Article

Melt Reintegration Modelling: Testing against a Subsolidus Reference Assemblage

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Abstract: Phase equilibria modelling incorporating melt reintegration offers a methodology to create hypothetical rock compositions that may have existed prior to melt loss, allowing the potential prograde evolution of rocks to be explored. However, melt reintegration modelling relies on assumptions concerning the volume of melt that was lost and is generally restricted by the absence of direct constraints on the pre-anatectic mineral assemblages. Mg-rich granulite in the 514–490 Ma Delamerian Orogen in southern Australia contains spinel–cordierite symplectic intergrowths that surround rare, coarse blocky domains of sillimanite. These sillimanite cores, as well as the widespread presence of andalusite in lower grade areas of the southern Delamerian Orogen, suggest that the subsolidus precursor to the granulite contained andalusite. This provides the opportunity to test if melt reintegration modelling of the granulite predicts subsolidus andalusite. Stepwise down-temperature melt reintegration modelling produces a water-saturated solidus after the addition of 12 mol% melt. When modelled at subsolidus conditions, the resulting rock composition produces andalusite-bearing assemblages with andalusite modes similar to the abundance of the sillimanite-cored spinel–cordierite intergrowths. The modelling results from this case study suggest that melt reintegration modelling is a valid method to recreate prograde subsolidus bulk rock compositions.

Keywords: P–T pseudosection; low pressure metamorphism; delamerian orogen; melt reintegration; melt loss

1. Introduction

Melting and melt loss during granulite facies metamorphism is a fundamental process that differentiates the continental crust and causes chemical depletion in the residual material [1–3]. Advances in thermodynamic modelling of mineral assemblages have enabled quantitative phase equilibria modelling of the melting processes operating in partially molten siliciclastic rocks in increasingly complex chemical systems [4–11]. These models have been used to investigate the effects of melting and melt loss on crustal rheology, composition and the preservation of geochronometers e.g., [1,5,12–15]. However, the application of these models has been limited to theoretical situations, as in almost all real cases there is little information about the composition of the rock prior to melting cf. [11]. This lack of information about the likely protolith composition also means that it is essentially impossible to determine the prograde P–T evolution of a granulite facies rock, because the composition of the rock in which prograde minerals developed has been irrevocably modified [16]. For this reason, quantitative petrological interpretation of mineral inclusions captured within prograde minerals
is difficult and often impossible. In trying to reconstruct potential prograde bulk compositions, metamorphic petrologists approach the problem via melt reintegation modelling, in which theoretical compositions of melt are integrated into a rock composition in order to create a hypothetical “protolith” bulk composition. A recent study by Bartoli [17] has shown that a number of methods of melt reintegation modelling, including the stepwise method used here, are able to produce plausible subsolidus bulk rock compositions. However, significant underestimations of melt productivity may still occur, as exact protolith compositions are unknown in most cases. While useful, this process relies on assumptions regarding the prograde metamorphic path and volume of melt that has been lost and is commonly done up to a point at which a water-saturated solidus is achieved [18,19]. The determination of a final composition is unconstrained, as the prograde compositions often no longer exist for comparison, except in some rare cases e.g., [20,21]. However, one way to assess the viability of melt reintegration modelling as a means of determining a reasonable prograde rock composition is to choose a granulite facies sample where the prograde subsolidus mineralogy is relatively well constrained.

The aim of this study is to use melt reintegration to reproduce the original bulk composition of a residual granulite facies rock from the Delamerian Orogen in southern Australia. The existence of andalusite in the subsolidus assemblage provides a test to see whether melt reintegration P–T modelling of the reintegrated bulk composition stabilises the observed modal abundance of subsolidus andalusite, giving some constraints on the validity of model-generated subsolidus bulk compositions.

2. Geological Setting

The footprint of the 514–490 Ma Delamerian Orogeny has a north–south extent from the Peake and Denison Inliers in the far north of South Australia, to the western tip of Kangaroo Island in the south (Figure 1). It extends to the east into western New South Wales, Victoria and Tasmania. The Antarctic continuation of the orogen is the Ross Orogen [22,23], which formed along the same paleo-continental margin but largely predates the Delamerian Orogeny. In South Australia, Victoria, western New South Wales and Tasmania, the Delamerian Orogen is primarily composed of Late Neoproterozoic and Early Cambrian sedimentary rocks that were deposited in a passive margin setting. In South Australia, the Early Cambrian sequences comprise the basal Normanville and overlying Kanmantoo Groups that were deposited in a passive margin setting. In South Australia, the Early Cambrian sequences comprise the basal Normanville and overlying Kanmantoo Groups that were deposited in the Stansbury Basin [24–26].

The early compressional stages of the Delamerian Orogeny involved west-vergent thrusting (D1) and the development of a low-angle fabric (S1), which was overprinted by two phases of tight to open upright folds during D2 and D3 [28–30]. The early stages of the event were synchronous with granitic magmatism throughout the region, with the earliest syntectonic granite intruded during the onset of orogenic crustal shortening at 514 ± 4 Ma [31]. Increasingly mantle-derived magmatism marked the end of the orogenic event (c. 490 Ma) and continued into the Early Ordovician [32–34]. Following the Delamerian Orogeny, an extensional setting was re-established and post-orogenic magmatism resulted in the intrusion of rhyolite and diorite dikes [27]. Within the Kanmantoo Group, Delamerian-aged metamorphism is characterised by mostly amphibolite facies low-pressure–high-temperature metamorphism, resulting in the formation of andalusite- to sillimanite-bearing assemblages in aluminous rocks [31,35]. Metamorphic grade increases in proximity to the late-tectonic granitic intrusives, culminating in extensive partial melting [27,36,37].

The Kanmantoo Group metasedimentary rocks at Reedy Creek consist primarily of extensively migmatised metapsammitic lithologies interlayered with thin discontinuous layers of metapelite. The hosting metapsammitic lithologies are primarily composed of quartz–plagioclase–cordierite–biotite ± sillimanite. The sequence contains a pervasive biotite-bearing foliation that is parallel to and cross-cut by migmatitic segregations. Abundant partial melting has resulted in the segregation of leucosome and melanosome. Thin leucosomes (c< 2 cm wide) have been folded with the original foliation, suggesting that some melting occurred in the early stages of the Delamerian Orogeny. However, some leucosomes are only slightly deformed and others cross-cut the original foliation, suggesting that melting continued
throughout the whole event. The migmatitic rocks were intruded by the Reedy Creek Granodiorite and associated diorites at 492.6 ± 1.1 Ma [27]. The granodiorite is weakly foliated parallel to the fabric in the migmatitic metasediments. Within the metasediments, decimetre-scale bodies of diorite have been converted to hornblende-bearing migmatites that contain K–feldspar-bearing melts.

Figure 1. Detail of the geology in the Delamerian Orogen as seen in the southern Adelaide Fold Belt. After Foden et al. [27].

Although the bulk of the metasedimentary sequence consists of metapsammitic lithologies with thermodynamically insensitive bulk compositions, thin discontinuous layers of cordierite-dominated Mg-rich (MgO/(MgO+FeO) = 0.66) metapelite—that were likely to have been mud layers in the original sequence—form sensitive mineral assemblages that can be used to investigate the $P$–$T$ evolution of the sequence. The discontinuous metapelite layers contain domains that are dominantly composed of orthopyroxene–biotite–cordierite–plagioclase (Figure 2a,b). Locally, these domains contain cordierite–spinel symplectites that rarely surround coarse/blocky aggregates of sillimanite (Figure 2c). In adjacent psammitic layers, cordierite occurs as porphyroblasts within felsic segregations and veins. We interpret the orthopyroxene and cordierite-bearing metapelitic layers to represent residual domains that have lost melt. These layers do not contain assemblages or chemistry typical of granitic compositions that would form leucosomes, due to their high Mg content and abundant orthopyroxene and cordierite. The peritectic mineral assemblages preserved in these layers are very fresh and relatively unretrogressed, indicating that melt has been drained from these layers [38]. Blocky, porphyroblastic aggregates of sillimanite surrounded by cordierite–spinel symplectites (up to 3 cm by 1 cm) (Figure 2c) in the pelite layers are interpreted to reflect the former presence of
andalusite, based on the extensive presence of andalusite in lower grade areas of the Kanmantoo Group (Figure 2d) [29,30,39]. Therefore, we infer the prograde sequence to have contained andalusite. Despite the intensity of the deformation, there is no evidence for distributed sillimanite within the foliation, suggesting that the proportion of blocky sillimanite aggregates represents the approximate proportion of andalusite in the protolith. Layers containing these sillimanite aggregates are relatively rare in the Kanmantoo Group sequence, but in the cordierite-dominated Mg-rich layers they comprise around 5% of the current rock volume.

Figure 2. (a) Mg-rich layer dominated by cordierite–biotite–orthopyroxene–plagioclase; (b) Thin, discontinuous orthopyroxene-bearing domain; (c) Rock sample of the thin layer that was sampled showing cordierite–spinel symplectites, amongst orthopyroxene and plagioclase; (d) Andalusite porphyroblasts enclosed within metapelite from a lower grade area of the Delamerian (54K 0316690 E, 6122806 S, datum WGS84).

Samples and Petrography

The sample used in this study (sample 14-RDC-04) was taken from a thin metapelitic layer (>10 cm in width), within thick metapsammitic layers in Reedy Creek (Figures 1 and 2a,b). Sample 14-RDC-04 contains orthopyroxene, biotite, cordierite, spinel, plagioclase, magnetite, ilmenite and minor sillimanite. Coarse-grained (1–6 mm) euhedral grains of orthopyroxene are common throughout the leucosomes and contain inclusions of biotite, ilmenite and magnetite. Coarse-grained, poikiloblastic cordierite (1–6 mm) contains inclusions of biotite, plagioclase, magnetite and ilmenite. Symplectites of spinel and cordierite (in patches 4–10 mm in diameter) are common (Figure 3a). In rare instances, these cordierite–spinel symplectites surround aggregates of sillimanite grains (3–6 mm; Figure 3b,c). Orthopyroxene grains occur in direct contact with biotite and are surrounded by cordierite and plagioclase (Figure 3d). There are two morphologies of biotite. Some biotite grains are aligned and define a weak foliation, whereas other biotite is randomly oriented. Biotite adjacent to cordierite
occurs as ragged anhedral grains. No biotite is present in the areas immediately adjacent to the cordierite–spinel symplectites (Figure 3a,c). Cordierite–spinel symplectites and sillimanite grains do not show any orientation and no foliation is observed in the biotite-absent regions. We interpret the peak assemblage present in this sample to be plagioclase + biotite + orthopyroxene + cordierite + spinel + magnetite + ilmenite. The rare, coarse-grained domains of sillimanite enclosed within spinel–cordierite symplectites are interpreted to be textural relics from replacement of prograde andalusite. Although sillimanite and orthopyroxene are in close proximity (Figure 3d), we do not interpret them to represent former stable coexistence.

Figure 3. Photomicrographs and thin section photographs of sample 14-RDC-04. (a) Spinel and cordierite symplectites; (b) Relict sillimanite grain, showing typical crosshatching andalusite replacement texture, surrounded by symplectic corona of spinel and cordierite; (c) Relict sillimanite surrounded by a cordierite–spinel corona; (d) Anhedral orthopyroxene grains adjacent to semi-oriented biotite. Biotite becomes unoriented and has an anhedral shape when in contact with cordierite.

3. Materials and Methods

The pelitic orthopyroxene-bearing domains that make up sample 14-RDC-04 are thin and irregularly shaped, meaning that rock slabs cut for whole rock geochemistry contain a mixture of residual pelite and the surrounding wall rock, and thus may not provide a valid composition for the purposes of phase equilibria modelling. Instead, a chemical composition of the orthopyroxene-bearing domains was determined by combining average measured mineral chemistry and density with estimates of mineral proportions in these domains. In the domain used for the determination of mineral modal abundances, sillimanite is almost completely replaced by blocky aggregates of cordierite–spinel, so was not considered as part of the composition. Estimations of the proportions of minerals present in the sample were determined by pixel counting minerals in thin section images using Adobe Photoshop. The area of thin section used for pixel counting was quite large, at approximately 7.5 × 3.5 cm (26.25 cm²). As the domains that were sampled were quite thin, we believe that this 2D
area is representative of the equilibrium volume. The use of individual mineral chemistry to determine a bulk composition is a relatively common practice e.g., [40,41]; however, it should be noted that this method may produce different results than compositions acquired using XRF (X-ray Fluorescence), see [42]. Mineral chemical analyses were obtained using a Cameca SXFive electron microprobe at Adelaide Microscopy, with standard operating procedures. The mineral modes and average mineral chemical analyses and densities used for the determination of the residual pelite composition are provided in Table 1.

Table 1. Mineral modal proportions and average mineral chemistry used to obtain a chemical composition of the orthopyroxene-bearing domains for the purposes of phase equilibria modelling. Individual mineral analyses can be found in Supplementary materials.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Mode (wt %)</th>
<th>Bi</th>
<th>Cd</th>
<th>Opx</th>
<th>Sp</th>
<th>Pl</th>
<th>Total</th>
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<td>SiO₂</td>
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<td>22.58</td>
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<tr>
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<td>98.87</td>
<td>101.22</td>
<td>100.05</td>
<td>99.99</td>
<td>101.08</td>
<td>100</td>
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One of the uncertainties of pseudosection modelling is determining the bulk composition that relates to the formation of the peak metamorphic mineral assemblage, particularly the amount of H₂O and Fe₂O₃. P–M 𝑂, P–M H₂O and T–M H₂O sections (where the oxidation state (M_O) or amount of water (M_H₂O) is varied as a function of pressure (P) or temperature (T); Appendix A were calculated to assess the sensitivity of the compositions of the sample to changing H₂O content and oxidation state (O), and determine appropriate values for the domain composition. The P–M O and P–M H₂O sections were drawn at a constant temperature of 750 °C, which was chosen based on a first pass estimate of the likely metamorphic conditions of the suprasolidus assemblage and the assumed sensitivity of the assemblage to pressure. A T–M H₂O section was modelled after the P–M H₂O to better assess the H₂O content of the sample due to the greater sensitivity of H₂O-dependent equilibria to temperature. The T–M H₂O section was calculated at a fixed pressure of 4.2 kbar, which was chosen based on the pressure of the peak field on the P–M O and P–M H₂O sections. Maximum H₂O values used on the x-axis of P–M H₂O and T–M H₂O sections were chosen to be 1.5 wt %. Compositions chosen along the x-axis from the peak fields of P–X and T–X diagrams for P–T modelling were selected based on modal proportions of minerals that best suited the estimated modes. Further information on the methods for the selection of compositions from these diagrams can be found in Appendix B.

Phase equilibria models were calculated for sample 14-RDC-04 using THERMOCALC 3.40 [43,44] in the chemical system MnO–Na₂O–CaO–K₂O–FeO–MgO–Al₂O₃–SiO₂–H₂O–TiO₂–O (MnNCKFMASTO), where “O” is a proxy for Fe₂O₃, using the internally-consistent thermodynamic dataset ds62’ [43] and activity–composition models reparametrised for Mn [8–10]. Completed pseudosections were contoured for the modal proportions of minerals using...
TCInvestigator [45]. The chemical composition of the discontinuous orthopyroxene-bearing pelitic layers (sample 14-RDC-04, see Figure 4) was used to constrain the peak metamorphic conditions at Reedy Creek.

Figure 4. (a) Calculated $P$–$T$ pseudosection using the composition of the residual orthopyroxene-bearing domain. The bulk composition (in mol%) is given above the pseudosection. The bold black line represents the solidus and the field outlined in yellow is the interpreted peak assemblage. (b) Overlay of modal contours on the peak field of Figure 4a. To avoid making the diagram overly complex, only the most abundant phases were included on the overlay. The contours of each mineral on the peak field of the $P$–$T$ section represent the modal abundances of the major phases in sample 14-RDC-04, including plagioclase, biotite, orthopyroxene and cordierite. Primary overlap of phases occurs between 2–4 kbar in the peak field. Individual modal proportion plots for each mineral can be found in Appendix B.

Melt reintegration was undertaken after calculation of the $P$–$T$ pseudosection using the original residual composition (Figure 4a) as a starting point, following the method of Korhonen, Brown, Clark and Bhattacharya [18]. This stepwise melt reintegration method is now well-established and has been used to remove melt or to rebuild protolith compositions in numerous studies e.g., [11,12,17–19]. This method adds 6 mol% melt to the system when the modal proportion of melt in the initial composition reaches 1 mol%, equivalent to the 7% melt connectivity transition (MCT) of [46]. This 7% threshold is interpreted as the point at which there is enough melt present in the rock to create a connected grain boundary network, resulting in the rapid decrease of rock strength and likely melt loss [47]. The composition of the melt that is reintegrated back into the rock corresponds to the composition of the 1% melt contour, which is interpreted to be the amount of melt remaining on grain boundaries after a melt loss event. The resulting composition is then normalised and can be used to calculate further $P$–$T$ pseudosections until the composition produces a wet solidus. It is assumed that as a rock crosses the solidus any remaining free $H_2O$ will be immediately used up during production
of the first vestiges of melt. Therefore, melt reintegration modelling aims to reintegrate melt until the solidus is just H$_2$O-saturated, to model the point at which a subsolidus composition would begin to melt. The $P$–$T$ path during the Delamerian Orogeny is not well constrained. Therefore, for the purpose of simplifying melt reintegration modelling, an isobaric heating path of 3.5 kbar was assumed, based on the peak pressures inferred from the $P$–$T$ pseudosection, the presence of andalusite pseudomorphs in the Reedy Creek area and the metamorphic pressures that are recorded elsewhere in the Delamerian Orogen e.g., [35,47,48].

4. Results

4.1. Peak P–T Conditions

The peak assemblage in sample 14-RDC-04 of plagioclase + biotite + orthopyroxene + cordierite + spinel + magnetite + ilmenite occurs at conditions of 1.0–4.4 kbar and 790–860 °C (Figure 4a). Mineral abundance contouring of this pseudosection indicates that peak $P$–$T$ conditions would have likely occurred between 2–4 kbar, where the modelled mineral modal abundance contours best correspond to the observed mineral abundances in thin section (Figure 4b). Observed abundances of minerals in thin section (determined through point counting) are provided in Table 1.

4.2. Melt Reintegration

Melt was reintegrated into the residual pelite composition in two stages (Figures 5 and 6), for a total of 12%. The addition of 12% melt results in a solidus that is water saturated at peak metamorphic conditions (1.0–4.4 kbar; Figure 6), with ~1.4 mol% H$_2$O present at 3.5 kbar. The addition of 12% melt results in a decrease in the temperature of the solidus at 3.5 kbar from ~810 °C to ~750 °C. After the reintegration of 12% melt, the peak assemblage above the solidus is still present and occurs at similar $P$–$T$ conditions, but subsolidus assemblages have been significantly changed. However, notably, subsolidus equilibria calculated using the H$_2$O content of the melt reintegrated composition do not contain andalusite (Figure 6a). The H$_2$O content of the melt reintegrated composition also does not allow for a free H$_2$O phase at temperatures below 570–670 °C (with increasing pressure). However, if H$_2$O is set to excess below the solidus, which is an approach commonly used for modelling subsolidus rocks, then mineral assemblages containing andalusite can be produced at temperatures below ~570 °C using the melt reintegrated composition (Figure 6b). Setting H$_2$O to be in excess also supresses the modelled stability of orthopyroxene.
Figure 5. Calculated $P$–$T$ pseudosection after the first melt reintegration (6% melt reintegrated). The bulk composition (in mol%) was calculated with the addition of 6% melt into the original granulite composition (Figure 4a) and is given above the pseudosection. The bold black line represents the solidus.

Figure 6. Calculated $P$–$T$ pseudosections after the second melt reintegration (12% melt reintegrated). The bulk composition (in mol%) was calculated with the addition of a further 6% melt into the composition after the first melt reintegration (Figure 4a) and is given above each pseudosection. The bold black line represents the solidus. (a) $H_2O$ was not set to excess below the solidus, instead subsolidus equilibria were calculated using the $H_2O$ content of the melt reintegrated composition. (b) $H_2O$ was set to excess below the solidus. Setting $H_2O$ to excess allows calculation of equilibria with a free water phase at subsolidus temperatures and results in significant changes to the stability of andalusite.
5. Discussion

The aim of this study is to test the veracity of melt reintegration calculations by reintegrating melt into a compositionally residual rock whose precursor mineralogy is partly known. We interpret andalusite to have been present during the subsolidus prograde evolution of the rock, based on blocky mineral aggregates cored by sillimanite that are rimmed by cordierite–spinel intergrowths, as well as the widespread presence of andalusite in lower grade stratigraphic equivalents. Sillimanite cores in the cordierite–spinel coronae are rare throughout the rock and were likely no more than metastable relics of the andalusite porphyroblasts that were unable to be completely replaced during metamorphism to higher temperatures due to sluggish diffusion of Al e.g., [49,50]. However, they do indicate that blocky domains of cordierite–spinel symplectites likely represent the complete replacement of andalusite porphyroblasts along the prograde path. As such, the localised occurrence of spinel–cordierite symplectites in the sample used in this study can be used to estimate the abundance of subsolidus andalusite. On this basis approximately 5% andalusite was present, allowing for volume loss associated with melt loss in the subsolidus protolith.

The $P$–$T$ pseudosection calculated for the original residual composition already has stable andalusite prior to any melt reintegration (Figure 7a). However, it is well established that residual bulk compositions are not appropriate for modelling the prograde evolution of a rock [12,18,51]. The melting and melt loss that has occurred in the field area would have altered the composition of the rocks so that the subsolidus part of the pseudosection does not represent real prograde assemblages. Therefore, we have used the presence of ~5% subsolidus andalusite as a target to evaluate the viability of the melt reintegration procedure for predicting subsolidus assemblages.

![Figure 7a](https://example.com/figure7a.png) ![Figure 7b](https://example.com/figure7b.png)

**Figure 7.** (a) TClInvestigator [45] plot of the modal proportion of andalusite present in the residual $P$–$T$ pseudosection (Figure 4a) prior to melt reintegration. (b) TClInvestigator [45] plot of the modal proportion of andalusite present in the ds55 $P$–$T$ pseudosection after 12% melt reintegration and with H$_2$O in excess.

Melt was reintegrated into the original residual composition in two separate batches of 6%, for a total of 12% melt until the solidus was H$_2$O saturated at peak conditions (1.0–4.4 kbar, and up to pressures of 6 kbar). The solidus in the final melt reintegrated bulk composition remains at a high temperature (750 °C at 3.5 kbar), which likely reflects the absence of quartz above 650 °C. It is common practice for petrologists to model subsolidus equilibria with H$_2$O in excess. Evidence of decreasing H$_2$O content with increasing metamorphic grade at subsolidus conditions can be found in most metamorphic terranes, suggesting that H$_2$O is typically lost from the system during prograde dehydration reactions, prior to melting [52–55]. Therefore, when the rock reaches the solidus, it is likely to contain very little free H$_2$O, and less H$_2$O...
than was present during the development of the lower temperature mineral assemblages [56]. Subsolidus andalusite could not be produced using the H$_2$O content of the final melt reintegrated composition except at very low temperatures; however, when H$_2$O was modelled to be in excess below the solidus, the reintegrated bulk composition was able to produce subsolidus assemblages that include andalusite (plagioclase–biotite–staurolite–chlorite–ilmenite–andalusite–quartz–H$_2$O, plagioclase–biotite–chlorite–ilmenite–andalusite–quartz–H$_2$O and plagioclase–biotite–cordsite–chlorite–ilmenite–andalusite–quartz–H$_2$O) with an andalusite abundance of 1–5%. The absence of andalusite in a fluid-undersaturated system suggests that H$_2$O plays a large role in its stability, at least using the dataset and activity models used in this study [8–10,43]. It should be noted that the use of different activity–composition models can result in differences in mineral stability [57]. Whereas modelling with subsolidus H$_2$O saturation arguably represents a closed system from which H$_2$O produced from dehydration reactions is unable to escape [56], a fluid-rich composition appears to be necessary to reproduce the subsolidus andalusite that we interpret was present. Therefore, to model the subsolidus andalusite-bearing mineral assemblages melt was reintegrated until a wet solidus was reached and H$_2$O was then assumed to be in excess below the solidus.

5.1. Limitations of the Modelling

While the modelling of melt reintegrated composition predicts the presence of prograde andalusite, there are several limitations to the modelling that was undertaken in this study. Some limitations relate to the uncertainties associated with the internally consistent thermodynamic data [43] and the activity–composition models [8–10] used in THERMOCALC to calculate the mineral stability fields e.g., [10]. The simplifications that are necessary to model phase equilibria in naturally complex chemical systems are also a limiting factor. The model chemical system used in this study does not take into account minor components, such as chlorine and fluorine in biotite and ZnO and Cr$_2$O$_3$ in spinel, which are present in minor amounts, see Table 1, [58,59]. Although the effects of these minor components are unlikely to greatly influence the outcomes of the modelling, they are real and will slightly modify the positions of reactions in P–T space [5]. The uncertainties in determining the effective chemical composition that relates to the formation of the peak mineral assemblages are also a limitation. The composition obtained by whole-rock geochemistry was not appropriate to model to the development of the pelitic orthopyroxene-bearing domains, as these occur in thin, discontinuous layers within metapassammitic rock. Instead, a composition for the purposes of phase equilibria modelling was determined by combining mineral abundances and average chemical compositions of minerals. A two-dimensional domainal composition is necessarily limited, but for the purposes of modelling we have assumed that it is a realistic composition.

The melt reintigration modelling is limited by simplifications to the modelled P–T path. An isobaric P–T path at 3.5 kbar was chosen for melt reintigration modelling on the basis of peak pressures reached by other locations during the Delamerian Orogeny and the presence of andalusite in lower grade successions within the Kanmantoo Group. In all likelihood, the sample used in this study would have experienced a change in pressure during prograde metamorphism, driven by the 10–20% shortening recorded in the southern part of the Delamerian Orogen [60]. However, due to the steep inclination of the solidus and liquid modal abundance contours (e.g., Figure 4a), minor changes in the pressure during heating do not significantly change the results of this study.

Despite successfully recreating subsolidus assemblages with andalusite in the presence of excess H$_2$O after melt reintegration, a more critical issue is that the orthopyroxene-bearing subsolidus assemblages in the final model produced in this study (Figure 6b) are unlikely to have existed, as subsolidus orthopyroxene does not appear to occur in sub-granulite grade metapelitic rocks. If it is assumed that the model shown in Figure 6b represents the amphibolite facies prograde metamorphic assemblages prior to melting and melt loss, subsolidus orthopyroxene would have formed on the prograde path. Tilley [61] described the apparent occurrence of subsolidus orthopyroxene in metapelites in the Comrie aureole in the Scottish Highlands. However, later evidence of subtle
partial melting was found [62], suggesting that the orthopyroxene-bearing assemblages formed under suprasolidus conditions. Although some studies have recorded mineral assemblages with prograde subsolidus orthopyroxene e.g., [63,64], these situations generally involve infiltration of a low H$_2$O activity fluid (aH$_2$O), such as CO$_2$-rich fluids or highly concentrated brines, and occur in rocks that have experienced granulite facies temperatures. An example of this is the formation of fine-grained orthopyroxene–andalusite symplectites under low aH$_2$O conditions during retrogression of pelitic granulites, described by Ballèvre, et al. [65]. The sources of low aH$_2$O fluids could include magmatic dykes, volatile-rich carbonates, or tectonic structures such as fractures or shear zones that may act as channels for fluid to move over large distances [66–68]. However, such sources are not relevant to the prediction of orthopyroxene as shown in Figure 6b, since the modelling does not incorporate a low aH$_2$O fluid and therefore it is unlikely that subsolidus orthopyroxene would form in the abundances predicted by our models in prograde metapelites at amphibolite facies conditions.

Instead, we suggest that the appearance of subsolidus orthopyroxene in our models is an artefact created by the absence of an activity–composition model for orthoamphibole in ds62 [10]. This means that Mg and Al cannot be incorporated into orthoamphibole and instead stabilises other minerals, resulting in the formation of unrealistic assemblages. Orthopyroxene can form by the breakdown of orthoamphibole in K-poor protoliths [69]. Occurrences of orthoamphibole in andalusite-bearing metapelites are rare; however, one such occurrence is also within the Kanmantoo Group, in the Springton region, approximately 30 km north of Reedy Creek [70]. Although the Springton region rocks are thought to have been metasomatised, their final bulk compositions contain low proportions of K$_2$O and CaO, and also contain a high Na$_2$O/K$_2$O ratio [70], which is similar to the composition of the sample used in this study.

The modelling undertaken in this study used the most recent internally-consistent thermodynamic dataset ds62 [43]. However, our attempts to incorporate orthoamphibole into the ds62 modelling failed to reproduce any subsolidus equilibria that could be rationalised with the incoming solidus or mineral assemblages that could plausibly have existed at subsolidus conditions. Therefore, at this stage, we cannot incorporate orthoamphibole into the modelling with ds62 with any confidence. It should also be noted that low–P subsolidus equilibria involving porphyroblastic cordierite like in this study have been known to be problematic to model in ds62 e.g., [9,57,71]. Instead, the final composition produced after melt reintegration modelling (Figure 6b) was used to model the subsolidus assemblages in the previous THERMOCALC dataset ds55 [72], to investigate the effects of incorporating orthoamphibole into the modelling e.g., [69,73]. The following activity relationships were used for modelling in ds55: silicate melt and biotite [5]; chlorite [74,75]; cordierite [72]; plagioclase [76]; gedrite [73]; orthopyroxene, spinel and magnetite [77]; ilmenite [6]. Using ds55, the subsolidus assemblages were again modelled with H$_2$O in excess (Figure 8). While we acknowledge the apparent logic disconnect in using a melt reintegrated composition created using ds62 followed by modelling in ds55, no subsolidus orthopyroxene was found to be stable using ds55, and instead, orthoamphibole (gedrite)-bearing assemblages were modelled to be stable (Figure 8). Despite the differences between the ds62 and ds55 subsolidus models, the melt reintegrated composition in ds55 was able to produce a number of assemblages containing andalusite at similar conditions to the ds62 model (1.0–4.1 kbar and 400–550 °C; Figure 8) and in similar modal proportions (Figure 7b; 1–6%). Furthermore, although the melt reintegrated composition is specific to the sample we have used, the prediction of gedrite-bearing assemblages is consistent with the presence of orthoamphibole-bearing assemblages in lower grade areas of the Kanmantoo Group [70], suggesting that while ds62 is now widely used, ds55 still provides realistic predictions.
5.2. Implications of the Melt Reintegration Modelling

Phase equilibria modelling is a powerful tool for investigating the chemical changes that occur as a result of partial melting and melt loss during high-grade metamorphism. It is one of the few methods that allows investigation of the petrological and chemical evolution of deep-crustal rocks, as it provides an opportunity to define equilibrium relationships between melt and solid mineral species for particular bulk compositions at a given set of P–T conditions e.g., [13,14]. Several previous studies have utilised this technique to explore the effects of reintegrating melt back into lower-crustal rocks [16,18,19,41,78,79]. However, rarely do studies attempt to assess the veracity of the melt reintegration process to create valid subsolidus rock compositions. Reedy Creek presents an opportunity in which a partially known subsolidus mineral assemblage can be used as a reference point for the assessment of the outcome of melt reintegration modelling on a granulite facies rock.

The Reedy Creek metasedimentary rocks preserve intact relict, centimetre-scale sedimentary layering. This suggests that melt loss from the rock was likely to have been incremental and efficient, without the accumulation of large amounts of melt that would have resulted in catastrophic melt loss and disruption of the sedimentary layering [80,81]. The melt connectivity transition approximately 7%, [46] was used as a maximum for the melt loss that would have occurred in any one melt extraction.
event. However, we acknowledge that in the presence of syn-anatectic deformation, melt extraction may well occur at a lower threshold [5,18,81,82].

In the modelled sample, the sillimanite is not interpreted to have been part of the equilibrium mineral assemblage at peak metamorphic conditions, or likely at any stage throughout the sample’s prograde history. In most cases, the cordierite–spinel symplectites that are seen surrounding sillimanite on rare occasion have completely replaced sillimanite, indicating that it is not part of the peak assemblage. It cannot be said for certain that sillimanite was never stable throughout any of the rock’s history. However, the models in this study (all of which were derived from the composition of the peak mineral assemblage in the sample) are not able to form stable sillimanite at any point, suggesting that it may not have occurred at all on the prograde path. Instead, we suggest that the sillimanite that we observe in the sample is simply a compositional relic of incomplete continuous reactions involving the progressive formation of cordierite and orthoamphibole at the expense of andalusite, biotite and chlorite as the rock traversed the narrow set of fields between andalusite-bearing assemblages and orthoamphibole–cordierite-bearing assemblages. We suggest that residual metastable aluminosilicate in micro-compositional domains underwent a reaction that cannot be sequentially tracked across phase fields in our $P$–$T$ models. The calculation of $P$–$T$ models relies on the assumption that all phases are in equilibrium at any given point across the diagram. However, in reality, this is rarely the case. Reactions may not proceed to completion, leaving behind minerals that should have reacted out to play a role in new reactions and restricting the growth of new minerals. This results in further reactions that involve metastable minerals that are not reflected on $P$–$T$ diagrams e.g., [83–86]. This overstepping displaces the location of the boundary of a metamorphic reaction in $P$–$T$ space, resulting in a phase diagram that does not necessarily reflect the sequence of prograde reactions that occurred in the rock. In this study, we interpret the small relics of aluminosilicate remaining in the granulite to be metastable with respect to the modelled bulk compositions. However, they play a critical role in our analysis because they allow us to confidently interpret the presence of a subsolidus mineral. Furthermore, the spinel–cordierite symplectites form clearly delineated domains in the rock that allow us to make an estimate of the approximate amount of subsolidus andalusite.

The predicted occurrence of andalusite at subsolidus conditions indicates that melt reintegreation modelling is a viable way to reproduce protolith bulk rock compositions. However, it should be noted that in cases where low-grade mineral assemblages are unclear or textural evidence of compositional domains not preserved, there may still be a significant amount of uncertainty involved in selecting a potential prograde $P$–$T$ path.

An important application of melt loss modelling is the interpretation of inclusion assemblages. A large number of studies have used inclusion assemblages to provide information about prograde history, even in rocks that have undergone partial melting and melt loss e.g., [87–89]. While these inclusions may offer an insight into the existence of potential prograde minerals, interpreting prograde conditions based on modelling a bulk composition that existed prior to modification due to melt loss is illogical, and unlikely to result in meaningful constraints. Furthermore, a number of workers have also used the compositions of individual minerals to make inferences about prograde histories e.g., [90–92]. Where rocks have experienced high temperature evolutions, any such compositions will only be fortuitously the same as they were at the time of capture due to diffusion-driven compositional modification of the inclusion. However, melt reintegreation modelling does offer an avenue for the interpretation of inclusion assemblages providing a window onto the prograde conditions of rocks, despite extensive melt extraction compositional modification.

6. Conclusions

Phase equilibria modelling undertaken on the Kanmantoo sediments at Reedy Creek indicates that peak $P$–$T$ conditions of 790–860 °C and 1.0–4.4 kbar were reached. From these peak conditions, melt reintegreation modelling of the pelitic granulites was able to reproduce several viable subsolidus mineral assemblages containing andalusite, which was known to have existed in these rocks at lower
grade. This indicates that current melt reintegartion methods may have the potential to effectively recreate valid protolith bulk rock compositions, but do require some knowledge of the protolith to be certain. However, the absence of orthoamphibole in the current ds62 model does, in this case, restrict our ability to accurately predict many subsolidus reactions that may have occurred in this rock.

Supplementary Materials: The following are available online at www.mdpi.com/2076-3263/7/3/75/s1, Mineral compositions obtained on MicroProbe SXFive.

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Appendix A

Figure A1. (a) P–M-O model calculated to constrain the oxidation state of sample 14-RDC-04. The bold line represents the solidus. The field outlined in yellow represents the peak assemblage observed in the sample. The dashed line is the composition on the x-axis that was chosen for further modelling. (b) TCI output of modal proportion of orthopyroxene. The peak field is outlined in bold. (c) TCI output of modal proportion of cordierite. The peak field is outlined in bold.
Figure A2. (a) $P-M_{H_2O}$ model calculated to constrain the $H_2O$ content of sample 14-RDC-04. The bold line represents the solidus. The field outlined in yellow represents the peak assemblage observed in the sample. (b) $T-M_{H_2O}$ model calculated to constrain the $H_2O$ content of sample 14-RDC-04. The bold line represents the solidus. The field outlined in yellow represents the peak assemblage observed in the sample. The dashed line is the composition on the x-axis that was chosen for further modelling.

A $P-M_O$ model (Figure A1a) was calculated to constrain the oxidation state (O) of the composition of sample 14-RDC-01. The oxidation state was chosen to be to the left of the centre of the field primarily to increase the proportion of orthopyroxene (Figure A1b) and decrease the proportion of cordierite (Figure 1c) to better adhere to the estimated modal proportions of these minerals in the sample. A $P-M_{H_2O}$ section (Figure A2a) was originally calculated to constrain the $H_2O$ content of sample 14-RDC-04. However, $H_2O$ is not very sensitive to changes in $P$ as opposed to $T$, and at the constant temperature that was chosen (750 °C) only a $H_2O$-saturated solidus could be achieved, meaning that no melt reintegration could take place. A $T-M_{H_2O}$ section (Figure A2b) was then calculated to assess the sensitivity of $H_2O$ with changing temperature. The left part of the peak field was chosen to represent the $H_2O$ content of the sample, as it resulted in a composition that was not $H_2O$-saturated at the solidus and would allow for melt reintegration modelling.
Appendix B

Figure A3. Mineral modal abundance contour diagrams produced using TCInvestigator [45] for the major phases that occur in the peak assemblage of sample 14-RDC-04. These diagrams were used to construct Figure 4b in the main text.

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