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Titanium isotopes constrain a magmatic transition at the Hadean-Archean boundary in the Acasta Gneiss Complex

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Plate subduction greatly influences the physical and chemical characteristics of Earth’s surface and deep interior, yet the timing of its initiation is debated because of the paucity of exposed rocks from Earth’s early history. We show that the titanium isotopic composition of orthogneisses from the Acasta Gneiss Complex spanning the Hadean to Eoarchean transition falls on two distinct magmatic differentiation trends. Hadean tonalitic gneisses show titanium isotopic compositions comparable to modern evolved tholeiitic magmas, formed by differentiation of dry parental magmas in plume settings. Younger Eoarchean granitoid gneisses have titanium isotopic compositions comparable to modern calc-alkaline magmas produced in convergent arcs. Our data therefore document a shift from tholeiitic- to calc-alkaline–style magmatism between 4.02 and 3.75 billion years (Ga) in the Slave craton.

INTRODUCTION

Modern manifestations of plate tectonics, such as seafloor magnetic reversal records (1), are absent from much of Earth’s geological record, and researchers must instead rely on indirect evidence to recognize the action of plate tectonics in the past. Arc magmatism that occurs at subduction zones is one of the hallmarks of modern plate tectonics (2–4). Therefore, geochemical comparisons between ancient rocks and modern arc-derived igneous rocks has been the focus of many studies aimed at understanding when Earth entered into a geodynamic regime involving subduction. Ancient subduction may have differed in many respects from present-day occurrences, so interpretation of the rock record can be difficult. For example, age estimates for the initiation of plate tectonics range from ~0.85 to >4.2 billion years (Ga) (5–13), spanning more than two-thirds of Earth’s history.

Modern calc-alkaline magmas are uniquely associated with subduction zone environments. Characteristic features of calc-alkaline magmas are their higher water contents, and both relative Fe depletion, and greater oxidation than magmas in other tectonic settings (2, 14–18). Tholeiitic magmas, on the other hand, are characterized by sharp FeO and TiO₂ enrichments during early magmatic differentiation, followed by more protracted depletions in these elements (2, 15). The reason for the unique geochemical evolution of calc-alkaline magmatic series relative to the tholeiitic series is primarily the water-rich and oxidized nature of their parental melts, which suppresses crystallization of plagioclase and tends to promote the early crystallization of oxides and their subsequent loss as cumulates (19). On the modern Earth, the above conditions are encountered almost exclusively near subduction zones, where oxidized and water-rich fluids from subducted oceanic crust flux-melt the overlying mantle and trigger arc magmatism.

Ancient rocks are exceedingly rare, with >3.6-Ga outcrops representing only a few parts per million of Earth’s exposed crust (20), and those that are preserved have often undergone multiple metamorphic events, each with the potential to chemically overprint primary magmatic signatures. This is true for Earth’s oldest presently known rocks, found in the Acasta Gneiss Complex (AGC, Slave craton, Canada). These rocks have experienced multiphase metamorphism, deformation, and igneous intrusion, which has resulted in the partial obliteration of primary textures and possibly bulk compositions, so that inferring primary chemical compositions is somewhat complicated. Reimink et al. (21, 22) identified several rock units, most notably the 4.02-Ga Jdi whàa (translates to “ancient times” in the language of the Tjîchop people who are indigenous to this area) tonalite gneiss (ITG) and volumetrically dominant 3.6- to 2.9-Ga granitoid rocks within the AGC, whose bulk compositions point to a transition between ~4.0 and 3.6 Ga from Fe-rich to Fe-poor rock series. The rocks from the ITG are a part of the AGC, and for the remainder of the text, we use younger AGC to denote granitoids younger than ~3.9 Ga. A plausible interpretation for the difference between ITG and younger AGC rocks is that ITG rocks differentiated from a dry magma, while AGC granitoids differentiated from wet magmas produced by melting of hydrated basalt (21–23). Trace element and radiogenic isotope compositions of the AGC have indicated changes in the modes of crustal generation at 3.55 Ga (22, 24–27), including evidence of a transition from shallow to deep-seated melting as well as a shift in the melt source from Hadean protocrust to hydrated primitive mantle. The rocks from the AGC have experienced a complex history, which makes a straightforward interpretation of elemental compositions of ancient rocks difficult. The diagrams used for classifying rocks into calc-alkaline and tholeiitic series and for identifying Fe enrichments typically involve plotting the samples in Mg/FeO total versus SiO₂ (28) or Na₂O + K₂O-FeO-MgO (14) diagrams. Many of the elements used for classification are fluid mobile (Mg, K, and Na), and a potential concern is that the protoliths of the Acasta granitoids were affected by fluid–rock interactions that modified the bulk chemical composition of the rocks. Given the amphibolite metamorphic grade of these rocks, disturbance of primary chemical signatures is likely, but difficult to discern. Characterizing the conditions of differentiation in the AGC therefore requires geochemical

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proxies with memory of the path taken to a given rock chemical composition. Stable isotopic proxies are ideally suited to tackle this problem.

Recent work has shown that Ti isotopes can distinguish between rocks that underwent magmatic differentiation along the calc-alkaline trend and those that fall on the intraplate tholeiitic trend (29–32). The $^{49}$Ti/$^{47}$Ti isotopic ratio (reported in $\delta^{49}$Ti notation; the per mil deviation relative to the $^{49}$Ti/$^{47}$Ti ratio of the OL-Ti reference material) increases with SiO$_2$ content in both calc-alkaline and tholeiitic rocks (29–32), but with the latter showing a much more rapid increase in $\delta^{49}$Ti with SiO$_2$. Titanium is highly fluid immobile and unlikely to be substantially mobilized by metamorphic disturbance. The isotopic composition of Ti can therefore provide clues on the crystallization of Fe-Ti oxides on the liquid line of descent, which is a function of the water content, oxygen fugacity, and the initial geochemical composition. Thus, whole-rock Ti isotopes provide a valuable tool for identifying processes governing igneous rock formation and even geodynamic setting back in time, even on samples that have been multiply metamorphosed and possibly metasomatized.

Iron isotopic ratios are also fractionated by magmatic differentiation, and $^{56}$Fe/$^{54}$Fe ratios (reported as $\delta^{56}$Fe, the per mil deviation relative to the $^{56}$Fe/$^{54}$Fe ratio of the IRMM-014 reference material) can also potentially trace Fe-Ti oxide crystallization (fig. S1). Here, we present Ti and Fe isotopic compositions of a suite of crustal magmatic rock samples ranging in age from 4.02 to <3.6 Ga with the aim of better characterizing the transition of magmatism in the AGC. The iron isotope data show notable scatter and are not straightforward to interpret. These data are not discussed further below, but more details are provided in the Supplementary Materials.

RESULTS
We observe variations in the Ti isotopic composition of ITG and younger AGC sample rocks (Fig. 1B). We also present the development of a new Ti isotope proxy that allows us to parameterize the affinity of ancient, altered rocks for intraplate tholeiitic versus calc-alkaline trends. In contrast to the indistinguishable evolution of TiO$_2$ content with increasing SiO$_2$ in the sample set analyzed (Fig. 1A), we find that the $\delta^{49}$Ti compositions of AGC samples fall on two distinct trends (Fig. 1B) that both show increasing $\delta^{49}$Ti values, ranging from ~+0.08 to 0.96‰, with increasing SiO$_2$. Titanium isotope data for the 4.02-Ga ITG and younger ≤3.75-Ga AGC granitoids appear to plot along different trajectories, the former comparable to modern intraplate tholeiites and the latter on the trend characteristic of calc-alkaline differentiation (Fig. 1B). The $\delta^{56}$Fe values of the younger AGC span a range (from +0.03 to +0.17‰), and $\delta^{56}$Fe values of the 4.02 Ga tonalites show a slightly decreasing trend with SiO$_2$ content (fig. S1A).

DISCUSSION
Given the complex history of the rocks from the AGC, we consider the effects of fluid-rock interaction and metamorphic disturbance on the Ti isotopic compositions observed here. Like other high-field strength elements (HFSEs), Ti is very insoluble in most fluids of geological relevance (33). The TiO$_2$ content in schists does not change with increasing metamorphic grade (34), indicating little mobilization and redistribution of Ti by alteration processes such as metamorphism and water-rock interaction. Titanium and HFSEs can only be mobilized by localized alkaline, fluoride-bearing fluids (35) restricted to epithermal metal veins.

Ti isotopes and the Ti isotope magmatic index
Several geochemical metrics have been used to distinguish between calc-alkaline and tholeiitic magmas (2, 3, 28), but they often rely on examining the behavior of Fe and Mg relative to Si or alkali elements. As discussed above, these classification schemes use fluid mobile elements like K, Na, and Mg, which can be affected by fluid-rock interactions such as weathering or hydrothermal and metasomatic events. Furthermore, these schemes are most reliably used when the rocks sample a range of SiO$_2$, a scarcity in the ancient rock record. Last, many of these diagrams tend to converge in the most evolved rocks, and the classification schemes are most useful when intermediate SiO$_2$ rocks are available.

Titanium isotopes combined with major element compositions of rocks can be applied to overcome some of these problems and gauge the tholeiitic affinity of evolved rocks. We introduce a new empirical Ti isotope magmatic index (TIMI), which uses the relationship between the base 10 logarithm of the ratio between the SiO$_2$ and TiO$_2$ weight percentage log(SiO$_2$/TiO$_2$) and $\delta^{49}$Ti of rocks to define two distinct trends between tholeiitic rocks from Iceland, Afar hot spot, and Hawaii (30, 31) and calc-alkaline rocks from Agung and Kos (29, 36) (Fig. 2A) that contain >52 weight % (wt %) SiO$_2$ and are saturated in Fe-Ti oxides. The benefit of using log(SiO$_2$/TiO$_2$) versus $\delta^{49}$Ti is that the magmatic series define two distinct linear trends that can be fitted with a simple linear relationships (log is the base 10 logarithm, and SiO$_2$ and TiO$_2$ are concentrations in wt %)

$$\text{Tholeiitic: } \delta^{49}\text{Ti} = 1.14 \log_{10}\left(\frac{\text{SiO}_2}{\text{TiO}_2}\right) - 1.32$$

$$\text{Calc-alkaline: } \delta^{49}\text{Ti} = 0.64\log_{10}\left(\frac{\text{SiO}_2}{\text{TiO}_2}\right) - 1.09$$
for intraplate tholeiitic and subduction-related calc-alkaline rocks, respectively (Fig. 2A). The linear correlations converge at log(SiO$_2$/TiO$_2$) = 0.46 and $\delta^{49}$Ti = −0.80. The slope (TIMI) of the segment between a rock and the convergence point, defined above, provides a measure of how tholeiitic a melt is. The slope defines an equation in the form

$$\text{TIMI} = \frac{\delta^{49}\text{Ti} + 0.80}{\log(\text{SiO}_2/\text{TiO}_2) - 0.46}$$

The steeper the slope (higher TIMI), the more Fe-Ti oxides crystallized at low SiO$_2$ concentration, which is one of the characteristics of tholeiitic magmatic differentiation. We use the classic subduction-related calc-alkaline [Agung, Sunda Arc and Kos, Aegean Arc (29, 36)] and intraplate tholeiitic [Iceland, Afar, and Hawaii (30, 31)] rocks to define TIMI, with an average TIMI value of 0.64 for calc-alkaline and 1.15 for intraplate tholeiitic rocks. Alkaline rocks from an intraplate setting [Ascension, Heard, and Afar (different rock series than the calc-alkaline rock average (TIMI = 1.15). Rocks dated between 4.0 and 3.6 Ga have intermediate TIMIs (0.99), and AGC rocks younger than 3.94 Ga average at 0.71. Unknown age rocks (metagabbros and an amphibolite) have lower TIMI values suggesting calc-alkaline affinity. Blue and red dashed lines indicate first-order TIMI thresholds for calc-alkaline (TIMI ≤ 0.64), sub-calc-alkaline (TIMI = 0.64 to 1.15), and tholeiitic (TIMI ≥ 1.15) rocks. Calc-alkaline and tholeiitic MELTS model output and calculations for TIMI (see Materials and Methods for details on MELTS modeling and Ti isotopic composition of MELTS output) are shown as schematic representations of the trajectories (colored arrows).

In summary, TIMIs from subduction-related rocks are below 0.85, with lower values signifying stronger calc-alkaline affinity. Tholeiitic and alkaline rocks from intraplate settings have TIMIs higher than 0.90, and 85% of these are above 1.00. Therefore, this index allows us to identify the calc-alkaline affinity of an individual evolved rock and the identification of rocks from intraplate or subduction origin. As an additional benefit, it only involves a major element (SiO$_2$) and an insoluble HFSE (Ti) and is therefore a robust method that should “see through” fluid-rock interaction. Silicification of rocks is a common feature in the Hadean and early Archean; however, the rocks studied here do not show evidence of silicification (26). Silica, which is a major component of igneous rocks, is quite immune to fluid-rock interactions if obvious secondary quartz veins are avoided. Furthermore, small variations in SiO$_2$ content do not substantially affect the TIMI. For example, a calc-alkaline rock sample (AGU25) from (29) has a TIMI of 0.66 at a SiO$_2$ concentration of 65 wt %. Even when strongly altering the SiO$_2$ concentration of this sample to 55 or 75 wt % and altering the TiO$_2$ concentration for the loss or gain of SiO$_2$ results in a slightly modified TIMI of 0.72 and 0.61, respectively.

Studies probing the fundamental differences in tholeiitic and calc-alkaline magma series ascribe the timing and extent (with respect to SiO$_2$ increase) of plagioclase and oxide crystallization as key controls on the behavior of Fe (and Ti) during magmatic differentiation (15, 40). The rate of plagioclase crystallization is important because, in the tholeiitic series, the dominance of plagioclase removal early on in crystallization passively enriches total Fe (and Ti) contents, while SiO$_2$ content of the melt remains approximately constant (15). By contrast, the presence of water and higher oxygen fugacity in the calc-alkaline magma series encourages early spinel and oxide crystallization and suppresses plagioclase crystallization, which drives the SiO$_2$ content of the melt to higher values (2, 15). Because TIMI relies on the ratio of SiO$_2$ to TiO$_2$ and the $\delta^{49}$Ti composition of a rock, it isolates the controls of oxide (changes to TiO$_2$ in melt) and plagioclase (changes to SiO$_2$ in melt) saturation and fractionation during magmatic differentiation (Fig. 2B).

The relationship between $\delta^{49}$Ti and log(SiO$_2$/TiO$_2$) for samples from the AGC is shown in Fig. 2A. The 4.02-Ga ITG granitoids fall on a similar trend to that defined by modern tholeiitic rocks (mean TIMI ITG rocks = 1.07), whereas the younger ≤3.75-Ga AGC granitoids fall close to the trend defined by modern calc-alkaline rocks (mean TIMI AGC rocks = 0.71). Of note, rocks from <3.2-Ga Barberton TTG suite align with the modern calc-alkaline rock array as well (Fig. 2B). The observed switch in the TIMI value of AGC rocks suggests that, in this region, a change in the formation mechanism of granitoid rocks occurred at 3.8 Ga. Older samples were generated via tholeiitic magmatism, whereas younger samples were produced in a calc-alkaline magmatic setting, with the latter being the dominant type of granitoid formation since that transition.
A magma series transition at ~3.8 Ga in the Slave craton

The clear compositional difference between the 4.02-Ga ITG and ≤3.75-Ga AGC granitoids (Fig. 1A) in δ\text{49}Ti can provide insight into their formation mechanisms. The two distinct TIMI values of ~1.07 and ~0.71 of the ITG and AGC, respectively, compared to TIMI values of ~1.15 and ~0.64 for modern intraplate tholeiitic and calc-alkaline rocks suggest the onset of hydrated, calc-alkaline–like magmatism and associated Fe–Ti oxide fractionation at around 3.8 Ga (Fig. 3). In the younger magmatic differentiation regime, fractional crystal- 

Proliferation in hydrated and potentially higher-pressure environment, which is consistent with existing petrological and geochemical data (22, 41). On modern Earth, nearly all calc-alkaline rocks are formed in magmatic arcs that overlie subduction zones. Therefore, a uniformitarian approach would suggest that the simplest explanation of our data is that a subduction-like setting existed on Earth at ~3.75 Ga. However, there are notable major element differences between Archean TTGs and modern arc granitoids (42). These unique Archean TTG signatures can be explained by partial melting of a down-going slab, in contrast to the modern “dehydration—mantle wedge melting—magmatic differentiation” that typically produces arc granitoids. Our Ti isotope data alone cannot discriminate between these two processes, but as both require the downward transport of hydrated material to melting depths, the mechanisms that produced ≤3.75-Ga TTGs in Acasta may have occurred in horizontal tectonic regime akin to subduction. The shift from dry magmatism, likely in a plume setting, to wet magmatism may be a short-lived regional occurrence, evidence of a pre-subduction regime (43–45), for a subduction plate tectonic (an intermediate stage between plate tectonics and stagnant lid convection) (43, 46, 47), or an alternative signature of modern-style convergent margin magmatism.

Origin of the 4.02-Ga ITG through impact-induced melting or magmatic differentiation?

In contrast to initial interpretations (21), recent work using phase equilibria modeling suggested that the 4.02-Ga ITG suite was produced via high-temperature partial melting of Fe-rich hydrated basalts at low pressures (48). Such low-pressure, high-temperature conditions were suggested to only be attainable at crustal conditions following meteorite impacts (48). Unlike fractional crystallization, single-stage equilibrium partial melting is not expected to produce large Ti isotopic fractionation because any distillation effect should be expressed in the residue rather than the melt (29, 36, 49). The maximum isotopic fractionation will occur in a single stage during incipient melting and will most likely be small if equilibrium prevails [an assumption used in partial melting calculations (50)]. Any melt produced will, with greater melt fraction, progressively assume the isotopic composition of the starting material as more Ti is transferred to the melt.

To quantitatively test whether the ITG was formed via impact-induced single-stage partial melting event or during magmatic differentiation, we calculate the Ti isotopic composition of magma and cumulate minerals during partial melting and after the onset of Ti-bearing mineral crystallization using a Rayleigh distillation model approach (Fig. 4) (29, 51). At all steps, in the model for partial melting, we calculate the remaining Ti and the δ\text{49}Ti of the residual solid, instantaneous melt, and bulk melt. Likewise, for the fractional crystallization scenario, we calculate the remaining Ti and the δ\text{49}Ti of the melt, instantaneously formed crystals, and bulk cumulate (Fig. 4).

To assess the effects of partial melting of the crust, we follow the approach of (43), meaning that we assume that TTGs are the product of partial melting of basaltic protoliths and use average Ti isotopic compositions from 2.98-Ga TTGs from the Kaapvaal craton (31) and enriched MORBs as the source (49) (Fig. 4A). This approach uses the calculated abundance of Ti-bearing minerals at various melt fractions modeled at 0.1 GPa used in trace element modeling of an impact melt event (48). We assume a Ti isotopic fractionation of +0.34‰ between melt and residue, which is the fractionation factor between residue (E-MORB) and felsic melt (TTGs) (49). This Ti isotopic fractionation factor between melt and solid residue is also consistent with that estimated using the method of (32), which is based on the correlation between FeO/TiO\text{2} and SiO\text{2} concentrations (i.e., ~0.32‰ at 1000°C and ~0.42‰ at 800°C). For low-degree partial melting, the calculated Ti isotopic fractionation is, at most, +0.34‰ (at negligible melt fraction), which is significantly lower than the δ\text{49}Ti values of +0.54 to +0.95‰ observed in the ITG samples. The heavy Ti isotopic values measured in ITG samples therefore provide evidence against the

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**Fig. 3.** Calculated TIMI for rocks from ITG and AGC (this study), same as in Fig. 2, and Barberton TTGs (36). Dashed red and blue lines show the average TIMI value for modern tholeiitic (30, 31) and calc-alkaline rocks (29, 36).

**Fig. 4.** Rayleigh distillation model of formation scenarios of the 4.02-Ga jidi whāa gneiss. (A) Partial melting of the mantle or basaltic crust scenario. The Δ\text{49}Ti\text{melt} or residue-source was calculated using the Ti isotope data of depleted MORB mantle (29, 49) and TTG data from (36) and assuming E-MORBs as source (49). Range of δ\text{49}Ti predicted by partial melting of Ti-bearing minerals at 0.1 GPa by (48) indicated by red lines. (B) Effect of Ti-bearing oxide crystallization on the evolution of magma and cumulate minerals. Δ\text{49}Ti\text{cumulate} was calculated following the approach of (29, 51) using Δ\text{49}Ti\text{melt-pyroxene} of ~0.118‰ [at T = 1125°C (29)] and ~0.39‰ [at T = 1000°C (31)]. Black line represents Earth’s mantle composition (29), and gray shaded box represents the range of Ti isotopic compositions of ITG rocks (this study). The highest δ\text{49}Ti values of the jidi whāa gneiss cannot be explained by partial melting and are most likely due to fractional crystallization.
The elevated Ti isotopic compositions of the ITG are better explained by fractional crystallization. To model the Ti isotopic composition during fractional crystallization of Ti-bearing minerals from a magma, we follow the approach of (29, 51) using oxide melt Ti isotope fractionation factors from literature data (29, 31). The larger Ti isotopic fractionation produced during magmatic differentiation ($\delta^{49}$Ti of residual melts varies from +0.00 to +1.80‰ at 99% Ti removed from the melt) can achieve values consistent with the ones we see in the ITG, suggesting that the rocks from the ITG were likely produced through fractional crystallization and not impact-induced partial melting.

### Vertical versus lateral tectonics in the AGC

The absence (or presence) of plate tectonics on Earth before 3.0 Ga is currently debated in both geochemical and modeling studies. Notably, Archean TTGs have been suggested to be formed via intraplate sagduction. Sagduction may have occurred where magma from the base of an oceanic plateau becomes detached and sinks into the mantle, thereby melting to produce deep-seated TTGs. Recent work modeling the thermal evolution of Archean crust formation shows that sagduction fails to bring water and basaltic material to the great depths required to produce a large volume of felsic crust (52). Models are still capable of TTG production via intraplate sagduction, but they yield small felsic crustal volumes (53). Given the small geographic area of exposed crust at the AGC (fig. S3), it is possible that the calc-alkaline trend observed here is a by-product of sagduction. Other geodynamic models have shown that plume-induced subduction is a possible mechanism for initiating the first subduction zone before a globally active plate tectonic regime (50). Our Ti isotope data indicate the presence of calc-alkaline magmatism, which supports the notion that hydrated crust was transported to relatively great depths. This may indicate that subduction-like tectonic behavior could have initiated on at least a local scale, early in Earth history (54), or less likely that intraplate sagduction produced a small volume of felsic crust.

### MATERIALS AND METHODS

#### Samples

To better understand the history of crystallization for the ITG and younger AGC granitoids, we have measured $\delta^{49}$Ti and $\delta^{56}$Fe values in rocks that span the Hadean-Archean transition at 4.0 Ga (see the Supplementary Materials for analytical details). We report 20 $\delta^{49}$Ti and 15 $\delta^{56}$Fe compositions of individual rock samples (see table S1) from the AGC spanning in age from 4.02 to <3.6 Ga and ranging in silica content from metagabbro ($\text{SiO}_2 = 52$ to 53 wt %) to tonalite ($\text{SiO}_2 = 57$ to 72 wt %). We also report the Ti isotopic compositions of two A-type granites from the Lachlan Fold Belt, whose petrogenesis has been proposed to be the result of melting of meta-igneous granulite left behind from a previous melt extraction (53–57), melting of granulitic crust metasomatized by mantle derived fluids (58), differentiation of anhydrous magmas (59, 60), or partial melting of tonalitic-granodioritic rocks (61, 62). See the Supplementary Materials (fig. S1) for more discussion regarding A-type granite affinities.

Calc-alkaline and tholeiitic rocks have different behavior with regard to TiO$_2$ enrichment before saturation and crystallization of Fe-Ti oxides. Typically, the TiO$_2$ content during magmatic differentiation can provide clues about the petrogenetic lineage of a sample; however, in the AGC samples, no differences are discernible (Fig. 1A). A likely reason for this is that all samples with robust zircon U-Pb ages are differentiated beyond the onset of Fe-Ti oxide crystallization (>~55 wt % SiO$_2$), at which point the trends of the tholeiitic and calc-alkaline series tend to converge (>60 wt % SiO$_2$; compare the red and blue curves in Fig. 1A). This convergence in composition illustrates the difficulty of ascribing a magma series grouping when rocks sampling a complete differentiation suite are not available.

#### Methods

Whole-rock samples used in this study have been extensively described in previous work (21, 23, 26). Aliquots of the same sample powder were used here. Twenty bulk powder samples were prepared for Ti isotopic analysis following the methods described in (51), which involves alkali-flux fusion to ensure sample homogenization and complete dissolution of Ti before ion exchange chromatography. Following flux fusion, samples were digested and double-spiked with $^{47}$Ti-$^{49}$Ti solution in a fixed proportion to Ti in the sample. We use the double-spike technique to correct the data for mass bias introduced in the instrument (63). Titanium from the dissolved samples and geostandards was then purified using ion exchange chromatography following procedures described in (64). The Ti isotopic composition of each sample and geostandard was subsequently analyzed on a Neptune multi-collector inductively coupled plasma mass spectrometer (MC-ICPMS) at the University of Chicago in medium-resolution mode to resolve interferences of $^{30}$Si/$^{46}$Ti (63). The sample measurements were introduced in the instrument using an Aridus 1 desolvating nebulizer and bracketed by standard measurements with $^{46}$Ti concentrations matched within 10% of the sample. Following a sample block of five samples, a clean acid solution was measured to correct for on-peak baseline. The samples were introduced in 0.3 M nitric and 0.005 M hydrofluoric acid. The Ti isotopic composition of each sample is reported in $\delta^{49}$Ti notation (see table S1), which is the per mil deviation of $^{49}$Ti/$^{47}$Ti ratio of each sample relative to the Origins Laboratory Ti reference material (OL−Ti).

$$\delta^{49}\text{Ti} (\text{‰}) = \left( \frac{^{49}\text{Ti}_{\text{Sample}}}{^{49}\text{Ti}_{\text{OL−Ti}}} - 1 \right) \times 1000$$

The uncertainty on each measurement was evaluated according to (65) and encompasses both the measurement session and the long-term external reproducibility. The $\delta^{49}$Ti isotopic composition of each sample was calculated from the raw data by double-spike reduction using Mathematica following methods outlined in (63). All geostandards analyzed for Ti are reported in table S1 and agree with their recommended values.

The chemical digestion and separation procedure for Fe followed the methods outlined in (66). Weights of the sample powder aliquots ranged from 2 to 70 mg for both geostandards and samples. After complete dissolution via acid digestion, the samples were purified using ion exchange chromatography. The Fe isotopic composition of each sample was then analyzed on a Neptune MC-ICPMS at the University of Chicago in medium-resolution mode to resolve interferences of ArN, ArO, and ArOH. Instrumental mass fractionation was corrected for using standard-sample bracketing. Iron isotope
TIMI applied to plutonic rocks

The $\delta^{56}$Ti composition and log(SiO$_2$/TiO$_2$) relationship relative to the TIMI of modern tholeiitic, calc-alkaline, and various granitoid rocks are plotted in fig. S2. The $\delta^{56}$Ti composition of I- and S-type granites is from (36), and the A-type granites are data presented here. Previous studies have debated the petrogenesis of A-type granites, with hypotheses ranging from melting of anhydrous granulitic met igneous residue left behind from a previous melt extraction (55–57), melting of granulitic crust metamixed by mantle-derived fluids (58), differentiation of anhydrous magmas (59, 60), or partial melting of crustal tonalitic-granodioritic rocks (56, 57). On the basis of the TIMI thresholds defined for calc-alkaline (TIMI ≤ 0.64), intermediate (TIMI = 0.64 to 1.15), and tholeiitic (TIMI ≥ 1.15) rocks, A-type granites display a tholeiitic affinity, whereas I- and S-type granites display a calc-alkaline affinity (fig. S2). On the basis of quantitative modeling of Ti isotopic compositions during partial melting and magmatic differentiation using a Rayleigh distillation model approach from (29, 51) (see main text; Fig. 4), it is likely that the A-type granites measured here formed via fractional crystallization of an anhydrous parental melt.

**MELTS modeling**

We performed Rhyolite-MELTS (63) modeling for a primitive tholeiitic and calc-alkaline magma composition and fractionally crystallized the melt starting above the liquidus temperature ranging from 1300° to 885°C at 5°C increments at 0.6 kbar (tholeiitic) and 1117° to 762°C at 1 kbar (calc-alkaline) (29). The initial oxygen fugacity of the tholeiitic and calc-alkaline melts was set to quartz-fayalite-magnetite (QFM) −0.3 (67) and +0.6 (17, 68), respectively, and the melts evolved with no oxygen fugacity buffer and at constant pressure. The MELTS calculation stops at a point where ~93 to 90% of the liquid has crystallized and the SiO$_2$ content of the remaining melt has reached ~68 and 73 wt % for the tholeiitic and calc-alkaline magma, respectively.

For estimating the Ti isotopic composition of the MELTS model outputs, we follow procedures outlined in previous studies (29–31). For calc-alkaline magmas, we use the empirically derived relationship of the stable Ti isotopic fractionation factor of $\delta^{49}$Ti$_{\text{melt-oxide}} = 0.23 \times 10^6 / T^2$ from (29), where $10^6 / T^2$ is in K$^{-2}$. For tholeiitic magmas, we use fractionation factors from (30, 31) and a multiple linear regression analysis from (31) to obtain a Ti isotopic fractionation factor of $\delta^{49}$Ti$_{\text{melt-oxide}} = (−0.5 + 0.017 \times [\text{SiO}_2]) \times 10^6 / T^2$, where $10^6 / T^2$ is in K$^{-2}$ and [SiO$_2$] is the SiO$_2$ wt % concentration of the melt. The Ti isotopic composition is modeled using a simple Rayleigh fractionation briefly outlined for tholeiitic magmas in (31).

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/50/eabc9959/DC1

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