Strengths and limitations of \textit{in situ} U-Pb titanite petrochronology in polymetamorphic rocks: An example from western Maine, USA.

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ABSTRACT

Titanite is a potentially powerful U-Pb petrochronometer that may record metamorphism, metasomatism, and deformation. Titanite may also incorporate significant inherited Pb, the correction for which may introduce inaccuracies and result in geologically ambiguous U-Pb dates. Here we present laser ablation inductively coupled mass spectrometry (LA-ICP-MS)-derived titanite U-Pb dates and trace element concentrations for two banded calc-silicate gneisses from south-central Maine, USA (SSP18-1A & -1B). Single spot common Pb-corrected dates range from 400 to 280 Ma with 12–20 Ma propagated 2SE. Titanite in sample SSP18-1B exhibit regular core-to-rim variations in texture, composition, and date. We identify four titanite populations: 1) 399 ± 5 Ma (95 % CL) low Y + HREE cores and mottled grains, 2) 372 ± 7 Ma high Y + REE mantles and cores, 3) 342 ± 6 Ma cores with high Y + REE and no Eu anomaly, and 4) 295 ± 6 Ma LREE-depleted rims. We interpret the increase in titanite Y + HREE between ca. 400 and ca. 372 Ma to constrain the timing of diopside fracturing and recrystallization and amphibole breakdown. Apparent Zr-in-titanite temperatures (803 ± 36 °C at 0.5 ± 0.2 GPa) and increased $X_{Di}$ suggest a thermal maximum at ca. 372 Ma. Population 3 domains dated to ca. 342 Ma exhibit no Eu anomaly and are observed only in compositional bands dominated by diopside (> 80 vol %), suggesting limited equilibrium between titanite and plagioclase. Finally, low LREE and high U/Th in Population 4 titanite date the formation of hydrous phases, such as allanite, during high $X_{H2O}$ fluid infiltration at ca. 295 Ma. In contrast to the well-defined date-composition-texture relationships observed for titanite from SSP18-1B, titanite grains from sample SSP18-1A exhibit complex zoning patterns and little correlation between texture, composition, and date. We hypothesize that the incorporation of variable amounts of radiogenic Pb from dissolved titanite into recrystallized domains resulted in mixed ages spanning 380–330 Ma. Although titanite may reliably record multiple phases of metamorphism, these data highlight the importance of considering U-Pb data along with chemical and textural data to screen for inherited radiogenic Pb.
1. INTRODUCTION

Titanite (Ca[Ti,Al,Fe$^{3+}$]SiO$_4$[O,OH,F,Cl]) is a common accessory phase and regularly participates in reactions with major phases (see review in Kohn, 2017). Similar to zircon and monazite, the incorporation of U and a variety of trace elements (e.g., Zr, Nb, Sr, Y+REE) make titanite a potentially useful petrochronometer (Frost et al., 2000; Kohn, 2017). In particular, Zr-in-titanite thermometry and TZARS (titanite with anorthite, (clino-)zoisite, rutile, and quartz) barometry allows for the estimation of both pressures ($P$) and temperatures ($T$) of metamorphism (Hayden et al., 2008; Kapp et al., 2009). However, the reactivity of titanite with major phases and fluids during metamorphism presents both benefits and drawbacks. Titanite U-Pb dates have been interpreted to reflect a variety of processes, including neocrystallization and growth (Castelli & Rubatto, 2002; Corfu, 1996; Kohn & Corrie, 2011; Rapa et al., 2017; Scott & St-Onge, 1995; Spencer et al., 2013; Stearns et al., 2015; 2016; Verts, Chamberlain, & Frost, 1995; Walters & Kohn, 2017), fluid driven recrystallization or alteration (Corfu, 1996; Garber et al., 2017; Holder & Hacker, 2019; Marsh & Smye, 2017), Pb volume diffusion and cooling (Cherniak, 1993; Kirkland et al., 2016; Mattinson, 1978; Mezger et al., 1991, 1993; Spear & Parish, 1996; Tucker et al., 1987), and deformation-induced recrystallization (Bonamici et al., 2015; Gordon et al., 2021; Papapavloou et al., 2018; Spencer et al., 2013; Timms et al., 2020). Additionally, disequilibrium between titanite and zircon (or quartz) may result in overestimated Zr-in-titanite temperatures (Cruz-Uribe et al., 2018; Johnson, Cottle, & Larson, 2020). Combined, these factors greatly complicate linking titanite zoning textures, compositions, and U-Pb dates to metamorphic, metasomatic, and tectonic processes.

Historically, titanite U-Pb geochronometry was interpreted to date cooling, as opposed to peak metamorphism, in high grade metamorphic terranes. Early studies of Pb diffusion rates in titanite proposed relatively low closure temperatures ($T_c$) of 450–600 °C based on natural samples (e.g., Heaman & Parrish, 1991; Gascoyne, 1986; Mattison, 1978; Mezger et al., 1993). Later, experimentally-derived Pb-in-titanite diffusion rates determined by Cherniak (1993) suggested a $T_c$ of 600 °C for a 100 µm titanite grain cooling at a rate of 10 °C/Myr (Kohn, 2017). Variations in morphology, color, and U-Pb ratios of titanite from contact aureoles were interpreted to reflect variable diffusive resetting of Pb isotope compositions (e.g., Mezger et al., 1991, 1993; Tucker et al., 1987). In contrast, studies by Corfu & Muir (1989), Corfu & Ayers (1991), Verts, Chamberlain, & Frost, (1995), and others suggested that variations in titanite U-Pb ratios may reflect multiple or protracted metamorphic and/or metasomatic events. A growing number of studies of natural titanite over the past decade suggest that volume diffusion rates of Pb are greatly overestimated (Gao et al., 2012; Garber et al., 2017; Hartnady et al., 2019; Holder & Hacker, 2019; Holder et al., 2019; Kohn & Corrie, 2011; Kohn, 2017; Smye et al., 2018; Spencer et al., 2013; Stearns et al., 2016; Walters & Kohn, 2017). Revised estimates for Pb diffusion coefficients range from $1 \times 10^{-20}$ to $1 \times 10^{-26}$ (m$^2$ s$^{-1}$) (Holder, Hacker, Seward, & Kylander-Clark, 2019; Kohn, 2017), equivalent to the experimentally derived diffusion rate for Sr in titanite (Cherniak, 1995). These data suggest
that volume diffusion is unlikely to disturb U-Pb dates in titanite metamorphosed at temperatures <850 °C at most metamorphic time scales.

In the absence of Pb diffusion, disturbance of titanite U-Pb is anticipated to occur through fluid- or deformation-induced recrystallization. Consequently, multi-generational titanite with both recrystallized and secondary domains are commonly observed (e.g., Castelli & Rubatto, 2002; Corfu, 1996; Corfu & Ayers, 1991; Corfu & Muir, 1989: Garber et al., 2017; Holder & Hacker, 2019; Johnson, Cottle, & Larson, 2020; Spencer et al., 2013; Tucker et al., 2004; Verts, Chamberlain, & Frost, 1995). Recrystallized zones are typically patchy to lobate and often oriented along fractures and cleavage planes, consistent with interface-coupled dissolution-reprecipitation reactions (ICDR) (Bonamici et al., 2015; Garber et al., 2017; Holder & Hacker, 2019; Marsh & Smye, 2017; Walters & Kohn, 2017). During ICDR, minerals in disequilibrium with a fluid will dissolve until the chemical potential of the solid assemblage and fluid phase are equal, followed by precipitation at the reaction front once the fluid has reached super-saturation (see review in Putnis, 2009). Alternatively, depolymerization of the precursor grain may form an intermediate amorphous phase which may then recrystallize (Konrad-Schmolke et al., 2018).

One of the pitfalls of titanite U-Pb geochronology is that significant Pb may be incorporated upon crystallization and a correction for the initial common Pb composition is often required (see discussions in Kirkland et al., 2018; Bomamici & Blum, 2020). Isotopic homogenization of Pb during metamorphism is required to obtain accurate U-Pb ages (Romer, 2001); however, Pb may be partially to wholly retained during ICDR, such that the initial \(^{207}\text{Pb}/^{206}\text{Pb}\) ratio of the recrystallized domains is a mix of common Pb and radiogenic Pb. Inherited radiogenic Pb has been identified in both neocrystallized titanite grains replacing precursor high-U phases (Kirkland et al., 2017, 2018; Romer & Rötzler, 2003), such as rutile, and in titanite recrystallized by ICDR (Castelli & Rubatto, 2002). Even phases that typically incorporate little initial Pb, such as monazite, may inherit old radiogenic Pb during ICDR recrystallization (e.g., Seydoux-Guillaume et al., 2003; Weinberg et al., 2020). Thus, the degree to which titanite dates reflect the timing of recrystallization requires careful consideration.

Here we present titanite textural observations coupled with in situ compositional and U-Pb isotopic data by electron probe microanalysis (EPMA) and laser-ablation inductively coupled mass-spectrometry (LA-ICP-MS) for two calc-silicate gneisses from western Maine, USA. In one sample (SSP18-1B), titanite textures, major and trace element compositions, and U-Pb dates strongly correlate. We identify four composition-age populations at ca. 400, 372, 342, and 295 Ma, three of which correspond to previously dated metamorphic and igneous activity in the area. Variations in major and trace element concentration between these populations are linked to different stages of high-temperature metamorphism, deformation, and fluid infiltration. In contrast, titanite analyses from a second sample (SSP18-1A) exhibit a decoupling of U-Pb dates from textures and compositions, indicative of inherited radiogenic Pb from precursor titanite domains. These two samples, collected at the same outcrop, illustrate both the strengths and weaknesses of U-Pb titanite petrochronology in polymetamorphic terranes.
2. GEOLOGIC SETTING

2.1. Regional Overview

The geology of Central and Western Maine is dominated by extensive metasedimentary sequences of the Central Maine Trough and Connecticut Valley-Gaspé Trough, which extend from Connecticut through Newfoundland (Fig. 1A) (Hibbard et al., 2006). In the Late Silurian to Early Devonian, these sediments were deposited into extensional basins within the Ganderian margin following accretion to Laurentia (e.g., Rankin et al., 2007). These sediments were subsequently metamorphosed during the Acadian (420–350 Ma) orogeny (Robinson et al., 1998; van Staal et al., 2009), resulting in a state-wide metamorphic gradient spanning from greenschist-facies rocks in northeastern Maine to a high-T migmatite terrane in southwestern Maine (Fig. 1A) (see review in Guidotti, 1989). Temperatures calculated by Holdaway et al. (1997) at 310 MPa range up to 626–649 °C in the highest-grade rocks. Holdaway, Dutrow, & Hinton (1988) noted a N-S increase in pressure, ranging from 235 ± 25 MPa for cordierite- and andalusite-bearing assemblages exposed in the northern lobe of the Lexington pluton to 400–500 MPa or possibly higher for the migmatitic rocks near Lewiston (Fig. 1A). Higher $P$ is supported by estimates of 550–640 MPa at 490–540 °C for kyanite-bearing rocks exposed just south of the Sebago pluton (S, Fig. 1A; Thomson & Guidotti, 1989).

Figure 1. (a) Map of Maine showing the distribution of metamorphic grades from sub-greenschist facies rocks in the north to migmatites in the southwest (after Guidotti, 1985; with updates from Solar & Brown, 1999). Major intrusions are highlighted, including the Lexington (L), Mooselookmeguntic (MIC), and Sebago (S) plutons. Also shown are the boundaries of the Connecticut Valley-Gaspé Trough (CVT) trough, Bronson Hill (BT) terrane, and Central Maine (CMT) trough. (b) Geologic map of the sample locality near Lewiston, Maine, and (c) Cross-section from A to A’ (after Hussey, 1983). Metasedimentary units include the Vassalboro (Vf), Waterville (Wf), Smalls Falls (Sff), and Sangerville formations. The latter is divided into pelitic (Sp), calc-silicate rich (Scs), and Taylor Pond (Stp) endmembers. Granite (G) and granodiorite (Gd) intrusions are observed north of the sample locality.
The complex multi-phase nature of the high-grade metamorphism has been intensely debated over the past half century (e.g., Holdaway, Dutrow, & Hinton, 1988; Guidotti, 1970; Lux & Guidotti, 1985; Solar & Brown, 1999; Solar & Tomascak, 2016). Early tectonometamorphic models for Maine describe high-grade regional metamorphism in the Acadian followed by pulses of contact metamorphism driven by pluton emplacement (see review in Guidotti, 1989). Indeed, the regional gradient is locally deflected around plutons (Fig., 1A). Additionally, pluton and dike crystallization ages range from ca. 420 Ma for coastal plutons to 270–260 Ma for abundant gem pegmatites (Bradley et al., 2000). Dates of ca. 400 Ma throughout western and central Maine and New Hampshire are consistent with large scale high-T Acadian metamorphism (Gerbi & West, 2007; Pyle et al., 2005; Moecher, McCulla, & Massey, 2020; Sanislav, 2011; Smith & Barreiro, 1990; Solar et al., 1998; Wing, Ferry, & Harrison, 2003). In the region surrounding the Mooselookmeguntic Igneous Complex (Fig. 1A), Acadian metamorphic conditions are thought to have exceeded 450 MPa and 700 °C in the migmatites, resulting in syntectonic melt generation, percolative flow, and emplacement of granite magmas (Brown & Solar, 1999; Johnson, Brown, & Solar, 2003).

The timing and conditions of high-grade metamorphism to the south in the vicinity of the Sebago pluton are more enigmatic. Emplacement of the Sebago ‘batholith’ was long considered responsible for much of the late high-grade metamorphism in western Maine (Fig. 1A; see review in Guidotti, 1989). However, gravimetric and field mapping has revealed a much smaller Sebago pluton that sits at the core of an extensive migmatite terrane intruded by numerous smaller granitic bodies (Behn, Eusden, & Notte, 1998; Solar & Tomascak, 2016). Tomascak, Krogstad, & Walker (1996) reported a U-Pb monazite thermal ionization mass spectrometry age of 293 ± 2 Ma for crystallization of the Sebago pluton. The crystallization ages of smaller granite and pegmatite bodies in the vicinity of the Sebago pluton range from 308 to 269 Ma (Solar & Tomascak, 2016; Tomascak, Krogstad, & Walker, 1996). These ages contrast with a pooled zircon rim age of 376 ± 14 Ma collected from three migmatite melanosomes (Tomascak & Solar, 2016), suggesting that the emplacement of the Sebago pluton is not responsible for partial melting in the surrounding region. Proposed mechanisms for metamorphism and melting in broader New England invoke the development of a thick crust and extended topographic high between ca. 400 and ca. 380 Ma (Hillenbrand et al., 2021), with thermal highs between ca. 380 and ca. 330 Ma that may have been generated by lithospheric delamination and asthenospheric upwelling (e.g., Pyle et al., 2005; Moecher, McCulla, & Massey, 2020; Robinson et al., 1998). The extent that regional, as opposed to contact, metamorphism is reflected in western Maine remains blurred by the murky temporal relationship between pluton emplacement, melting, and high-T metamorphism. Here we use titanite petrochronology to investigate the timing of high temperature metamorphism in west-central Maine and distinguish between thermal sources.

2.2 Sample Locality

Three samples were collected in Lewiston, Maine, from the Sangerville formation, which in this area comprises three lithologic units: 1. Migmatic paragneiss with the dominant assemblage
Bt + Ms + Qtz + Pl ± Sill ± Grt ± Kfs (Sp, Fig. 1b & S1), 2. The Taylor Pond endmember paragneiss with the assemblage Bt + Qtz + Pl ± Hbl and thin interbeds of calc-silicate (Stp), and 3. Banded calc-silicate and metapsammitic gneiss (Scs, Fig. S1c; Hussey, 1983). Two samples were collected from the calcareous bands (SSP18-1A & -1B) and a third was collected from a rare garnet-bearing horizon in the paragneiss (SS18-1D, Appendix S1). All abbreviations are after Whitney & Evans (2010). Two larger intrusive bodies are observed to the northwest of the sample locality (Fig. 1b). The larger two-mica granite pluton (G) exhibits sharp and discordant contacts with the metasediments and no obvious contact metamorphism (Hussey, 1983). Although the granite is undeformed, the fine to medium-grained granodiorite body (Gd) is moderately to strongly foliated (Hussey, 1983). The outcrop includes at least two dike populations. The first population is parallel to the foliation of the migmatites and boudinaged (Fig. S1a & b), whereas the second population crosscuts the foliation and exhibits sharp contacts with the host rock (Fig. S1c & d). Pegmatite dikes are also common in the field area but are not observed in the sampled outcrop (Hussey, 1983).

3. ANALYTICAL METHODS

3.1 Major and minor element analysis

Major and minor element data on silicate phases were collected through electron probe microanalysis (EPMA) using a Cameca SX-100 electron microprobe equipped with five wavelength dispersive spectrometers, housed at the University of Maine. Standard spot analyses utilized a beam size of 5 µm, accelerating voltage of 15 keV, and 10 nA beam current. On-peak counting times were 20 s for all elements, except for Na, where counting time was 10 s. Backgrounds were measured for half the on-peak counting time. Data are reported in Table S1 as the mean and 2SE of multiple analyses collected within compositionally homogenous domains. Ferric iron in garnet and clinopyroxene was determined using method #2 of Schumacher (1991), in which all Fe is considered Fe³⁺ in titanite and epidote-group minerals. Amphibole structural formula calculations follow Leake et al. (1997) and classified following Hawthorne et al. (2012). Structural formulae of other minerals were determined by assuming a total number of cations or anions where appropriate and the methods listed are given in Appendix S1.

Qualitative and quantitative compositional maps were also collected by EPMA. Qualitative garnet (Fig. S2) and full thin section (Fig. S3) X-ray intensity maps were collected using a 20 keV accelerating voltage, 100 nA beam current, and a dwell time of 30 ms. Single- and multi-phase quantitative maps were collected using at a 15 keV accelerating voltage, 100 nA beam current, and 300 ms dwell time with a focused beam and a step size of 2 µm. X-ray intensity maps were quantified using 15–36 spot analyses per phase and the program XMapTools (Lanari et al., 2014; 2018).

3.2 Titanite U-Pb geochronology and trace element analysis

In situ titanite U-Pb and trace element analyses were conducted on polished thin sections by laser-ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) in the the
Analyses of unknowns were bracketed by those of titanite reference materials MKED1 (1518.87 ± 0.32 Ma; Spandler et al., 2016), BLR-1 (1047.1 ± 0.4 Ma; Aleinikoff et al., 2007), and McClure Mountain (523.26 ± 0.65 Ma; Schoene & Bowring, 2006). Analyses of NIST SRM610, NIST SRM612, and GSD-1G glasses (Pearce et al., 1997; Jochum et al., 2011) were conducted for trace element determination. MKED1 was used as the primary reference material for U-Pb isotope ratios and Ca, Ti, and Zr mass fractions, whereas NIST610 was used as the primary reference material for all other trace elements. All titanite U-Pb isotope and trace element data for unknowns are reported in Tables S2 and S3, respectively. Single spot dates are given with uncertainties reported as 2SE, whereas the uncertainties of pooled dates are reported as the 95 % confidence interval corrected for overdispersion (Vermeesch, 2018).

Measured \(^{207}\)Pb/\(^{206}\)Pb ratios of clinopyroxene from SSP18-1A and -1B are 0.9654 ± 0.0393 (2SE, n = 5) and 0.9554 ± 0.0495 (n = 10), respectively. In contrast, feldspar \(^{207}\)Pb/\(^{206}\)Pb ratios are 0.8730 ± 0.0064 (n = 15) and 0.8781 ± 0.0061 (n = 10), respectively. Common Pb isotope analyses are reported in Table S4. We estimate that 2–20 % of the measured Pb is inherited ‘common’ unradiogenic Pb (\(f^{207}\)Pb, Table S2). As a result, dates that are uncorrected, corrected using the two-stage crustal evolution model of Stacey & Kramers (1975), and corrected with measured feldspar or pyroxene \(^{207}\)Pb/\(^{206}\)Pb ratios are equal within their respective uncertainties. Given the higher Pb concentrations in feldspar relative to clinopyroxene and smaller scatter we calculated all dates using the feldspar Pb ratios. We discuss potential systematic inaccuracies introduced by our preferred method of common Pb correction in section 7.2.
The measured dates and trace element concentrations for secondary reference materials are consistent with published values and are reported in Tables S5 and S6, respectively. Weighted mean ages of 1044 ± 14 Ma (n = 20, MSWD = 0.71) and 1065 ± 23 Ma (n = 16, MSWD = 2.40) for BLR-1 calculated for our two sessions are within uncertainty of the published dates. These data suggest that ages are both precise and true relative to reference values within ± 1–2% uncertainty (95% CI). Additional dates of 516 ± 37 (n = 10, MSWD = 2.38) and 537 ± 16 (n = 10, MSWD = 0.83) for the McClure Mountain titanite are also consistent with the published age. We find that the trace element concentrations of NIST SRM612 and GSD-1G are within 3 and 14%, respectively, of reference values from Jochum et al. (2011). Zr concentrations exhibit a larger bias (6 and 17%, respectively); however, MKED1 titanite was used as a reference material for Zr and the bias is likely due to matrix differences. Our trace element concentrations for BLR are less consistent with published values (Table S6). For example, our Zr concentrations are ~270 µg g⁻¹ higher than those reported by Mazdab (2009). It is likely that these differences represent trace element heterogeneity in the BLR-1 titanite.

3.3 Titanite LA-ICP-MS Mapping

Trace element maps of a titanite grain from SSP18-1A were also collected at the University of Maine. Maps were collected using fast washout tubing and an ESI dual concentric injector ICP-MS torch. Analytical conditions consisted of a square 7x7 µm spot, 88 µm s⁻¹ raster rate, 5 J cm⁻² beam energy density, 200 Hz repetition rate, and a He carrier gas flow rate of 1100 mL/min. Monitored isotopes and dwell times (ms) were ²⁸Si (1), ⁴⁹Ti (1), ⁸⁹Y (1), ⁹⁰Zr (1), ²⁰²Pb (20), ²⁰⁷Pb (30), and ²³⁸U (10). The total ICP-MS duty cycle of 80 ms was synchronized to the laser pulse width following Van Malderen et al. (2018) and van Elteren et al. (2019) to achieve pixel dimensions of 7x7 µm. Two lines of BLR titanite were rastered at the same conditions before and after the unknown titanite grain. The time-resolved signals were processed using the VizualAge UComPbine and Trace Elements data reduction schemes in Iolite (v4) using ²⁸Si (14.2 wt. %) as the internal standardization element for mass fraction calculations. Trace element maps were produced using CellSpace in Iolite4 (Paton et al., 2011; Paul et al., 2012; Woodhead et al., 2007). Final maps were reprocessed in XMapTools (Lanari et al., 2014; 2018; Raimondo et al., 2017).

3.4 Major mineral LA-ICP-MS analysis

In addition to titanite, minor and trace element analyses (Session 3) were conducted on diopside, amphibole, epidote group minerals, plagioclase, and alkali feldspar and are reported in Table S7. Analyses of diopside, amphibole, and epidote group minerals were conducted with a spot diameter of 30 or 40 µm, a ~3 J cm⁻² beam energy density, and a repetition rate of 8 Hz, whereas feldspar analyses were conducted with a 40 µm spot diameter, a 6.7 J cm⁻² beam energy density, and a repetition rate of 8 Hz. A gas flow rate of 800 mL/min of He was used, and a ‘Squid’-type signal smoother was employed. Measured isotopes and dwell times (ms) were ²⁷Al (3), ²⁹Si (3), ⁴⁴Ca (3), ⁴⁹Ti (30), ⁵⁶Fe (20), ⁵⁸Sr (14), ⁸⁹Y (5), ⁹⁰Zr (10), ⁹³Nb (10), ¹¹⁷Ba (20), ¹³⁹La (20), ¹⁴⁰Ce (15), ¹⁴¹Pr (20), ¹⁴⁶Nd (20), ¹⁴⁷Sm (20), ¹⁵³Eu (20), ¹⁵⁷Gd (20), ¹⁵⁹Tb (20), ¹⁶³Dy (20), ¹⁶⁵Ho (20), ¹⁶⁶Er (20), ¹⁶⁹Tm (20), ¹⁷²Yb (20), ¹⁷⁵Lu (20), ¹⁷⁸Hf (20), ¹⁸¹Ta (20), and ²⁰⁸Pb (10). All
analyses consisted of 40 s ablation with 15 s background collection during laser warm up and 10 s of washout on the ICP-MS. Unknown analyses were bracketed with analyses NIST SRM610, NIST SRM612, and GSD1G reference glasses. We used NIST SRM610 as the primary reference material and the data were reduced using Iolite (v4). Silicon contents measured by EPMA were used as the internal standards and are reported in Table S7. Trace element concentrations for plagioclase and amphibole were calculated using compositional extremes. Major mineral LA-ICP-MS results are summarized in Table S7.

3.5 Electron backscatter diffraction analysis

For electron backscatter diffraction (EBSD) analysis of titanite grains in Sample SSP18-1A, the sample was polished mechanically with 0.3 μm alumina suspension and chemically in 0.02 μm colloidal silica suspension, and then was coated with a thin layer of carbon to prevent electron charging. EBSD patterns of titanites were collected in the Tescan Vega II Scanning Electron Microscope equipped with an EDAX-TSL EBSD system at the University of Maine. Operating conditions for the EBSD analysis were an acceleration voltage of 20 kV, a working distance of 25 mm, a beam current of ~6 nA, a 70° sample tilt, high-vacuum conditions, and a step size of 2 μm.

Collected EBSD patterns were indexed using the crystallography data of a natural titanite sample (E2312; Hawthorne et al., 1991). EDAX-TSL OIM Analysis 6 software was used to post-process titanite EBSD data based on confidence index (CI) and neighboring orientations. Non- and poor-indexed pixels (CI < 0.02) were replaced with well-indexed neighboring pixels of CI ≥0.02. The indexing rate is more than 99.8 %. Relative misorientation maps display the minimum misorientation relative to a reference orientation (see Figs. S7–8).

4. PETROGRAPHY AND MAJOR ELEMENT CHEMISTRY

Below we describe the mineral assemblages, textures, and major element compositions for calc-silicate samples SSP18-1A and -1B and infer possible reactions. Additional petrographic description is given for the metapelitic sample (SSP18-1D) in Appendix S1, along with photomicrographs (Fig. S9) and garnet EPMA transects (Fig. S10).

4.1 Calc-silicate sample SSP18-1A

Sample SSP18-1A is a gneiss with alternating calcisilicate and metapsammitic bands comprising the peak mineral assemblages Di + Kfs + Pl + Ttn + Ap and Bt + Pl, respectively (Fig. 2a–f). Minor and trace minerals in the calcisilicate band include pyrrhotite, calcite, and zircon, whereas only zircon is present in the psammitic bands. Plagioclase (An40–46), alkali feldspar, diopside, titanite, and apatite in the calcisilicate band exhibit a granular texture (Fig. 2a–b), consistent with equilibrium. Major mineral compositions are summarized in Figure S11. Backscattered electron (BSE) images of alkali feldspar reveal intersecting linear high Ba zones (up to 1 wt. %, Fig. 2c). Diopside grains exhibit slight undulatory and patchy Fe-Mg zoning in BSE (Fig. 2e). Calcite is rare (<< 1 vol %) and quartz is not observed. The absence of quartz and rarity calcite in the peak assemblage is consistent with the reaction proposed by Ferry (1976):
Figure 2. Plane polarized light (a), cross-polarized light (g-h), and backscattered electron (BSE, b-f, d) images of minerals and textures in sample SSP18-1A and -1B. SSP18-1A: (a) Large titanite grains adjacent to diopside, plagioclase, and alkali feldspar, (b) Granular texture of plagioclase, alkali feldspar, and diopside adjacent to titanite, with small grains of clinozoisite and pyrrhotite, (c) High-Ba BSE-bright linear features observed in alkali feldspar, (d) A patch of intergrown clinozoisite, low-An Pl, and calcite adjacent to high-An plagioclase, (e) Calcic-amphibole and complex lobate zoning in diopside, and (f) prehnite replacing biotite in a metapsammitic band. SSP18-1B: (g) Complex and sharply interlocking plagioclase grains are observed along with diopside and plagioclase, (h) oscillatory zones high-An plagioclase is cut by veins of low-An plagioclase, and (d) clinozoisite with allanite cores and rare quartz overprint earlier formed diopside, titanite, and plagioclase.

(1) Qz + Cal + Tr = Di + CO₂ + H₂O

While alkali feldspar is abundant (Fig. S3), we observe no evidence of precursor K-bearing phases, such as biotite, in the calcisilicate band itself.

Retrograde minerals are observed only locally and make up <1–2 vol % or less of the sample. First, a second phase of An₀₆ plagioclase and clinozoisite replace An₄₀–₄₆ plagioclase and...
calcite (Fig. 3D). Second, diopside grains are rimmed by calcic amphibole, which range in composition from Mg-hornblende to actinolite (Fig. 2e & S6a–b). These textures are consistent with the following reactions in the CaO–MgO–Al₂O₃–SiO₂–H₂O–CO₂ system:

(2) \( \text{An} + \text{Cal} + \text{H}_2\text{O} = \text{Czo} + \text{CO}_2 \)

(3) \( \text{Di} + \text{En} + \text{Qz} + \text{H}_2\text{O} = \text{Tr} \)

In particular, retrogression via Reactions 2 and 3 were likely limited by the rarity of calcite and quartz. In the psammitic bands, biotite grains are partially replaced by prehnite (Fig. 2f). No clear Mg- or Fe-rich product phase is observed, and possible reactions require mobilization of these components into a fluid phase.

Titanite (~10 vol %) in the calcisilicate band occur as large (≤ 2.5 mm) grains. Misorientation maps show up to 10° of internal deformation that sweep gradationally across grains and larger misorientations occur associated with fractures, grain boundaries, and twinning (Figs. S7–S8). Misorientations are not correlated with compositional zoning in BSE (Figs. S4–S5). Titanite BSE images reveal a complex patchwork of overlapping zones with sharp to gradational boundaries (Figs. S4–S5). Sector zones, with their characteristic fir-tree shape, and cyclical zones are observed in some grains (Fig. 3a). Titanite endmember fractions range from Ttn₉₂ to Ttn₉₄, with considerable variation in Al, Fe³⁺, and F shown in major element compositional maps (Fig. 3b–c). Aluminum and Fe³⁺ are positively correlated within grain interiors, whereas BSE dark rims exhibit elevated Al + F and low Fe³⁺. The observed zoning is thus consistent with the substitution (Al,
Fe$^{3+}$ + (OH, F) = Ti$^{3+}$ + O$^{2-}$ (Kohn, 2017). Possible titanite forming reactions proposed by Frost et al. (2000) and Rapa et al. (2017) are:

(4) Qz + Rt + Cal = Ttn + CO$_2$,
(5) Ti-Phl + Cal + Qz = Kfs + Tr + Ttn + H$_2$O + CO$_2$,
(6) Ti-Phl + Cal + Qz = Kfs + Di + Ttn + H$_2$O + CO$_2$.

Additionally, if $X_{CO2}$ was sufficiently low during prograde metamorphism, the reaction
(7) Qz + Rt + Zo = Ttn + An,
may have occurred. Other phases, such as amphibole and epidote group minerals, may contain significant Ti (Frei et al., 2004; Hawthorne et al., 2012) and their breakdown during prograde metamorphism may also promote titanite formation. However, the lack of prograde reaction textures and inclusions in titanite preclude clear petrographic identification of initial titanite forming reactions.

4.2 Calcsilicate Sample SSP18-1B

Sample SSP18-1B is a gneiss with alternating diopside- or plagioclase-rich bands (Fig. S12). The peak assemblage Di + Kfs + Pl + Cal + Ttn is observed (Fig. 2g-i & 4). Minor and trace phases include pyrrhotite and zircon. Calcite is rare, making up <10 vol % of plagioclase-rich bands. Plagioclase grains exhibit irregular interlocking grain boundaries with sharp spire-like intergrowths (Fig. 2g) and more rarely cyclical zones (Fig. 2h). These grains are often fractured and infilled by lower An plagioclase. Compositional maps and spot analyses reveal a range from An$_{28}$ to An$_{80}$ (Figs. 4d & S11). Diopside grains exhibit intense fracturing and rehealing textures: Early diopside exhibits the composition Di$_{53-54}$ which increases to Di$_{58-70}$ in the later fracture-fill clinopyroxene (Fig. 4b). Rare amphibole inclusions are observed in larger diopside grains, consistent with Reaction 1. Alkali feldspar is observed in fractured regions of diopside (Fig. 4), suggesting the reaction
(8) Phl + Cal + Qz = Tr + Kfs + CO$_2$ + H$_2$O,
which may have occurred following the fracturing of the early diopside grains. However, no inclusions of biotite or relict matrix grains are observed.

Extensive amphibole and clinzoisite retrogression are observed in sample SS18-1B (Fig. 2h). Amphibole compositions range from Mg-hornblende to actinolite from core to rim. Allanite occurs as cores in clinzoisite, suggesting LREE depletion from the reactive bulk composition as retrogression progressed (Fig. 2i). Epidote compositions range from Ep$_{29}$ to Ep$_{60}$, with higher Fe contents adjacent to diopside (Fig. 4c). In some areas, clinzoisite is associated with quartz (Fig. 2i), suggesting the hydration reaction
(9) Di + An + H$_2$O = Qz + Czo + Tr.

Elsewhere, clinzoisite has nucleated adjacent to calcite, consistent with the decarbonation-hydration Reaction 2 (Fig. 4a).
Titanite grains (1–2 vol %) in sample SSP18-1B are relatively small (<800 µm) and lenticular in habit. Titanite grains are observed primarily in the matrix and as rare inclusions in diopside. The interiors of some rare grains exhibit oscillatory zoning or patchy cores, both with and without sectors (Fig. S6). However, most grains exhibit regular core to rim zoning, with BSE dark cores observed in some grains, followed by a BSE bright mantle and a BSE dark rim (Fig. S6). Many grains exhibit only bright cores and dark rims. Titanite compositions range from Ttn$_{94}$ to in the cores to Ttn$_{87}$ in the rims.

5. TRACE ELEMENT CHEMISTRY

Feldspar and diopside are the dominant peak metamorphic phases in our samples. Analyses of plagioclase reveal relatively high concentrations of Sr (160–1684 µg g$^{-1}$), Ba (17.7–944 µg g$^{-1}$), Fe (46.8–1476 µg g$^{-1}$), and Pb (0.291–23.9 µg g$^{-1}$). Plagioclase exhibits very low Eu concentrations (0.001–0.570 µg g$^{-1}$) but displays a positive Eu anomaly relative to Sm and Gd, which are below detection. Alkali feldspar in SSP18-1A exhibits elevated Ba (5296–7064 µg g$^{-1}$), Sr (790–1054 µg g$^{-1}$), Pb (33.4–66.7 µg g$^{-1}$), and Fe (18.4–40.0 µg g$^{-1}$), with a positive Eu
anomaly. Diopside grains have elevated Ti (9.5–212 µg g\(^{-1}\)) and Sr (0.86–95.6 µg g\(^{-1}\)), relative to other trace elements, and exhibits high Y + HREE relative to LREE. In Sample SSP18-1B, Ti, Y (+HREE), and Sr concentrations decrease between early fractured high \(X_{\text{Di}}\) and healed low \(X_{\text{Di}}\) diopside domains (Figs. 4 & 5).

Texturally late amphibole and epidote group minerals were also analyzed. Amphibole compositions range between Ti (730–2635 µg g\(^{-1}\)), Sr (8.84–37.8 µg g\(^{-1}\)), and Y (9.73–68.9 µg g\(^{-1}\)) and are depleted in LREE relative to HREE. Analyses of epidote and clinozoisite in Sample SSP18-1B reveal high Sr (690–1595 µg g\(^{-1}\)) and Pb (30.8–138 µg g\(^{-1}\)) relative to other trace elements, elevated LREE relative to HREE, and a strong positive Eu anomaly (\(\text{Eu}/\text{Eu}^* = 2.1–9.7\)). Analysis of an allanite core in epidote revealed high LREE (e.g., Ce ~ 37 wt %), Th (2024 µg g\(^{-1}\)), U (487 µg g\(^{-1}\)), and Ti (1288 µg g\(^{-1}\)), with sharply decreasing abundance from the allanite core through the epidote mantle and rim.

Titanite grains exhibit a wide range of compositions. For example, Fe, Y, and Sr concentrations range from 1227 to 2526 µg g\(^{-1}\), 94 to 5175 µg g\(^{-1}\), and 25.8 to 156 µg g\(^{-1}\), respectively (Fig. 6). Rare Earth element patterns vary considerably between and within samples. In SSP18-1B, titanite REE patterns range from steep MREE to HREE slope with high LREE to analyses with flat MREE to HREE and low LREE (Fig. 7a). Some domains exhibit no Eu anomaly whereas others exhibit a strong negative Eu anomaly. In contrast, sample SSP18-1A REE patterns range from relatively flat with strong negative Eu anomalies to MREE depleted patterns with positive Eu anomalies (Fig. 7b). The correlation of these trace element patterns with age are discussed in Section 7.

6. ZR-IN-TITANITE THERMOMETRY

Zr-in-titanite temperatures were calculated for samples SS18-1A and -1B using the formulation of Thomas et al. (2010). Temperatures were calculated at 500 MPa with an associated 2\(s\) uncertainty of ± 200 MPa, corresponding to the 300–700 MPa range predicted here of possible pressures proposed for various stages of metamorphism in western Maine (Holdaway, Dutrow, &
Figure 6. LA-ICP-MS analyses of titanite showing concentrations of (a) Fe, (b) Sr, (c) Y, (d) Th/Pb, and (e) U/Th plotted relative to apparent age, as well as (f) Ce concentrations plotted against Sr. Data for sample SSP18-1B are color coded by Population: 1) yellow, 2) green, 3) blue, and 4) purple, whereas data for sample SSP18-1A are given in grey. Arrows indicate trends with time for sample SS18-1B. Data plotted in f are labeled according to titanite location (diopside-rich or plagioclase-rich bands) and texture (BSE dark rims).

Hinton, 1988; Holdaway et al., 1997; Thomson & Guidotti, 1989; this study). The $a_{\text{TiO}_2}$ of titanite-bearing rutile-absent metamorphic rocks is thought to be >0.5 (Chambers & Kohn, 2012, Ghent & Stout, 1984); therefore, we assume a value of 0.75 ± 0.25 (2s). Temperatures were also corrected for $a_{\text{CaTiSiO}_5}$ using an ideal activity model and our measured $X_{\text{Ttn}}$ and 2s uncertainties of 0.93 ± 0.01 and 0.91 ± 0.05 for samples SSP18-1A and -1B, respectively. All calculations assume $a_{\text{SiO}_2}$ and $a_{\text{ZrSiO}_4}$ of unity. Zircon grains are rare in the calc-silicate band of SSP18-1A, whereas zircon inclusions in titanite are common in SSP18-1B (Fig. 2d & S6). Quartz is only observed in SSP18-1B associated with texturally late clinozoisite and sample SSP18-1A is quartz-absent; therefore, the majority of our Zr-in-titanite temperatures for SSP18-1A and those calculated from interior
TITANITE PETROCHRONOLOGY

Titanite single spot common Pb-corrected dates range from 400 to 280 Ma with 12-20 Ma propagated 2SE. In sample SSP18-1B, titanite displays distinctive trends in minor and trace element composition and rare earth element (REE) pattern with date (Fig. 6). Zoning in BSE is largely concentric, with the oldest dates obtained from rare BSE dark cores and the youngest ages obtained from BSE dark rims (Fig. S6). Analyses show an increase in Y and Fe through time, a decrease in Sr, variable Th/Pb, increasing U/Th, and a decoupling of Ce from Sr (Fig. 6). Chondrite normalized REE patterns also reveal increasing HREE through time, followed by a late depletion in LREE (Fig. 7a). These trends correspond to distinct zones in BSE, revealing four texture-chemistry-date populations (Fig. 8). First, BSE dark cores are dated to 399 ± 14 and exhibit low Y
+ Fe + Zr concentrations, and a steep MREE to HREE slope (Fig. 6 & 7a). Two titanite grains (1 and 2, Fig. S6a–b) show complicated internal zoning with mottled textures and occasional minor oscillations. These grains display dates, Y + Fe concentrations, and low HREE consistent with BSE dark core domains; however, some analyses of these regions give dates less than <380 Ma which may indicate disturbance of the U-Pb system. If the oldest analyses of mottled zones are included with BSE dark zones, a pooled age of 397 ± 5 is calculated for Population 1 (Table 1). BSE dark cores are surrounded by BSE bright mantles; however, in most grains only BSE bright cores are observed. The BSE bright domains are divided into Populations 2 and 3. Population 2 domains give a pooled date of 372 ± 7 and have elevated Y and Zr concentrations, a shallower MREE to HREE slope, and clear Eu anomaly (Table 1, Fig. 7a). Population 3 domains were observed in two grains with angular BSE bright cores and gives a pooled date of 342 ± 8. Unlike Population 2, Population 3 domains exhibit lower Zr and Sr concentrations and display no Eu anomaly (Table 1, Fig. 7a). Finally, Population 4 BSE dark rim and mantle domains are dated to 295 ± 6 and display high Y, low Zr + Sr, and a distinctive depletion in the LREE (Table 1, Fig. 7a).

### Table 1. Summary of titanite date, trace element concentration, and Zr-in-titanite temperature for sample SSP18-1B.

<table>
<thead>
<tr>
<th>Population</th>
<th>Pooled Concordia Date&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Mean Trace Element Concentrations (μg g⁻¹)</th>
<th>Zr-in-Titanite Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>95 % CL</td>
<td>n</td>
</tr>
<tr>
<td>1</td>
<td>397</td>
<td>5</td>
<td>14</td>
</tr>
<tr>
<td>2</td>
<td>372</td>
<td>7</td>
<td>13</td>
</tr>
<tr>
<td>3</td>
<td>342</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td>4</td>
<td>295</td>
<td>6</td>
<td>26</td>
</tr>
</tbody>
</table>

<sup>a</sup>Model 1 Discordia (Vermeesch, 2018), <sup>b</sup>weighted standard deviation adjusted for overdispersion.

In contrast to sample SSP18-1B, most analyses of titanite grains from SSP18-1A do not display clear trends with date. Most analyses produce a cloud of data in ²³⁸U/²⁰⁶Pb – ²⁰⁷Pb/²⁰⁶Pb space with associated dates spanning a range of 380 and 330 Ma (Fig. 8b). No correlation is observed between age and Y and Fe concentrations and REE pattern for dates within this range (Figs. 6 & 7). A comparison of analysis location, zoning in BSE, and trace element concentrations reveals zones with similar composition and varying date, but also crosscutting zones with distinct trace element compositions and overlapping dates (Figs. S4–S5). The latter case is demonstrated by LA-ICP-MS mapping of a titanite grain from SSP18-1A: Internal zones show large variation in Y and Zr concentrations at a nearly constant ²⁰⁷Pb/²⁰⁶Pb of 0.07 – 0.10 (Fig. 9). Like SSP18-1B, titanite grains in SSP18-1A display young BSE dark rims with lower Zr and a pooled date of 306 ± 7 Ma (MSWD = 3.50, n = 19). These rim domains exhibit more restricted, and generally lower, Fe, and Y relative to older interior domains, but do not display the LREE depletion observed in SSP18-1A (Figs. 6, 7, & 9). Rim domains consistently exhibit higher ²⁰⁷Pb/²⁰⁶Pb ratios (>0.10).
relative to interior domains (< 0.10), suggesting a greater incorporation of unradiogenic initial Pb (Fig. 9). Occasional sector zoning is observed in titanite grains from SSP18-1A (Fig. 3a). Bright and dark sectors in BSE exhibit equivalent $^{207}\text{Pb}/^{206}\text{Pb}$ ratios and U, Sr, and Nb concentrations, suggesting these elements were not partitioned between sectors during crystallization. In contrast, Y and REE are enriched in BSE bright sectors by 150–200 % and <50 %, respectively.

8. DISCUSSION

8.1 Linking titanite dates to metamorphic processes

Titanite in sample SSP18-1B reveal a 105 Myr history of metamorphism from ca. 400 to ca. 295 Ma. Titanite in this sample largely exhibit core-to-rim zoning in BSE, with a decrease in date between cores and rims (Fig. S6), consistent with multiple stages of secondary overgrowth (e.g., Castelli & Rubatto, 2002). Beginning with the earliest stage, Population 1 BSE dark cores and mottled domains are dated to ca. 400 Ma and give a range of 690–785 °C for maximum Zr-in-titanite temperatures. There is little textural evidence of this early metamorphic event. Initial neocrystallization of titanite may have formed by the breakdown of rutile or biotite (Reactions 4–6). No inclusions of biotite or rutile are observed in titanite or other phases, suggesting that these phases were not present following prograde metamorphism at ca. 400 Ma. Amphibole and clinzoisite/epidote in calcsilicate rocks may also contain 10s to 1000s of µg g$^{-1}$ of Ti (Table S7). The prograde breakdown of these phases may have also promoted Population
1 titanite crystallization. The wide range in Th/Pb ratios (5–25) observed in these domains (Fig. 6d) does give some indication of possible reactions at ca. 400 Ma. Allanite will strongly fractionate Th from Pb and the prograde breakdown of allanite may have increased the Th/Pb ratio of coexisting titanite (Fig. 10b). Considerable variation in LREE concentrations is also observed in Population 1 domains, consistent with allanite-breakdown (Fig. 7a) (Johnson, Cottle, & Larson, 2021). Our explanation is consistent with rock-buffered fluid compositions anticipated for prograde of calc-silicates: In a closed-system, prograde decarbonation reactions buffer the fluid from low X\text{CO}_2 conditions to higher X\text{CO}_2 conditions, resulting in the breakdown of early prograde epidote group minerals to form An-rich plagioclase (Fig. 10a) (e.g., Ferry, 1976; Groppo et al., 2017; Rapa et al., 2017).

The transition between 400 Ma (Pop. 1) and 372 Ma (Pop. 2) domains is marked by a sharp increase in Y+HREE (Fig. 6c & 7a) and a maximum Zr-in-titanite temperature of ~800 °C (Table 1). Garnet and amphibole may partition Y+HREE from titanite (Garber et al., 2017). Garnet is not observed as a prograde phase and only rarely occurs in the psammitic to pelitic bands in our field area; therefore, garnet breakdown at 372 Ma is unlikely to have enriched titanite in these elements. Instead, amphibole is observed together with titanite as inclusions in diopside but does not occur in the peak assemblage. We suggest that amphibole, rather than garnet, breakdown at ca. 372 Ma led to the increase in Y+HREE (Fig. 10b). Diopside grains also reveal a protracted metamorphic
Figure 10. (a) Petrogenetic T-XCO₂ grid calculated at 500 MPa for the CaO-Al₂O₃-SiO₂-H₂O-CO₂ (green), CaO-MgO-Al₂O₃-SiO₂-H₂O-CO₂ (black), and CaO-Al₂O₃-SiO₂-TiO₂-H₂O-CO₂ (orange) systems calculated in Perple_X (Connolly, 2005). Reactions involving MgO are also calculated with an activity of diopside (aDi) of 0.70. The grey field denotes the high-T moderate to high XCO₂ assemblage Di + An + Cal + Ttn, whereas the blue field denotes the stability field of Zo (or Czo). (b) Shows a schematic diagram illustrating the reaction sequences inferred from mineral textures, compositions, and titanite trace element compositions between ca. 400 and ca. 295 Ma (see text for details).

describes the history of an early low XDi phase followed by fracturing and recrystallization of diopside with higher XDi (Fig. 4b). In addition to a decrease in Al and Fe between fractured and healed diopside, our analyses also reveal lower in Ti, Y, and Sr in healed domains (Fig. 5). Given that titanite is the only high-T phase to partition Y+HREE over pyroxene in our rocks, mass balance requires recrystallization of fractured diopside formed at ca. 400 contributed to Y+HREE enrichment in titanite at ca. 372 Ma. Population 2 titanite likely formed through a combination of neocrystallization, as well as secondary overgrowth on Population 1 domains, driven by residual amphibole breakdown and diopside recrystallization (Fig. 10b). Lower activities of diopside shift the diopside-in reactions to lower T (Fig. 10a). The increase in XDi and high maximum Zr-in-titanite temperatures require that peak metamorphic conditions occurred at ca. 372 Ma.

Population 3 domains at ca. 342 Ma exhibit no Eu anomaly and lower maximum Zr-in-titanite temperature (~765 °C) relative to Population 2 domains (Fig. 7a & Table 1). Absolute Eu concentrations increase by ~30 µg g⁻¹ between ca. 372 and ca. 342 Ma, whereas Sm and Gd concentrations are constant. Plagioclase breakdown will release Eu relative to other REE. Plagioclase in SSP18-1B contains 1098–1282 µg g⁻¹ of Sr, and plagioclase breakdown would release significant Sr in addition to Eu. Titanite Sr concentrations decrease through time across all domains (Fig. 6b). Prograde metamorphism of calc-silicates at temperatures below scapolite stability produce increasing plagioclase modes at the expense of other Ca-bearing phases, such as calcite and clinzoisite (e.g., Ferry, 1976; Groppo et al., 2017; Walters & Kohn, 2017). We suggest that the decrease in titanite Sr concentrations with time represents increasing plagioclase modes. Therefore, plagioclase breakdown at ca. 342 Ma cannot account for the lack of the Eu anomaly in
Population 3 titanite domains. Interestingly, Population 3 titanite were only observed in a diopside-rich (~80 vol. %) band (Fig. 10b). Plagioclase incorporates Sr, but not LREE, whereas other phases, such as epidote group minerals, incorporate both Sr and LREE. Exchange between titanite and plagioclase will produce variable Sr but LREE will remain unaffected. Population 2 and 3 titanite domains from the diopside-rich band exhibit variable Fe, low Sr, and high Ce, whereas Population 3 domains from the plagioclase-rich band exhibit a variable Sr concentrations and restricted Ce concentrations (Fig. 6a & f). We suggest that poor intragrain mobility of trace elements between the diopside- and plagioclase-rich bands resulted in local equilibrium. Additionally, local equilibria must have controlled titanite (re-)crystallization at ca. 342 Ma, as domains of this date are not observed in the plagioclase-rich bands (Fig. 10b).

Finally, BSE dark Population 4 rims dated to ca. 295 Ma exhibit low Sr, Th/Pb, and LREE, high U/Th, and deep negative Eu anomaly (Figs. 6 & 7a). We link these trace element characteristics to infiltration of a hydrous during this final stage of titanite (re-)crystallization (Fig. 10b). A decrease in $X_{CO_2}$ to <0.10 ($X_{H_2O}$ > 0.90) would stabilize clinozoisite over plagioclase and result in the breakdown of diopside to amphibole, both of which texturally overprint the earlier higher $X_{CO_2}$ Di + Kfs + Pl + Cal assemblage in our samples. Clinozoisite-epidote Sr concentrations (771–1595 µg g$^{-1}$) overlap with those of plagioclase, consistent with plagioclase-breakdown and the low Sr concentrations in Population 4 titanite. Additionally, clinozoisite-epidote grains exhibit positive Eu anomalies. Allanite, which is observed in some clinozoisite-epidote cores, contains elevated Th/Pb and LREE, and low U/Th relative to titanite. Therefore, the low Sr, Th/Pb, and LREE, high U/Th, and deep negative Eu anomaly characteristic of Population 4 titanite may be entirely explained by exchange between titanite and epidote group minerals. Population 4 domains from both the plagioclase- and diopside-rich bands exhibit similar textures and compositions, suggesting a higher degree of trace element mobility at ca. 295 Ma. Garber et al. (2017) and Johnson, Cottle, & Larson (2021) also observed titanite with low LREE and Sr in titanite from allanite-bearing rocks. Rims dated in sample SSP18-1A exhibit similar pooled date of 306 ± 7 Ma ($n = 19$, MSWD = 3.50). In both samples, titanite rims are elevated in OH, F, and Al (Fig. 3), consistent with a high $X_{H_2O}$ fluid (Garber et al., 2017). Ferry (1976, 2016) observed a late influx of H$_2$O-rich fluids in rocks ~60 km northeast of our field area, suggesting potentially widespread metasomatism and retrograde metamorphism at ca. 295 Ma.

8.2 Decoupling of titanite U-Pb dates and trace element compositions

Lead-loss, U-loss, and/or inheritance of unsupported radiogenic Pb can result in apparent ages that do not reflect the timing of metamorphism (e.g., Bonamici & Blum, 2020; Castelli & Rubatto, 2002; Garber et al., 2017; Kirkland et al., 2016; 2018; Smye et al., 2018). As a result, U-Pb dates may decouple from major and trace element compositional zoning. Titanite from sample SSP18-1A exhibit no clear trends in texture, major and trace element concentrations, and date between ca. 380 and ca. 320 Ma (Fig. 6, 7b, & 9). Additionally, titanite grains display equivalent dates between texturally overprinting zones, as well as significant age variation within some chemically homogenous zones (Figs. S4, S5). Plotted in a Tera-Wasserburg diagram, the analyses
of grain interiors in sample SSP18-1A form a sub-horizontal array with a narrow range (0.08 to 0.10) in $^{207}\text{Pb}/^{206}\text{Pb}$ (Fig. 8b). In contrast, rim analyses of titanite in SSP18-1A, dated to 306 ± 7 Ma, exhibit more restricted trace element concentrations and are similar in composition and age to Population 4 rim analyses in Sample SSP18-1B. Based on these observations, we suggest that titanite dates for interior zones from sample SSP18-1A do not reflect the timing of (re-)crystallization.

The diffusion rate of Pb will differ from those of other trace elements with dissimilar size and charge; as a result, diffusive Pb-loss may decouple dates from trace element zoning patterns. Although experimental diffusion rates for Pb in titanite suggest a $T_c$ of 600 °C (Cherniak, 1993), natural data suggest Pb-loss by diffusion is unlikely to be efficient below 850 °C (Hartnady et al., 2019; Holder et al., 2019; Kohn, 2017; Stearns et al., 2016; Walters & Kohn, 2017). Despite undergoing at least four metamorphic episodes, we observe no evidence of Pb diffusion in our samples. In SSP18-1B, core-to-rim variations in U-Pb date correspond closely to variations in texture and trace element chemistry, inconsistent with diffusive Pb-loss. Our dates for SSP18-1B also correlate well with known metamorphic and igneous events in the region, such as regional Acadian metamorphism at ca. 400 Ma and Neocadian metamorphism to the northwest at ca. 376 Ma (Sanislav, 2011; Smith & Barreiro, 1990; Solar et al., 1998; Tomascak & Solar, 2016). Lead-loss cannot explain the different U-Pb date-textural relationship between our two samples, which were collected from the same outcrop and experienced the same metamorphic histories. Additionally, the LA-ICP-MS Pb map of titanite from SSP18-1A reveals relatively sharp zoning (Fig., 9d). Our data are therefore consistent with the reevaluated $T_c$ for Pb in titanite.

Incorrect assumptions about the initial $^{207}\text{Pb}/^{206}\text{Pb}$ may also result in a decoupling of titanite U-Pb dates from major and trace element patterns. Prograde, diagenetic, and detrital U-bearing phases, such as monazite, xenotime, allanite, and rutile, may break down during metamorphism (see discussions in Engi, 2017; Kohn, 2017) to release unsupported radiogenic Pb (e.g., Kirkland et al., 2018; Marsh & Smye, 2017). As a result, the initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio of a metamorphic mineral will represent a mixture of ‘common’ unradiogenic Pb and unsupported radiogenic Pb (Bonamici & Blum, 2020; Romer, 2001). Multiple analyses of one or more phases are commonly used to develop an isochron but assumes that each aliquot received the same initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio. Metamorphic titanite $^{238}\text{U}/^{206}\text{Pb}$ and $^{207}\text{Pb}/^{206}\text{Pb}$ data often form linear arrays on Tera-Wasserburg diagrams, through which a linear regression constrains the initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio (e.g., Bonamici & Blum, 2020; Marsh & Smye, 2017; Kirkland et al., 2017, 2018). However, multiple linear arrays with different initial $^{207}\text{Pb}/^{206}\text{Pb}$ intercepts are sometimes observed, suggesting a variable contribution of unsupported radiogenic Pb (e.g., Bonamici & Blum, 2020; Romer & Rötzler, 2003). Correction of initial Pb by isochron is therefore only possible if linear arrays can be confidently discerned. Additionally, if radiogenic Pb released during the breakdown of U-bearing phases does not reach isotopic equilibrium with Pb in other phases then the $^{207}\text{Pb}/^{206}\text{Pb}$ ratio of low U/Pb phases cannot be used to determine the initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio of titanite. Often the $^{207}\text{Pb}/^{206}\text{Pb}$ ratios measured on U-poor phases, such as plagioclase and alkali feldspar, fall within the value of initial $^{207}\text{Pb}/^{206}\text{Pb}$ defined by regression through titanite U-Pb data.
For Sample SSP18-1B, we observe that isochrons for pooled dates at ca. 400 and ca. 295 Ma (Pop. 1 & 2, respectively) demonstrate good agreement with the \(^{207}\text{Pb}/^{206}\text{Pb}\) ratio measured on plagioclase (Fig. 8a). In contrast, U-Pb data from sample SSP18-1A form a cloud on the Tera-Wasserburg diagram with a narrow \(^{207}\text{Pb}/^{206}\text{Pb}\) ratio (Fig. 10b). No isochrons are discernable for analyses of grain interiors, consistent with a variably mixed initial \(^{207}\text{Pb}/^{206}\text{Pb}\) ratio (Bonamici & Blum, 2020).

We suggest that partial incorporation of unsupported radiogenic Pb during interface-coupled dissolution-reprecipitation (ICDR) reactions is responsible for the decoupling of titanite textures, compositions, and dates in Sample SSP18-1A. During ICDR, dissolution of the reactant phase produces a thin interfacial boundary layer of fluid that is supersaturated with respect to product phases, which nucleate at the reaction front (Putnis, 2009). Porosity is generated if the volume of reprecipitated solid matter is less than the volume dissolved, allowing the reaction front to propagate through the crystal if porosity is maintained (Ruiz-Agudo, Putnis, & Putnis, 2014). Porosity is reduced at high temperatures by textural equilibration and, if the product phase is structurally similar to the reactant, crystallographic continuity may be preserved (Ruiz-Agudo, Putnis, & Putnis, 2014).

Titanite grains from SSP18-1A exhibit cross-cutting, lobate, uneven, and fragmentary zones which are consistent with extensive ICDR (Figs. 3 & 9; see review in Putnis, 2009). The lack of correlation between variations in titanite crystallographic orientation and chemical zoning is also consistent with ICDR (Figs. S7–S8). Lead liberated during titanite dissolution may be partially scavenged from the fluid before isotopic equilibrium is achieved with the bulk rock. For example, titanite formed at 400 Ma with 5 to 20 % common Pb may have recrystallized during subsequent metamorphic recrystallization at 340 or 300 Ma (Fig. 11). The entire range of titanite U-Pb ratios in SSP18-1A may be explained by 50 % or less retention of Pb during ICDR. Similar to our study, Castelli & Rubatto (2002) found that the mixed-dates of complexly zoned
metamorphic titanite from the Western Alps represented incomplete isotopic resetting during recrystallization. Inheritance of unsupported radiogenic Pb has also been observed in monazite. For example, Seydoux-Guillaume et al. (2003) observed a spread in monazite dates between initial growth at 2.5 Ga and recrystallization at 790 Ma, which they attributed unsupported Pb in recrystallized domains. Weinberg et al. (2020) found that monazite dates, spread over 50 myr, were decoupled from chemical zonation and reflected inherited radiogenic Pb from older domains during ICDR reactions. When taken together with our study, these data suggest that dates of phases sensitive to ICDR, such as monazite and titanite, may lack geological meaning in polymetamorphic rocks and should be carefully screened prior to interpretation.

In contrast to SSP18-1A, titanite from SSP18-1B are dominated by regular core-to-rim zoning in composition and date that is consistent with titanite crystallization (Castelli & Rubatto, 2002). We suggest that neocrystallization and secondary growth of titanite, which requires reactions with low U/Pb major mineral phases to liberate Ca, Ti, Al, and Si, will form titanite with an initial Pb composition dominated by common Pb. In contrast, recrystallization of titanite by ICDR may limit isotopic homogenization of Pb. Additionally, the larger grain size and higher modal abundance of titanite in SSP18-1A (<2.5 mm, ~ 10 vol %) relative to SSP18-1B (<800 µm, 1–2 vol %), suggests that a greater percentage of the bulk rock Pb budget is stored in titanite in SSP18-1A. These differences may have also played a role in the availability of radiogenic Pb relative to common Pb during titanite ICDR reactions.

8.3 Common Pb uptake during crystallization

Titanite may incorporate significant common Pb upon crystallization (e.g., Bonamici & Blum, 2020; Kirkland et al., 2018; Kohn, 2017). High ratios of common to radiogenic Pb require large extrapolations to concordia and result in U-Pb dates with high uncertainties. The initial U/Pb ratio inherited by titanite will be a function of the bulk rock U/Pb ratio and the relative partitioning of these phases between minerals during continuous and discontinuous metamorphic reactions (Römer, 2001).

Titanite in both our samples display less than 20 % inherited common $^{207}$Pb/$^{206}$Pb ($f_{207}$) (e.g., Kirkland et al., 2016; 2018). In Figure 12, we plot $f_{207}$ of titanite analyses from SSP18-1B as a function of date and Sr concentration. A decrease in $f_{207}$ is observed between ca. 400 and ca. 372 Ma followed by an increase from ca. 342 to ca. 295 Ma (Fig. 12a). Strontium concentrations in titanite are positively correlated with $f_{207}$ at ca. 400 Ma, uncorrelated at ca. 372 Ma, and negatively correlated at ca. 295 Ma (Fig. 12b). Plagioclase in SSP18-1B contains 20.5–23.9 µg g$^{-1}$ Pb, compared to 1.55–28.8 µg g$^{-1}$ measured in titanite (radiogenic + inherited Pb, Table S2 & S7). Plagioclase is considerably more abundant (up to ~80 vol % in plagioclase-rich bands) than titanite (1–2 vol %) and is therefore the largest Pb sink in SSP18-1B. The initial decrease in $f_{207}$ and Sr corresponds to an increase in plagioclase modes between ca. 400 and ca. 372 Ma, whereas the elevated $f_{207}$ at ca. 295 Ma corresponds to plagioclase breakdown (Fig. 10b & 12). We suggest that the presence of phases that fractionate Pb from U, such as An-rich plagioclase and alkali
Figure 12. Percent of common Pb ($\alpha^{207}$) in titanite plotted relative to (a) date and (b) Sr concentration. The average 2SE uncertainty of single spot dates (black scale bar) and trends with time (red arrow) are shown. In (a), plagioclase shows a trend of decreasing $\alpha^{207}$ attributed to plagioclase growth, followed by an increase in $\alpha^{207}$ corresponding to hydration, An-rich plagioclase breakdown, and the formation of epidote group minerals. In (b), Sr concentrations roughly mirror the trend with date, showing correlation between $\alpha^{207}$ and Sr during the early and final stages of metamorphism.

feldspar, is a key factor contributing to low common Pb and may lead to both more precise and accurate titanite U-Pb ages.

8.4 $P$–$T$ conditions of retrogression and Zr-in-titanite disequilibrium

Population 4 titanite rim domains dated to ca. 295 Ma in sample SSP18-1B give Zr-in-titanite temperatures of $\sim$750 °C. Titanite rims are associated with the retrograde assemblage amphibole, clinozoisite/epidote, allanite, albitic plagioclase, and quartz. The presence of zircon and retrograde quartz suggests that assumptions of unity for $\alpha_{zrSiO_4}$ and $\alpha_{SiO_2}$ are reasonable for Population 4 Zr-in-titanite temperatures. The $T$ estimate of 750 °C is inconsistent with the retrograde assemblage in the calc-silicates. The reaction of diopside and anorthite to quartz, clinozoisite, and tremolite (Reaction 9) observed in SSP18-1B is expected to occur at $\leq$550 °C at 500 MPa (Fig. 10a). Additionally, attempts to calculate pressures using the titanite, clinozoisite, anorthite, rutile, quartz (TZARS) barometer of Kapp, Manning, & Tropper (2009) at 750 °C produced unrealistically high pressures of $>$1.0 GPa, whereas more reasonable pressures of 600 MPa were calculated for 500 °C.

The Zr-in-titanite temperatures are also inconsistent with retrograde assemblages of metapelite and metapsammitic rocks from the sampled outcrop. At 500 MPa, vapor-present muscovite-dehydration melting occurs at temperatures above $\sim$650 °C (Fig. S11). Sillimanite is observed in garnet inclusions in metapelite sample SSP18-1D and no sillimanite is observed in the matrix, in contrast muscovite is abundant in the matrix but absent from inclusions. Rehydration of metapelites below the wet-solidus will produce muscovite at the expense of sillimanite (the reverse of the muscovite-dehydration reaction). Prehnite is observed to replace biotite in metapsammitic
bands in Sample SSP18-1A. These require relatively low $T$ retrogression and hydration of the metapelite and metapsammite rocks and suggest a shared history of fluid infiltration with the calcisilicate bands at ca. 295 Ma. Johnson, Cottle, & Larson (2020) suggested limited equilibrium between titanite and zircon in their samples resulted in Zr-in-titanite temperatures that are 100 to 150 °C higher than temperatures calculated from other techniques and inconsistent with the observed mineral assemblages. Here we also suggest that quartz and/or zircon cannot be in equilibrium with titanite at ca. 295 Ma and suggest caution when applying the Zr-in-titanite thermometer (e.g., Cruz-Uribe et al., 2018).

8.5 Implications for the tectono-metamorphic history of western Maine

The Acadian Orogeny (420 to 350 Ma) is the dominant tectono-metamorphic event in the northern Appalachian Mountains and corresponds to the collision of the Avalon terrane with the eastern margin of Laurentia (Robinson et al., 1998; van Staal et al., 2009). Based on stratigraphic constraints and intrusion ages, Bradley et al. (2000) suggested that the Acadian deformation front moved from coastal Maine into Québec between at 423 and 382.5 Ma. The northwestward migration of the deformation front is consistent with ca. 400 Ma zircon and monazite ages observed throughout west-central Maine and New Hampshire (Gerbi & West, 2007; Pyle et al., 2005; Moecher, McCulla, & Massey, 2020; Smith & Barreiro, 1990; Solar et al., 1998; Wing, Ferry, & Harrison, 2003). The pooled Population 1 titanite date of ca. 400 Ma therefore corresponds to widespread regional Acadian metamorphism.

Studies of metamorphism in western Maine have long proposed that metamorphism in western Maine during the Neoacadian (380–350 Ma) and Alleghanian (325–260) Ma orogenies (Robinson et al., 1998) was driven by pluton emplacement and contact heating, resulting in localized overprinting cycles of low-$P$ prograde and retrograde metamorphism (see review in Guidotti, 1989). The increase in $X_{\text{Di}}$ between recrystallized and fractured diopside, breakdown of residual amphibole ($\pm$ biotite), and higher maximum Zr-in-titanite $T$ indicate that the highest thermal conditions experienced by our samples occurred at ca. 372 Ma (Fig. 10). These data, in conjunction with the pooled ca. 376 Ma age of zircons from migmatites to the west of our field area (Tomascak & Solar, 2016), suggest that anatexis and deformation in this region occurred 20–30 Ma later than in the migmatites in the vicinity of the Mooselookmeguntic pluton (Fig. 1a) (Brown & Solar, 1999; Johnson, Brown, & Solar, 2003). Partial melting ceased prior to ca. 295 Ma, suggesting no connection between anatexis and the emplacement of the Sebago pluton. Instead, high-$T$ metamorphism at ca. 372 Ma overlaps with the 380–330 Ma period of partial melting observed in New Hampshire, central Massachusetts, and central Connecticut (see compilation in Hillenbrand et al., 2021). Retrograde metamorphism at ca. 295 Ma corresponds to the age of the Sebago pluton (Tomascak, Krogstad, & Walker, 1996), which may have been the source of the high $X_{\text{H}_2\text{O}}$ fluids.

Neoacadian and Alleghanian metamorphism in western Maine is thought largely to be post-deformational, following localization of deformation to the Norumbega Fault System by ca. 380 Ma (Hubbard et al., 1995; Gerbi & West, 2007; Ludman, 1998; West & Hubbard, 1997; West,
Metapelitic rocks in our field area exhibit stretched leucosomes that are oriented parallel to boudinaged dikes (Fig. S1a-b), which require that anatexis and dike emplacement was either pre- or syn-tectonic. A second generation of dikes is also observed, but these are undeformed and crosscut the foliation of the host gneisses (Fig. S1c-d). Based on these relationships, we also suggest that deformation in west-central Maine continued even after the initiation of dextral shear on the Norumbega Fault System, requiring that at least some strain was partitioned further inland.

9. CONCLUSIONS

Titanite is a highly reactive mineral that may record multiple generations of crystallization, growth, and dissolution-reprecipitation. In addition, the temperature-dependent incorporation of Zr and the slow diffusivity of Pb make titanite a potentially excellent petrochronometer. However, titanite may incorporate significant initial Pb, potentially resulting in inaccurate age determination. Here we demonstrate both the benefits and shortcomings of titanite chronometry. Our data for sample SSP18-1B reveals four phases of metamorphism spanning a period from ca. 400 to ca. 295 Ma. Most titanite grains in this sample exhibit systematic core-to-rim variations in trace element concentrations and U-Pb dates, consistent with neocrystallization and secondary overgrowth. Trace element concentrations and REE patterns reflect changing partitioning between titanite and other phases, revealing the breakdown of hydrous phases between ca. 400 and ca. 342 followed by an influx of hydrous fluids at ca. 295 Ma. Metamorphism at ca. 400 Ma corresponds to the northwestward migration of the Acadian front, whereas the highest temperature conditions correspond a period of metamorphism at ca. 372 Ma and are consistent with wide-spread partial melting observed throughout greater New England. Retrogression at ca. 295 Ma may be related to hydrous fluids exsolved from the crystallizing Sebago pluton to the SW of our field area. In contrast, titanite from a second sample (SSP18-1A) exhibits little correlation between trace element concentrations, REE patterns, texture, and date. We suggest date-composition decoupling is caused by the incorporation of variable amounts of unsupported radiogenic Pb in recrystallized domains, resulting in mixed geologically-meaningless ages. We suggest that titanite may reliably record multiple phases of metamorphism; however, U-Pb dates should be coupled with chemical and textural characterization to screen for unsupported radiogenic Pb in recrystallized domains.

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REFERENCES


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