however crystallising magmas may also experience subsolidus water loss.
 595 Large amounts of H₂O may be generated as a magma approaches and then crosses the
 596 solidus if a magma had an initially high H₂O content. **The magma is rapidly converted to**
 597 **solids + H₂O across the narrow melt + H₂O fields just above the wet solidus (Paths D & E**
 598 **in Fig. 8). Even at** upper crustal conditions the inter-grain porosity is likely to be small and
 599 the tendency for free H₂O to escape the now solid rock volume is likely to be high (Connolly,
 600 1997). The escape of free H₂O will cause the bulk H₂O contents to decrease towards the
 601 minimum saturation point. Further cooling will take the rock across the H₂O-out boundary
 602 and the H₂O contents will become fixed. An important consequence of this is that the final
 603 H₂O contents of the crystallised solid (e.g. a tonalite) will not reflect its H₂O content when
 604 it was a magma but rather reflects the water contents in the vicinity of the minimum
 605 saturation point. This would mean that analysing the H₂O contents of granites collected in
 606 the field may tell us nothing about the H₂O contents of the original magma.

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765 TABLES

Table 1. Starting compositions used in this study in wt.% and equivalent NCKFMASHTO mol.% values. The compositions are and the corresponding experimental studies are as follows: VH88P - high-Al metapelite from Vielzeuf & Holloway (1988), G95P - low-Al metapelite from Gardien *et al.* (1995), VM94G - metagreywacke from Vielzeuf & Montel (1994), and PD97T - metatonalite from Patino-Douce (1997).

Sample	VH88P		G95P		VM94G		PD97T	
	wt. %	mol. %	wt. %	mol. %	wt. %	mol. %	wt. %	mol. %
SiO ₂	64.35	66.64	70.04	72.14	69.99	72.27	61.48	66.56
Al ₂ O ₃	18.13	11.06	14.92	9.06	12.96	7.89	17.07	10.87
Fe ₂ O ₃	—	—	—	—	0.45	—	—	—
FeO	6.26	5.42	3.65	3.14	4.42	4.16	5.92	5.35
MgO	2.44	3.77	1.39	2.13	2.36	3.64	2.68	4.32
MnO	0.09	—	0.09	—	0.06	—	0.11	—
CaO	1.52	1.69	1.79	1.98	1.67	1.85	5.39	6.24
Na ₂ O	1.66	1.67	2.65	2.65	2.95	2.95	3.88	4.07
K ₂ O	2.56	1.69	3.18	2.09	2.41	1.59	2.57	1.77
TiO ₂	0.82	0.64	0.38	0.29	0.7	0.54	0.9	0.73
P ₂ O ₅	—	—	—	—	0.2	—	—	—
CO ₂	—	—	—	—	0.1	—	—	—
water	2.15	7.42	1.9	6.52	1.43	4.92	—	—
F	—	—	—	—	0.09	—	—	—
S	—	—	—	—	0.067	—	—	—
B(ppm)	—	—	—	—	20	—	—	—
Total	99.98	100.00	99.99	100.00	99.857	99.81	99.89	100.00

Table 2. Compositions used to construct pseudosections (in mol.%).

Figure	Composition	Diagram	H ₂ O	SiO ₂	Al ₂ O ₃	CaO	MgO	FeO	K ₂ O	Na ₂ O	TiO ₂	O
Figures 2 & 3												
(a)	VH88P	$T-M_{\text{H}_2\text{O}}$ lhs	0.11	71.90	11.93	1.82	4.07	5.85	1.82	1.80	0.69	0.01
		$T-M_{\text{H}_2\text{O}}$ rhs	14.02	61.88	10.27	1.57	3.50	5.03	1.57	1.55	0.59	0.01
(b)	G95P	$T-M_{\text{H}_2\text{O}}$ lhs	0.11	76.94	9.67	2.11	3.35	2.27	2.23	2.82	0.31	0.20
		$T-M_{\text{H}_2\text{O}}$ rhs	13.80	66.39	8.34	1.82	2.89	1.96	1.92	2.44	0.27	0.17
(c)	VM94G	$T-M_{\text{H}_2\text{O}}$ lhs	0.52	75.62	8.25	1.93	3.81	4.35	1.66	3.09	0.56	0.20
		$T-M_{\text{H}_2\text{O}}$ rhs	12.41	66.58	7.26	1.70	3.36	3.83	1.46	2.72	0.50	0.17
(d)	PD97T	$T-M_{\text{H}_2\text{O}}$ lhs	0.50	66.22	10.83	6.22	4.30	5.33	1.76	4.06	0.73	0.05
		$T-M_{\text{H}_2\text{O}}$ rhs	9.92	59.95	9.81	5.63	3.90	4.83	1.60	3.67	0.66	0.05
Figure 4												
(a)	VH88P - CSW	mode(X)-T	7.42	66.63	11.06	1.69	3.77	5.42	1.69	1.67	0.64	0.01
(b)	VH88P - SWL	mode(X)-T	3.45	69.50	11.53	1.76	3.93	5.65	1.76	1.74	0.67	0.01
(c)	G95P - CSW	mode(X)-T	6.52	72.00	9.05	1.97	3.13	2.12	2.08	2.64	0.29	0.19
(d)	G95P - SWL	mode(X)-T	4.08	73.88	9.28	2.03	3.22	2.18	2.14	2.71	0.30	0.19
(e)	VM94G - CSW	mode(X)-T	5.28	72.01	7.85	1.84	3.63	4.14	1.58	2.94	0.54	0.19
(f)	VM94G - SWL	mode(X)-T	2.78	73.91	8.06	1.89	3.72	4.25	1.62	3.02	0.55	0.19
(g)	PD97T - CSW	mode(X)-T	3.19	64.43	10.54	6.05	4.19	5.19	1.71	3.95	0.71	0.05
(h)	PD97T - SWL	mode(X)-T	2.57	64.84	10.61	6.09	4.21	5.22	1.72	3.97	0.71	0.05
Figure 5												
(a)	VH88P - SWL	mode(X)-T	3.47	69.48	11.53	1.76	3.93	5.65	1.76	1.74	0.67	0.01
(b)	G95P - SWL	mode(X)-T	3.55	74.29	9.34	2.04	3.23	2.19	2.15	2.73	0.30	0.18
(c)	VM94G - SWL	mode(X)-T	2.88	73.84	8.05	1.88	3.72	4.25	1.62	3.02	0.55	0.19
(d)	PD97T - SWL	mode(X)-T	2.48	64.90	10.62	6.10	4.22	5.22	1.73	3.98	0.72	0.05

766 **FIGURE CAPTIONS**

767 Fig. 1 Harker plots of mol.% oxides, molar K_2O / Na_2O and ASI (molar
 768 $Al_2O_3 / (CaO + Na_2O + K_2O)$) for the starting compositions: VH88P - metapelite, G95P
 769 - metapelite, VM94G - metagreywacke and PD97T - metatonalite. L, M and U
 770 represent the estimated average compositions of the lower-, middle- and upper-crust,
 771 respectively, of Rudnick & Gao (2003). All values are plotted on an anhydrous basis.

772 Fig. 2 Calculated NCKFMASHTO T - M_{H_2O} pseudosections for the compositions: (a)
 773 VH88P - metapelite, (b) G95P - metapelite and (c) VM94G - metagreywacke at
 774 10 kbar and (d) PD97T - metatonalite at 8 kbar (see Table 2). The dashed line
 775 indicates the solidus for each diagram. The white dot indicates the locus of the
 776 minimum saturation point for each diagram.

777 Fig. 3 Calculated contours for the modal abundances of melt (solid lines) and free
 778 H_2O (dashed lines) for the compositions: (a) VH88P - metapelite, (b) G95P -
 779 metapelite, (c) VM94G - metagreywacke and (d) PD97T - metatonalite (see Table 2).
 780 The dark grey areas represent the regions over which free-water is stable. The grey
 781 arrows represent the path followed for heating in a closed system with the
 782 H_2O contents fixed at the value reported for the experimental starting composition.
 783 The H_2O contents for the tonalitic bulk composition, (d), is unknown so the
 784 H_2O contents were determined so that 30% melt is generated at 950 °C as reported in
 785 Patino-Douce (1997). As with Fig. 2 the white dot indicates the locus of the minimum
 786 saturation point for each diagram. The black arrows represent heating combined with
 787 subsolidus water loss. The stars represent melt fraction estimates with increasing
 788 temperature derived from the corresponding melting experiments (assuming

789 uncertainty in the H₂O contents only; see text for discussion).

790 Fig 4 Calculated molecular proportions of phases (X) with increasing temperature for the
 791 four compositions: VH88P, G95P and VM94G at 10 kbar and PD97T at 8 kbar under
 792 conditions of conservation of water () and sub-solidus water loss (SWL). The dashed
 793 vertical line indicates the solidus for each diagram.

794 Fig 5 Calculated molecular proportions of phases (X) with increasing temperature for the
 795 four compositions: VH88P, G95P and VM94G at 5 kbar and PD97T at 4 kbar under
 796 conditions of sub-solidus water loss (SWL). The solidus is at the left edge for each
 797 diagram.

798 Fig 6 Comparison of the CSW and SWL melt fraction curves (shown in figure 5) with the
 799 available experimental estimates for the four compositions VH88P, G95P and VM94G
 800 at 10 kbar and PD97T at 8 kbar . The horizontal dashed lines indicate the melt
 801 fractions corresponding to the solid-liquid transition (SLT) and melt connectivity
 802 threshold (MCT; both after Rosenberg & Handy, 2005) and the melt escape threshold
 803 (MET; after Vigneresse *et al.*, 1996). The arrows indicate the difference in the
 804 temperature of the MCT between the CSW and SWL scenarios (see text for
 805 discussion).

806 Fig 7 Comparison of the melt fraction curves with increasing temperature at low P (light
 807 grey curve) and high P (dark grey curve) for the four compositions VH88P, G95P and
 808 VM94G (5 kbar and 10 kbar) and PD97T at (4 kbar and 8 kbar). The respective
 809 solidi at each pressure are indicated by the corresponding grey arrows. The position of
 810 the MCT is shown by the horizontal dashed line.

811 Fig 8 Schematic diagram showing the relationships between various melting paths (dark
812 grey dotted lines) and crystallisation paths (light grey dotted lines) as a function of
813 water contents. Thin grey lines represent melt contours (increasing to the top-right of
814 the diagram). The striped area is inaccessible because H_2O contents less than this
815 value are unobtainable without invoking changes in other compositional vectors (e.g.
816 via melt and or solid segregation; see text for discussion).

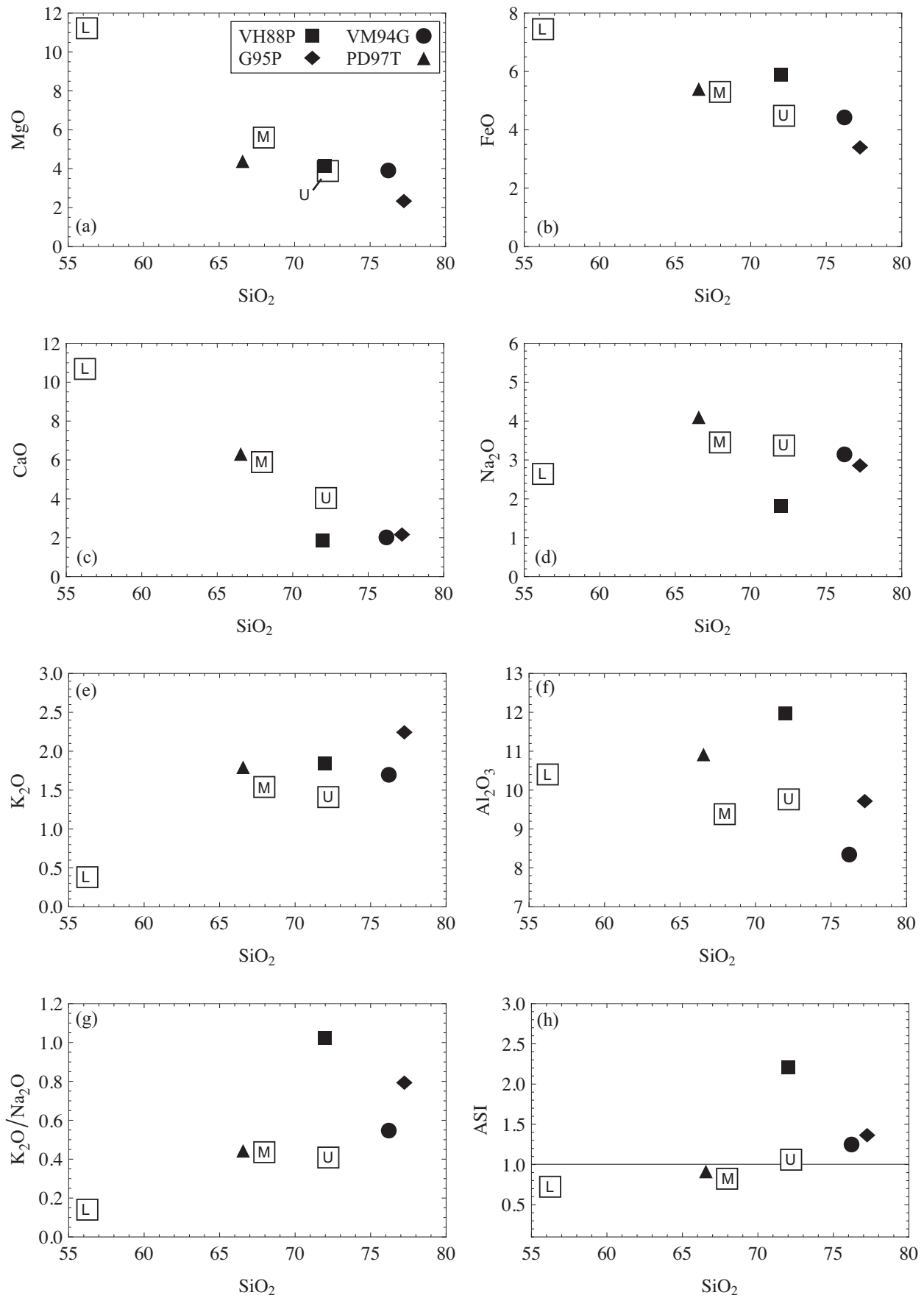


Figure 1.

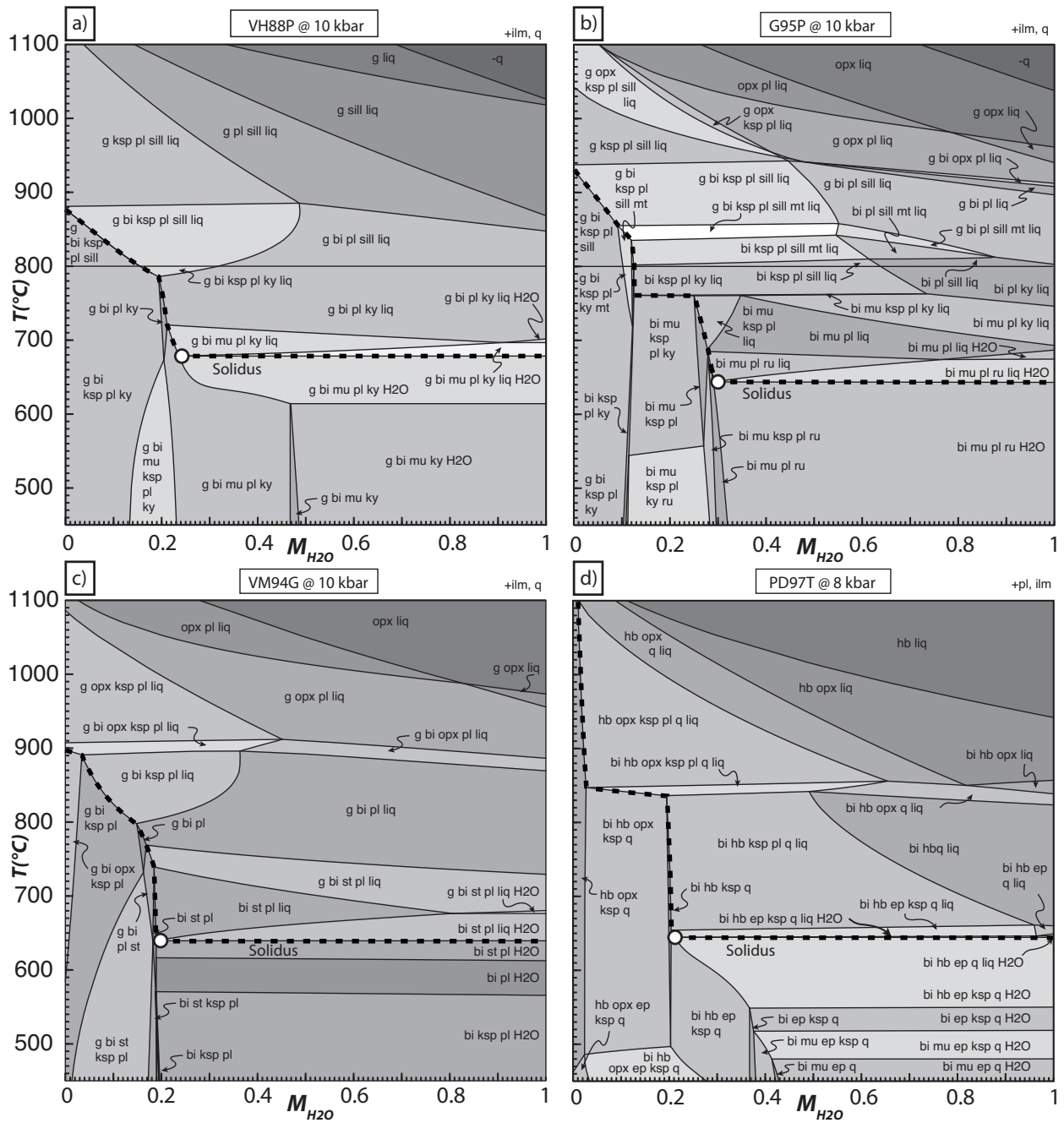


Figure 2.

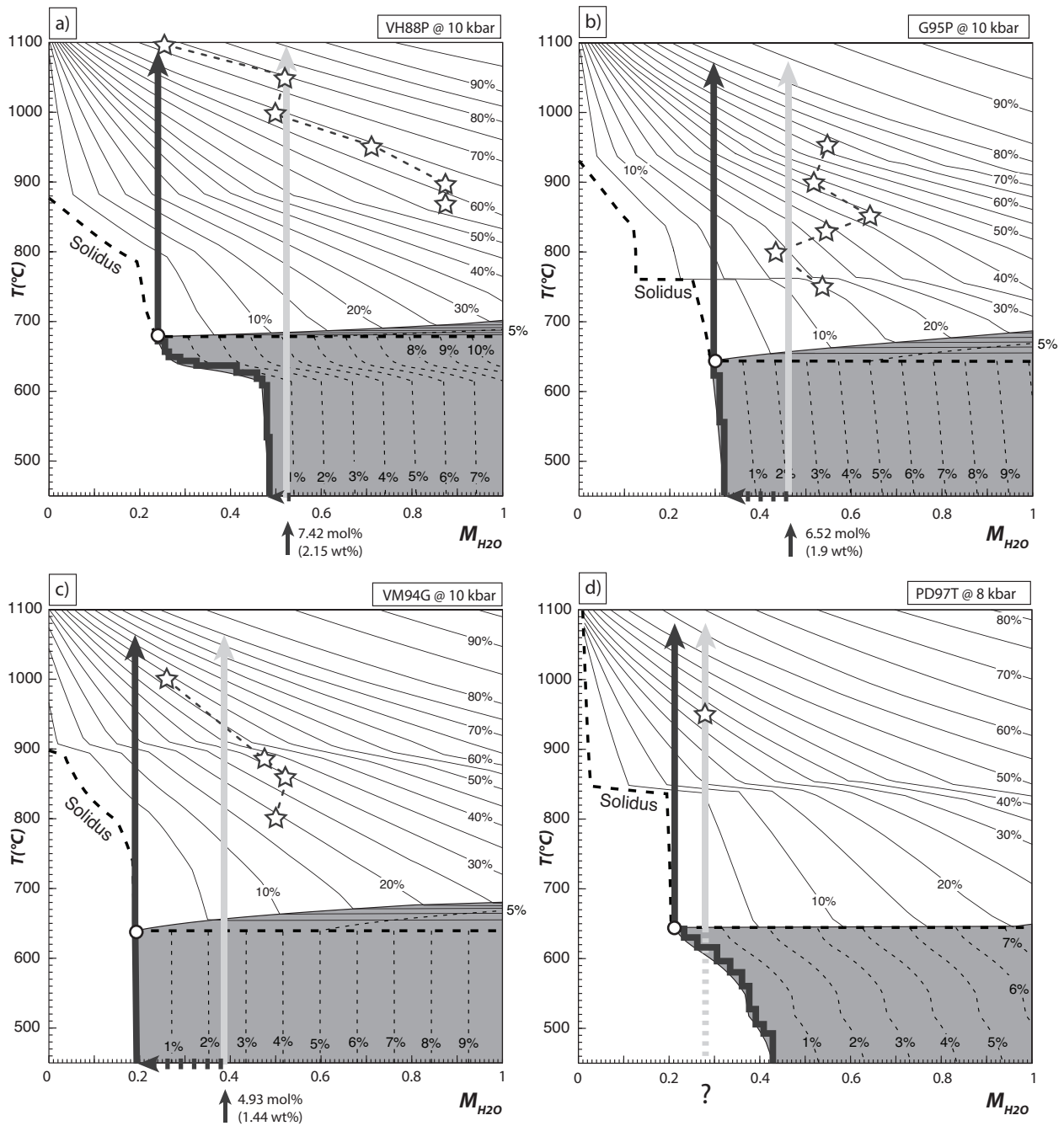


Figure 3.

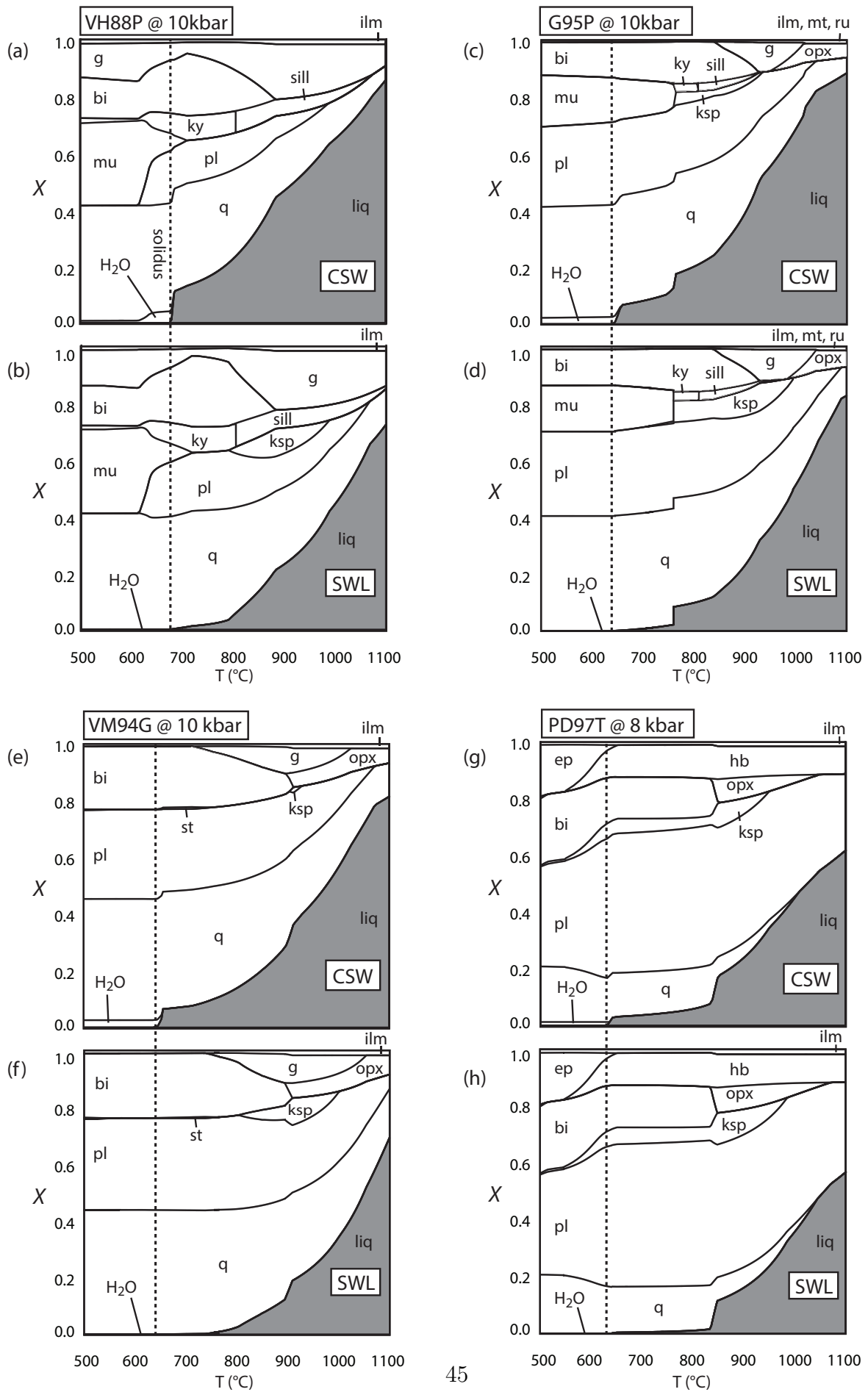


Figure 4.

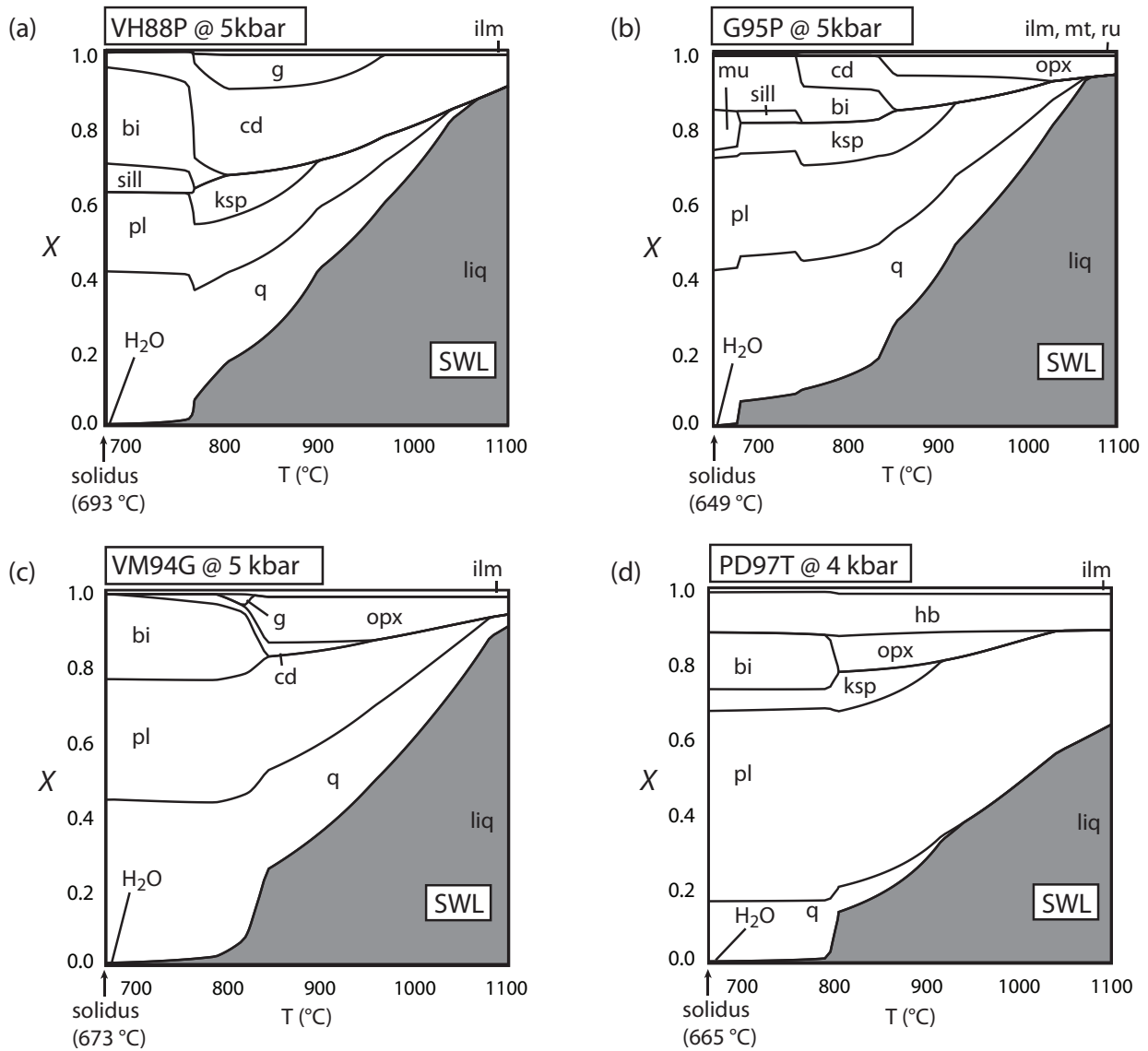


Figure 5.

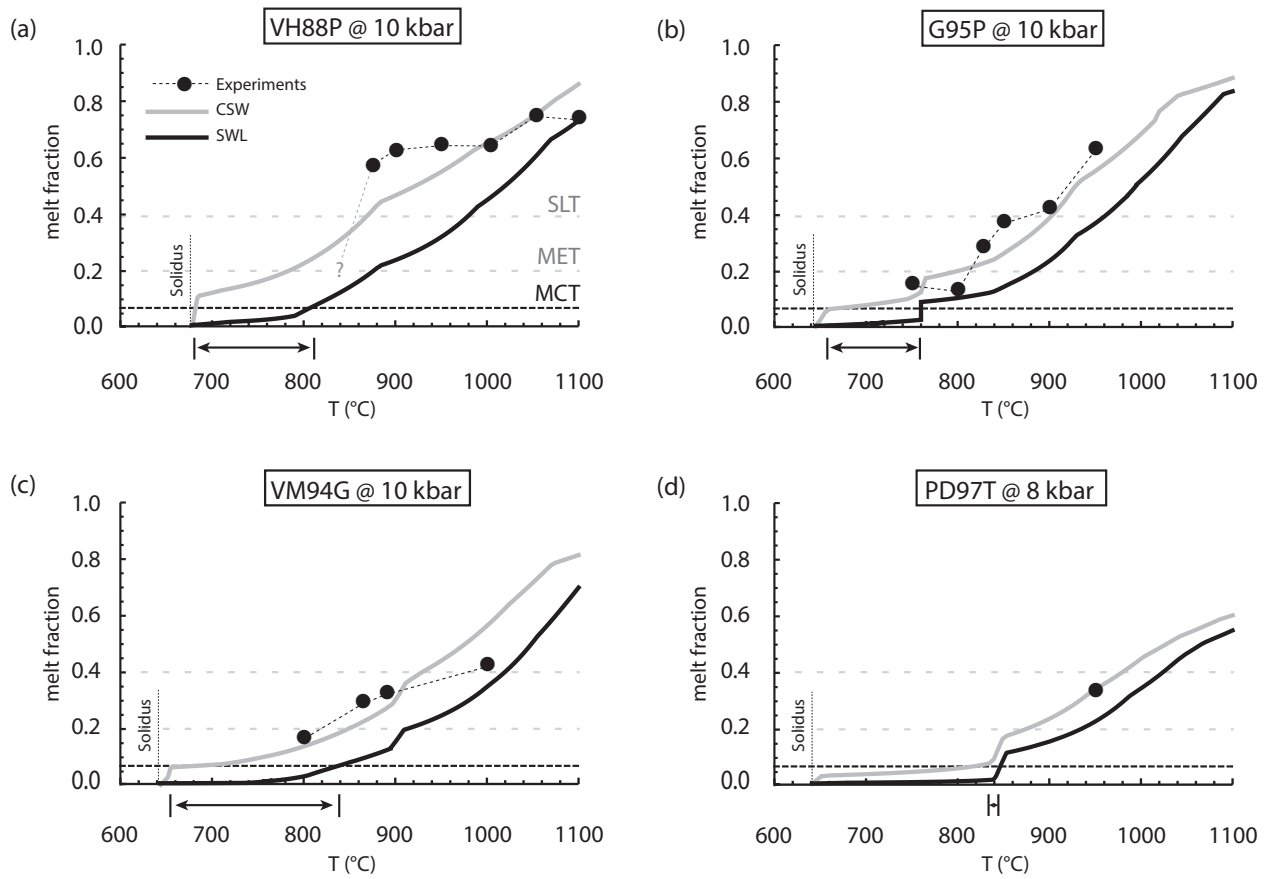


Figure 6.

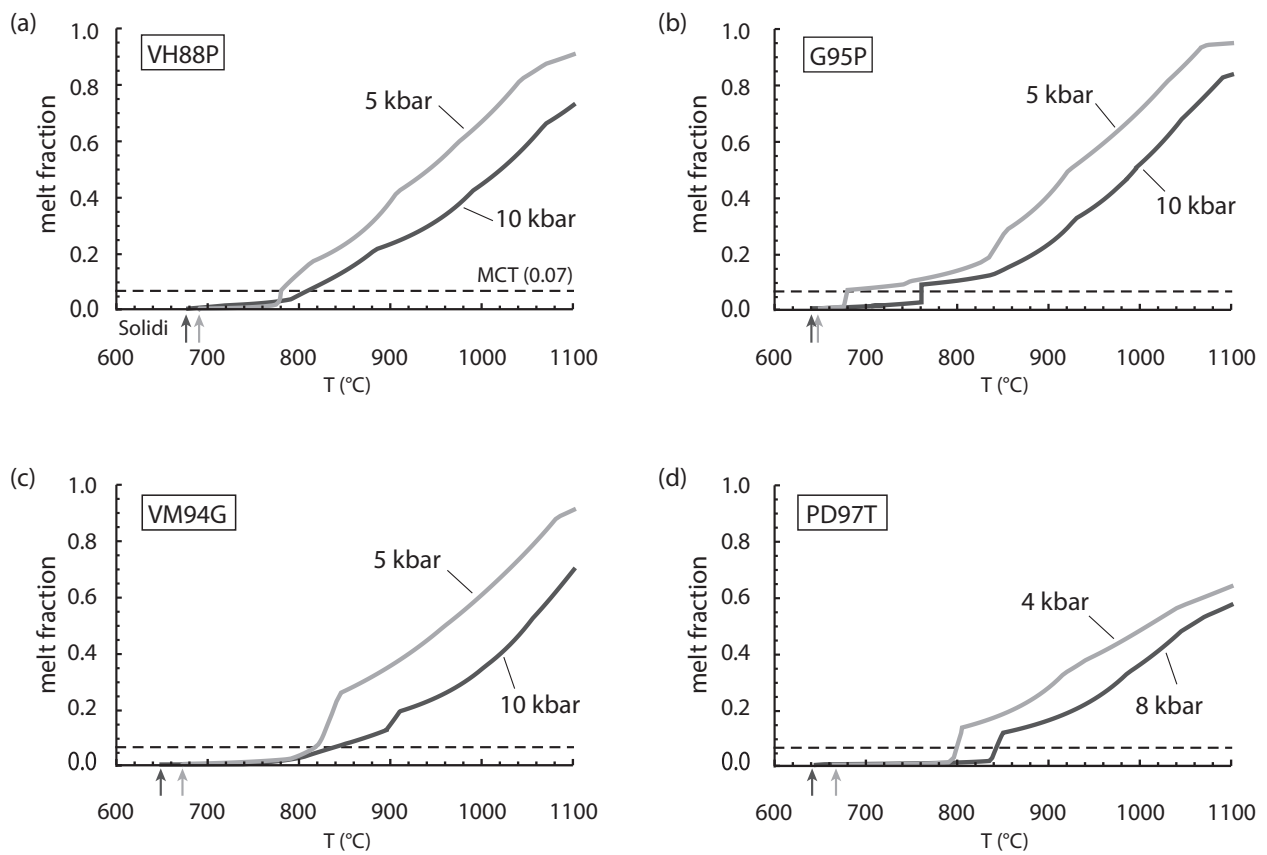


Figure 7.

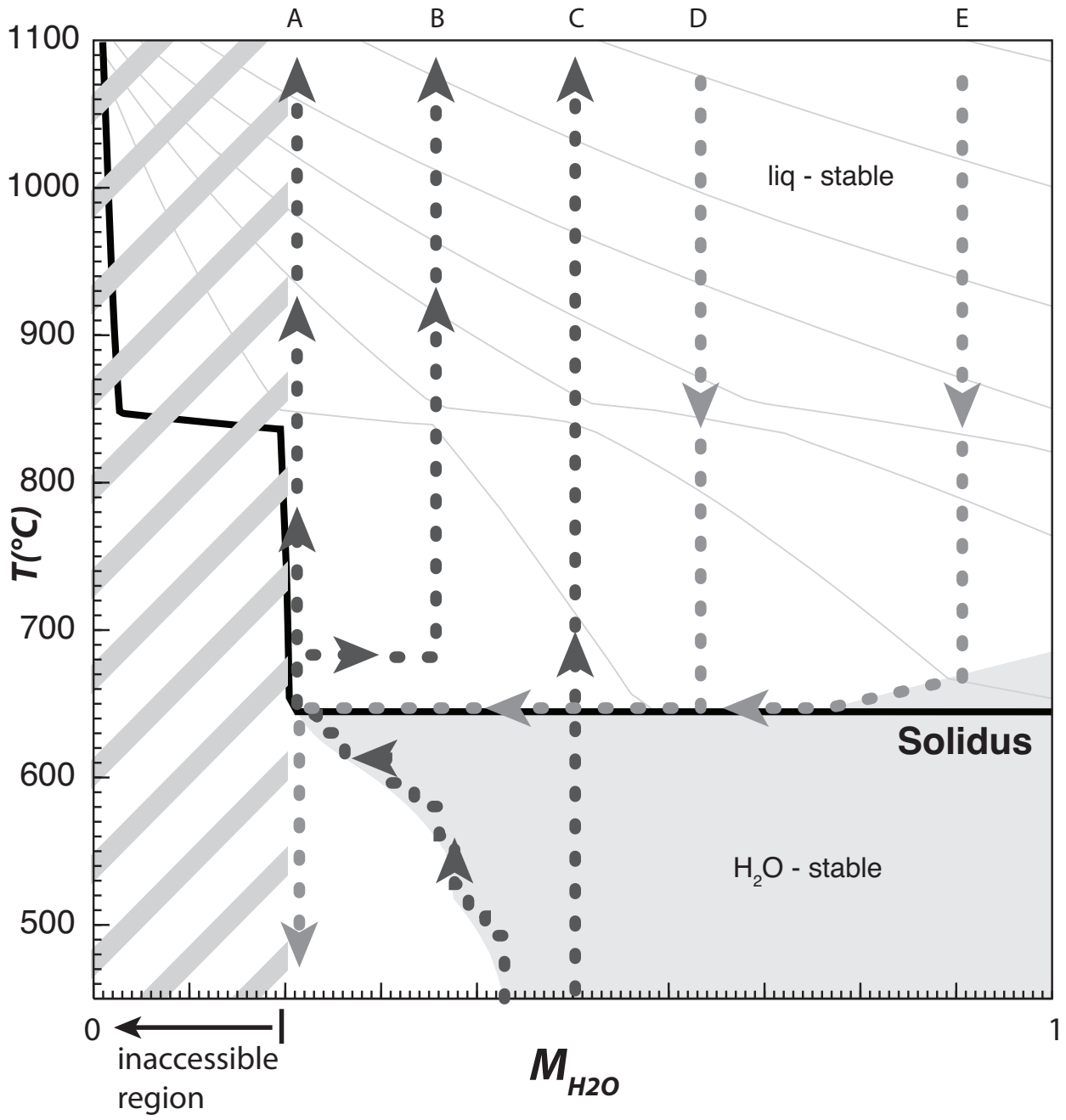


Figure 8.