A practical method for assigning uncertainty and improving the accuracy of alpha-ejection corrections and eU concentrations in apatite (U-Th)/He chronology

 Spencer D. Zeigler¹, James R. Metcalf¹, Rebecca M. Flowers¹
 ¹Department of Geological Sciences, University of Colorado Boulder, Boulder, CO, 80309, USA *Correspondence to*: Spencer D. Zeigler (spencer.zeigler@colorado.edu)

Abstract. Apatite (U-Th)/He (AHe) dating generally assumes that grains can be accurately and precisely modeled as geometrically perfect hexagonal prisms or ellipsoids in order to compute the

- 10 apatite volume (V), alpha-ejection corrections (F_T), equivalent spherical radius (R_{FT}), effective uranium concentration (eU), and corrected (U-Th)/He date. It is well-known that this assumption is not true. In this work, we present a set of corrections and uncertainties for V, F_T , and R_{FT} aimed <u>1</u>) at "undoing" the systematic deviation from the idealized geometry, and 2) at quantifying the contribution of geometric uncertainty to the total uncertainty budget on eU and AHe dates. These corrections and uncertainties
- 15 can be easily integrated into existing laboratory workflows at no added cost, can be routinely applied to all dated apatite, and can even be retroactively applied to published data. To quantify the degree to which real apatite deviate from geometric models, we selected 264 grains that span the full spectrum of commonly analyzed morphologies, measured their dimensions using standard 2D microscopy methods, and then acquired 3D scans of the same grains using high-resolution computed-tomography (CT). We
- 20 then compared our apatite 2D length, maximum width, and minimum width measurements with those determined by CT, as well as the V, FT, and RFT values calculated from 2D-microscopy measurements with those from the 'real' 3D measurements. While our 2D length and maximum width measurements match the 3D values well, the 2D minimum width values systematically underestimate the 3D values and have high scatter. We therefore use only the 2D length and maximum width measurements to
- 25 compute V, F_T, and R_{FT}. With this approach, apatite V, F_T, and R_{FT} values are all consistently overestimated by the 2D microscopy method, requiring correction factors of 0.74-0.83 (or 17-26%), 0.91-0.99 (or 1-9%), and 0.85-0.93 (or 7-15%), respectively. The 1s uncertainties on V, F_T, and R_{FT} are 20-23%, 1-6%, and 6-10%, respectively. The primary control on the magnitude of the corrections and uncertainties is grain geometry, with grain size exerting additional control on F_T uncertainty.
- 30 Application of these corrections and uncertainties to a real dataset (N = 24 AHe analyses) yields 1s analytical and geometric uncertainties of 15-16% on eU and 3-7% on the corrected date. These geometric corrections and uncertainties are substantial and should not be ignored when reporting, plotting, and interpreting AHe datasets. The Geometric Correction Method presented here provides a simple and practical tool for deriving more accurate F_T and eU values, and for incorporating this oft
- 35 neglected geometric uncertainty into AHe dates.

Deleted: 7	
Deleted: -	
Deleted: We then compared	
Deleted: calculated	

Deleted: We find that

Deleted: at

Deleted: (U-Th)

1 Introduction

Apatite (U-Th)/He (AHe) dating is a widely-applied thermochronologic technique used to decipher low-temperature thermal histories. In addition to analysis of parent and daughter isotopes, the conventional whole crystal (U-Th)/He method typically includes microscopy measurements of the analyzed grain. These measurements are combined with an assumed idealized grain morphology to estimate the grain volume (V) and surface area, which in turn are used to calculate three important parameters: the alpha-ejection correction (FT value), the effective uranium concentration (eU), and the equivalent spherical

- 50 radius. F_T values are required for accurate dates on crystals that are not fragments, because ⁴He atoms travel ~20 µm during α -decay and a correction is required to account for He lost by this effect (e.g., Farley et al., 1996; Ketcham et al., 2011). eU is important for accurate (U-Th)/He data interpretation because radiation damage scales with eU, which affects He retentivity (e.g., Shuster et al., 2006; Flowers et al., 2007). The equivalent spherical radius is used to approximate the diffusion domain of
- 55 whole crystals, and is a standard parameter needed for diffusion modeling (here we use a sphere with an equivalent F_T correction as the analyzed grain and refer to this parameter as R_{FT}).

It is well-recognized that there is both uncertainty and potentially systematic error associated with the microscopy approach to calculating geometric data and the parameters derived from them (Ehlers and

- 60 Farley, 2003; Herman et al., 2007; Evans et al., 2008; Glotzbach et al., 2019; Cooperdock et al., 2019; Flowers et al., 2022a). Throughout this paper we use "uncertainty" to refer to the reproducibility of measurements, and "error" to refer to a systematic deviation between a measured value and the true value (JCGM, 2012), Figure 1 shows how the commonly assigned hexagonal and ellipsoid grain geometries for apatite do not perfectly capture the true volumes and surface areas of real grains. Early
- 65 work suggested that these deviations could cause as much as ± 25% uncertainty on the F_T values for hexagonal, prismatic apatite grains of 50 μm width, decreasing to <2% for grains with cross-sections of >125 μm (Ehlers and Farley, 2003). Geometric uncertainties and systematic error have also been explored using x-ray micro- or nano-computed tomography (CT), a non-destructive method that creates 3D models of scanned objects (Herman et al., 2007; Evans et al., 2008; Glotzbach et al., 2019;
- 70 Cooperdock et al., 2019). These studies presented new, more comprehensive techniques for 2D <u>apatite</u> grain measurements (the 3D-He method of Glotzbach et al., 2019) and proposed a method to routinely acquire CT data for all dated apatite grains (Cooperdock et al., 2019).

<u>Rigorous quantification of uncertainties and corrections for systematic error on the geometric</u>

- 75 parameters are required to represent and interpret AHe data accurately. For example, appropriate uncertainties on single-grain dates are important for deciding if data are normally distributed and thus reasonable to represent and model as a mean sample date, or if the data are "overdispersed" (e.g., Flowers et al., 2022b). Similarly, appropriate uncertainties on other parameters such as eU are needed to properly decipher AHe date vs. eU patterns. However, despite the past work addressing geometric
- 80 uncertainties (e.g., Cooperdock et al., 2019; Glotzbach et al., 2019), the uncertainties on the grain's geometric information are not typically propagated into the uncertainties of the derived parameters (e.g., eU concentration, corrected (U-Th)/He date). Nor are data systematically corrected for potential error associated with grain measurements. This is largely because uncertainty and error in the geometric

Deleted: Conceptually, Deleted: can lead to deviations from the

	Moved (insertion) [1]
(Deleted: However, ri
~(Deleted: (U-Th)/

Deleted: 1	
Deleted: D	
Deleted: this	

parameters depend in large part on how much the real grain geometry deviates from that assumed, which may vary from grain to grain, depending on grain morphology, as well as possibly on grain size and other parameters. Moreover, although both the 3D-He method (Glotzbach et al., 2019) and the routine CT analysis approach (Cooperdock et al., 2019) would improve the accuracy and precision of geometric parameters, both add more time to the (U-Th)/He dating process, and in the case of the latter,

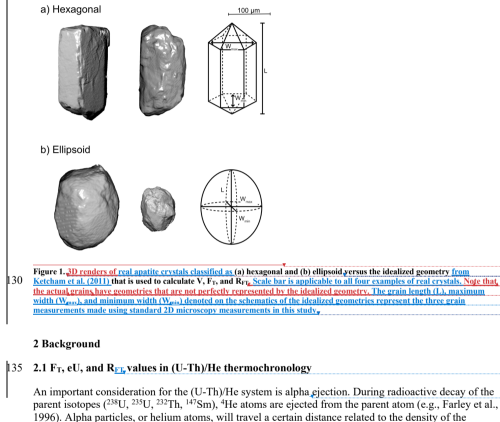
requires regular access to CT instrumentation.

To address this problem, we present a time-efficient and straightforward "geometric correction" method to routinely correct for systematic error and to assign uncertainties to F_T, eU, and R_{FT} values for the full spectrum of regularly analyzed apatite grain sizes and morphologies. This approach requires no

- 100 spectrum of regularly analyzed apatite grain sizes and morphologies. This approach requires no additional work or cost beyond what is already done as part of most existing (U-Th)/He dating workflows. Nor does it necessitate additional microscopy measurements or routine CT analysis of grains, so it is easily adoptable by any lab or data user. Additionally, this method can be applied retroactively to previously collected data, even after the grains themselves have been dissolved and are
- 105 no longer available for additional work. We first developed a simple classification system for apatite grains of varying <u>shape</u> and surface roughness. For 237 apatite crystals characterized by a wide range of morphology, size, age, and lithologic source, we then compared V, F_T, and R_{FT} estimates calculated from 2D microscopy measurements with those determined by CT scans of the same grains at 0.64 μm resolution. We use these data to derive corrections for systematic error and to determine uncertainty
- 110 values that can be applied to 2D V, F_T, and R_{FT} values depending on the geometry and size of the analyzed apatite. These outcomes allow analysts to 1) correct geometric parameter values for systematic error, 2) propagate the F_T uncertainty into the reported uncertainty on corrected (U-Th)/He dates, 3) propagate the V uncertainty into the reported uncertainty on eU values, and 4) report R_{FT} value uncertainties that have potential to be included in thermal history modeling. We conclude by illustrating
- 115 this approach with real (U-Th)/He data and discuss the implications for the accuracy and precision of (U-Th)/He datasets more broadly.

Moved up [1]: However, rigorous quantification of uncertainties and corrections for systematic error are required to represent and interpret (U-Th)/He data accurately. For example, appropriate uncertainties on single-grain dates are important for deciding if data are normally distributed and thus reasonable to represent and model as a mean sample date, or if the data are "overdispersed" (e.g., Flowers et al., 2022b). Similarly, appropriate uncertainties on other parameters such as eU are needed to properly decipher AHe date vs. eU patterns.⁴

Deleted: morphology



1996). Alpha particles, or neutim atoms, will traver a certain distance related to the density of the mineral through which they travel and the ejection energy from the parent atom. For apatite, the average stopping distances for ²³⁸U, ²³⁵U, ²³²Th, and ¹⁴⁷Sm are 18.81 μm, 21.80 μm, 22.25 μm, and 5.93 μm respectively (e.g., Ketcham et al., 2011). If the parent atom is positioned within the ejection range of the grain edge, then the He atom has a non-zero chance of being ejected from the crystal entirely. The

probability of retention increases with increasing distance of the parent from the grain edge. Overall, the smaller the grain, the higher the surface area to volume ratio of the grain, and the greater percentage ofHe that is lost via the ejection process.

and a second	a) Hexagonal				
	b) Ellipsoid				
	\longrightarrow $\xrightarrow{\iota}$				
	Apatite grain selected Idealized (2D) ell for analysis geometry				
	Deleted:				
J	Deleted: Examples of				
h	Deleted: realistic				
2	Deleted: apatite				
	Deleted: geometries				
\sum	Deleted: rains				
$\langle \rangle$	Deleted: (left column				
	Deleted:)				
	Formatted: Not Superscript/ Subscript				
	Formatted: Not Superscript/ Subscript				
	Formatted: Not Superscript/ Subscript				
	Formatted: Not Superscript/ Subscript				
	Formatted: Not Superscript/ Subscript				
	Formatted: Subscript				
	Formatted: Subscript Deleted: (middle column). In the right column, the 'real' grain				
$\left \right $	is overlaid onto the idealized geometry in red. The shaded [1]				
Deleted: For both hexagonal and ellipsoid shapes, the ideal geometry overestimates the real apatite volumes.					
11	Deleted: s				
$\langle \rangle \rangle$	Deleted: -				
()	Deleted: and				
	Deleted: and				
	Deleted: , and for ¹⁴⁷ Sm it is				
1	Deleted: 5.03 um				

To obtain an intuitively more meaningful date, (U-Th)/He dates on crystals that retain their original grain edge are typically corrected for the He lost by alpha ejection to obtain a "corrected (U-Th)/He

- date". This alpha-ejection correction (or F_T value) is the fraction of He that is retained in the crystal, such that an F_T value of 0.70 means that an estimated 30% of He was lost from the crystal by ejection. F_T is typically calculated based on the stopping distances of He in each mineral for each parent isotope, the proportion of the parent isotopes, the crystal dimensions, and an assumed idealized crystal geometry
- 175 that enables one to use the crystal measurements to estimate the surface area and volume of the crystal (Farley et al., 1996). F_T corrections typically assume a uniform distribution of parent isotopes; parent isotope zonation in crystals can introduce additional uncertainty into the F_T correction (Farley et al., 1996; Meesters and Dunai, 2002; Hourigan et al., 2005). Additional uncertainty can also arise for broken or abraded crystals, where the magnitude of the appropriate correction can be unclear_(Rahl et
- 180 al., 2003; Brown et al., 2013; He and Reiners, 2022),

¢U₄ is important for (U-Th)/He thermochronology because it can be used as a proxy for radiation damage, which can have a large effect on the mineral He retentivity (e.g., Shuster et al., 2006; Flowers et al., 2007). Radiation damage can cause positive correlations between AHe date and eU for thermal

185 <u>histories characterized by slow cooling, partial resetting, or long residence in the helium partial retention zone.</u> Accurate eU values depend on accurate grain volumes, because volumes are used to calculate grain masses, which in turn are used to compute parent isotope concentrations and eU (e.g., Flowers et al., 2022a).

- The equivalent spherical radius is relevant for (U-Th)/He thermochronology because mineral diffusion depends on grain size. Grain size is therefore included in the diffusion modeling used to decipher thermal histories from (U-Th)/He data. The equivalent spherical radius parameter can be reported either as a sphere with the same surface area to volume ratio as the analyzed grain (R_{SV}), or as a sphere with the same F_T value as the analyzed grain (R_{FT}, Ketcham et al., 2011; Cooperdock et al., 2019). Use of
- 195 R_{FT} is preferred, because during thermal history modeling this value yields outcomes more similar to those using the real 3D grain geometries (Ketcham et al., 2011).

2.2 Use of CT for $F_{T},\,eU,\,and\,R_{FT}$ value determinations

Computed tomography (CT) is a high-resolution (sub-micrometer), non-destructive, 3D imaging technique based on the attenuation of x-rays through a sample. 2D cross sections ('slices') of the sample are created as x-rays pass through the sample and are then processed into 3D models. These models can be analyzed with software like Drogonfly and Plob2D to arteset high quality 3D geometric data like

be analyzed with software like Dragonfly and Blob3D to extract high quality 3D geometric data like volume and surface area (Ketcham, 2005; Dragonfly, v.2020.2).

CT has been applied to improve the accuracy of geometric parameters in (U-Th)/He chronology in four studies (Herman et al., 2007; Evans et al., 2008; Glotzbach et al., 2019; Cooperdock et al., 2019). Initial work used CT data at a 6.3 μm resolution to derive F_T values for 11 irregularly shaped detrital apatite grains (Herman et al., 2007). This study then dated the crystals by (U-Th)/He and combined the 3D CT models of the dated grains with an inversion algorithm to constrain a range of thermal histories.

	Deleted: (Rahl et al., 2003; Brown et al., 2013).
	Deleted: The effective U concentration (
5	Deleted:)
	Deleted: in the case of
ı	
1	
er	
	Deleted: ; Flowers et al., 2022a
ole	
ın	
	Deleted: dimensional and

The subsequent studies have directly compared geometric parameters determined from 2D microscopy data with 3D CT measurements of the same grains. Evans et al. (2008) <u>calculated "effective F_T " values for 9 euhedral to subhedral, detrital and volcanic apatite and zircon grains using CT scans at 3.8 µm resolution and eroding the outer 20 µm of the scanned grain in 3D_r Glotzbach et al. (2019) developed an</u>

- 220 improved microscopy method, called the 3D-He approach, to estimate F_T values using dimensions measured from a suite of photomicrographs to simulate a 3D grain model. They acquired CT data at 1.2 μm resolution for 24 apatite grains, including rounded, pitted, broken, anhedral, subhedral, and euhedral crystals. Cooperdock et al. (2019) presented a method for regular CT characterization of grains at 4-5 μm resolution and acquired CT data for a suite of 109 high quality euhedral apatite crystals from two
- 225 plutonic samples. These three studies found that the 2D data variably over- or under_estimated the 3D data for V, F_T, and R_{FT}, and estimated a range of scatter for the different parameters. These previous results are discussed in greater detail in Sect. 6.2 where we compare the outcomes of our study with this past work.

3 Selecting and Characterizing a Representative Apatite Suite

230 3.1 Strategy

235

We designed our study to ensure that we captured the range of representative apatite crystals commonly dated by the (U-Th)/He method. Our goal was to include the full spectrum of grain qualities in realistic proportions so that the study outcomes are relevant for the complete range of routinely analyzed grains rather than being biased to apatite morphologies specific to a single sample type. As described in more detail below, grain selection focused primarily on including crystals from samples encompassing a

spectrum of lithology and age (Sect. 3.2), with a range of sizes (Sect. 3.3), and with variable morphology (Sect. 3.4). We ultimately selected 400 apatite grains for analysis, from which we obtained high-quality CT data for 264 crystals.

3.2 Selecting a Representative Sample Suite

- Apatite grains were selected from eight samples that include six igneous and metamorphic rocks and two clastic sedimentary rocks with ages from Oligocene to Archean (Table 1). All samples were separated using standard crushing, density, and magnetic separation techniques. Most samples were dated previously by AHe, in the CU TraIL (Thermochronology Research and Instrumentation Lab). The Oligocene Fish Canyon Tuff (sample FCT) from the San Juan Mountains in Colorado, USA is
- commonly used as a (U-Th)/He reference standard, with AHe dates younger than emplacement (e.g., Gleadow et al., 2015). The Eocene granitic Ipapah pluton is from the Deep Creek Range (sample DCA) of east-central Nevada, USA and yields Miocene AHe dates (unpublished data). The Cretaceous Whitehorn granodiorite (sample BF16-1) is from the Arkansas Hills in Colorado, USA and has Eocene AHe dates (Abbott et al., 2022). The Cambrian McClure Mountain syenite (sample MM1) from the Wet
 Mountains of south-central Colorado yields Mesozoic AHe dates (Weisberg et al., 2018). A Proterozoic
- granitic dike from the Baileyville drill core (sample Bail933) in northeastern Kansas, USA js



Dele	ted: scanned
Dele	ted: a
Dele	ted: , used the CT data to
Dele	ted: e
Dele	ted: and recalculated the "effective Fr" (Evans et al. 2008)

D	eleted: and to
D	eleted: avoid
De	eleted: ing
D	eleted: the results towards
D	eleted: lithology or age
D	eleted: 7

-(Deleted: apatite (U-Th)/He (
(Deleted:)
(Deleted: h
(Deleted: h
(Deleted: yield
(Deleted: h
(Deleted: h
(Deleted: yields

<u>characterized by</u> Paleozoic AHe results (Flowers and Kelley, 2011). An Archean gneiss from the Superior craton in Canada (sample C50) yields Cambrian AHe dates (TraIL unpublished data). The two detrital samples (samples 16MFS-05 and 15MFS-07) have Cretaceous depositional ages, are from the Kaikoura Range on the South Island of New Zealand, and <u>have</u> late Miocene to Pliocene AHe dates (Collett et al., 2019; Harbert, 2019).

Table 1. Apatite sample information,

275

Sample				Longitude	Latitude	GEM		
Name	Unit and Lithology	Sample Age	Locality	(*W)	(°N)	Categories	N ^a	Additional Geochronologic and Thermochronologic Data
	Fish Canyon Tuff		San Juan Mountains,					Zircon U-Pb 28.172 ± 0.028 Ma (2s) (Schmitz and Bowring, 2001); AHe 20.8 ± 0.4 Ma (1s) (Gleadow et al
FCT	dacite	Oligocene	Colorado, USA	-106.93	37.76	A1, A2, B1	30	2015) 2001); AHE 20.8 ± 0.4 Ma (15) (Gleadow et a
	Ipapah		Deep Creek Range East-			A1, A2, B1,		Zircon U-Pb 39 Ma ± 1 Ma (Rodgers, 1989); AHe 14.3-9
DCA	monzogranite	Eocene	Central Nevada, USA	-113.92	39.83	B2, C2	30	Ma* (TRaIL unpublished data)
	Whitehorn		Arkansas Hills, Colorado,			A1, A2, B1,		Zircon U-Pb 67.31 Ma + 0.57/-0.78 Ma (2s) (Abbey et a
BF16-1	granodiorite	Cretaceous	USA	-105.90	38.50	B2	25	2017); AHe 47.4 ± 4.2 Ma (1s) (Abbott et al., 2022)
								Hornblende 40 Ar/ 39 Ar 523.2 \pm 0.9 Ma (1s) (Spell and
мм	McClure	Cambrian	Wet Mountains, South- Central Colorado, USA	-105.47	38.35	A1, A2, B1, B2	36	McDougall, 2003); AHe 150-70 Ma* (Weisberg et al. 2018)
IVIIVI	Mountain syenite	Cambrian	Central Colorado, USA	-105.47	38.33	BZ	30	2018)
	Baileyville drill					A1, A2, B1,		Zircon U-Pb ca. 1400 Ma (Van Schmus et al., 1987); A
Bail	core granitic dike	Proterozoic	Northeast Kansas, USA	-96.20	39.90	B2	22	150-70 Ma* (Flowers and Kelley, 2011)
	Superior Craton					A1, A2, B1,		Zircon U-Pb 2720-2680 Ma (Hoffman, 1988); AHe 559
C50	tonalitic gneiss	Archean	Superior Craton, Canada	-92.99	51.76	B2	47	461 Ma* (TRaIL unpublished data)
	Marlborough							
	Fault System		Kaikõura Ranges, South			A1, A2, B1,		Deposition 100 ± 20 Ma (1s) (Rattenbury et al., 2006
16MFS-05	sandstone	Cretaceous	Island, New Zealand	173.69	-42.29	B2, C1, C2	45	AHe 4.2 ± 0.6 Ma (1s) (Collett et al., 2019)
	Marlborough Fault System		Kaikõura Ranges, South			B1, B2, C1,		Deposition 120 ± 22 (1s) (Harbert, 2019; Mean AHe 5.4
15MFS-07	grevwacke	Cretaceous	Island, New Zealand	173.22	-41.78	C2	29	0.2 (1s) (Harbert, 2019)

280 "The number of grains for which high quality CT data were acquired. Not all grains in this dataset were included in the regressions; see Sect. 4.4.

3.3 Selecting a Representative Crystal Size Distribution

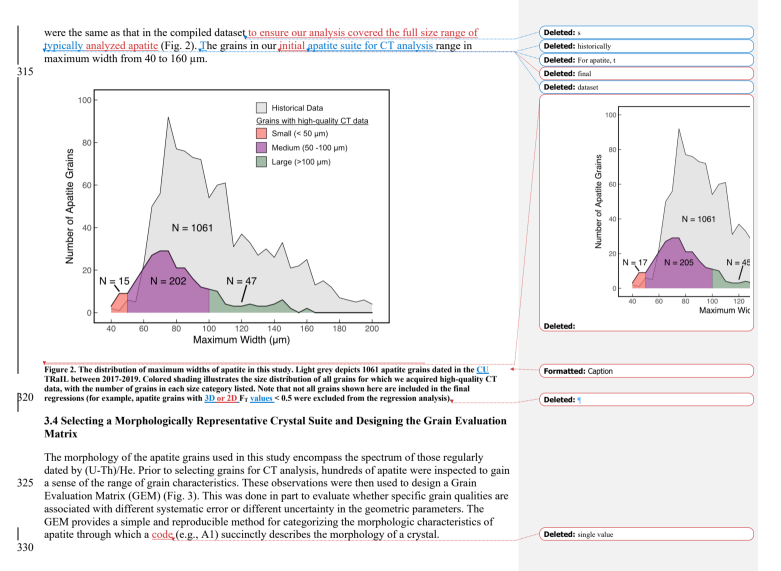
The size distribution of grains analyzed in this study is based on the size distribution of apatite grains routinely analyzed for (U-Th)/He dates. We first plotted the maximum width (the larger of the two measured widths; Fig. 1; see also Sect. 4.2) of apatite grains (N = 1061; Fig. 2) analyzed in the CU TraIL over a two-year period. The grains in this dataset were from a variety of sources and were selected and measured by TRaIL staff, TRaIL students, and visitors. Our analysis focused on crystal width because the smaller grain dimension (i.e., the width) is the chief control on alpha ejection due to the long stopping distances of alpha particles. Maximum width was used because for apatite it can be particularly difficult to reliably and accurately measure the minimum width (see Sect. 4.2 and 5.1). These lab analyses were subdivided into small (< 50 μm maximum width), medium (50-100 μm maximum width) and large (>100μm maximum width) size categories (shading in Fig. 2). From the samples described above we then picked suites of apatite crystals for CT with size distributions that

7

(C	Deleted: r
	Deleted: r
	Deleted: st
(Deleted: -
(Deleted: second (
	Deleted:)
-(C	Deleted:
· (C	Deleted: see
(Deleted: Appendix _
(Deleted: accurately
-	

Deleted: h	Deleted: h								
Deletea: n									
Deleted: yield									
Deleted: h									
Deleted: et al, in re	view								
Deleted: .									
Sample Name	Sample Age	Lithology	Locality						
Fish Canyon Tuff	Oligocene	Dacite	San Juan Mountains, C						
Deep Creek	Eocene	Monzogranite	Deep Creek Range East-Ce						
Whitehorn	Cretaceous	Granodiorite	Arkansas Hills, Colo						
McClure Mountain	Cambrian	Syenite	Wet Mountains, South-Cen						
Baileyville drill core	Proterozoic	Granite	Northeast Kansa						
Superior Craton	Archean	Tonalite	Superior Craton,						
	Cretaceous	Sandstone	Kaikõura Ranges, South Isla						
Mariborough Fault System	cretaceous								

Dalahada



340 345	The GEM has two axes (Fig. 3): a "geometric classification" x-axis and a "roughness index" y-axis. Geometry and surface roughness were chosen for the GEM because apatite inspection revealed that these are the morphological features most likely to contribute to a grain's deviation from the idealized hexagonal or ellipsoid geometry used to calculate 2D geometric parameters. In the GEM, geometry is described as A (hexagonal), B (sub-hexagonal), or C (ellipsoid), where A and B grains assume a hexagonal geometry and C grains an ellipsoidal geometry for 2D calculations (Ketcham et al., 2011). Surface roughness is described as 1 (smooth) or 2 (rough).	(Deleted: tical
350	Grains with missing terminations are sometimes analyzed by (U-Th)/He, so a subset of grains with one or two missing terminations was selected for CT analysis. For apatite, grains with missing terminations are approximately similar in proportion to those in the overall apatite sample suite.		
I	For each apatite GEM category, grains from at least two samples and as many as eight samples were selected for CT analysis to ensure a range of subtle differences among grain types (Fig. B1). The number of grains selected for CT analysis in each GEM category was approximately proportional to the abundance of grains in that category in the entire sample suite. For example, because B1 (sub-	(Formatted: Normal (Web)
355	hexagonal, smooth) apatite crystals were more common than C2 (ellipsoid, rough) crystals in the apatite suite, more B1 than C2 apatite were analyzed by CT.	(Deleted: 1
360	Grain roughness (the y-axis of the GEM) was ultimately determined to have no bearing on the corrections or uncertainties <u>derived in this study</u> . Despite this, the GEM retains this axis because the GEM is a simple, coherent, and consistent tool for identifying and communicating grain characteristics that can influence the (U-Th)/He date. Noting the roughness of an apatite grain is useful for evaluating overall sample quality and can aid in identifying and evaluating dispersion in a (U-Th)/He dataset. The GEM provides a way to easily report the overall morphologic character and quality of dated apatite grains. It also is a useful teaching tool to show newcomers to mineral picking the wide variety of morphologies possible for apatite grains.		Deleted: reported Deleted: the Deleted: Additionally, the Deleted: is Deleted: al Deleted: in studies reporting apatite (U-Th)/He data and Deleted: grain
			Deleted: for newcomers to mineral picking

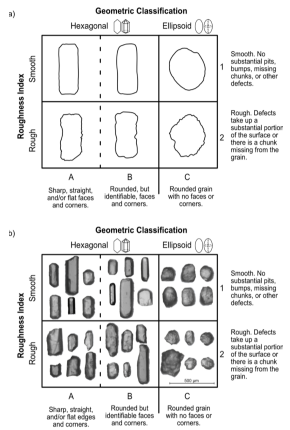


Figure 3. The apatite Grain Evaluation Matrix (GEM) in (a) schematic form and (b) with images of real grains analyzed in this study. The geometric axis classifies grains as A (hexagonal), B (sub-hexagonal), or C (ellipsoid). Both A and B apatite grains assume an idealized hexagonal prism geometry while C apatite grains assume an idealized ellipsoid geometry for 2D calculations (Ketcham et al., 2011). The roughness index classifies grains as 1 (smooth) or 2 (rough). Grains can be described by combining a geometric value and a roughness value (eg. A1, B2).

4 Measurement and Data Reduction Methods

4.1 Strategy

The goals of this work are to develop corrections for systematic error and assign appropriate

- 385 uncertainties to conventional "2D" microscopy estimates of the geometric parameters by comparing 2D values with "3D" values derived from CT data. To do this we first measured our suite of representative apatite crystals using the 2D microscopy approach (Sect. 4.2) and then acquired high-resolution (0.64 µm) CT data for these grains (Sect. 4.3). We then examined the 2D-3D relationships, linearly regressed them to determine corrections depending on grain geometry that make the 2D measurements as close to
- 390 the 3D values as possible, and calculated uncertainties (Sect. 4.4). This analysis assumes that the 3D values are accurate (Sect 4.3). The final corrections and uncertainties are most appropriate for grains with characteristics like those used in this calibration study, with geometries like those in Figure 3 and microscopy measurements and 2D calculations done as described below. The apatite grains in this work have length/maximum width ratios of 0.8-3.6 and maximum width/minimum width ratios of 1-1.7, FT
- 395 uncertainties include only those uncertainties associated with grain geometry and not those due to parent isotope zonation, grain abrasion, or crystal breakage.

4.2 Microscopy measurements and 2D calculation methods

Apatite grains were hand-picked under a Leica M165 binocular microscope under 160X magnification. Each grain was photographed on a Leica DMC5400 digital camera, manually measured using <u>either</u> the

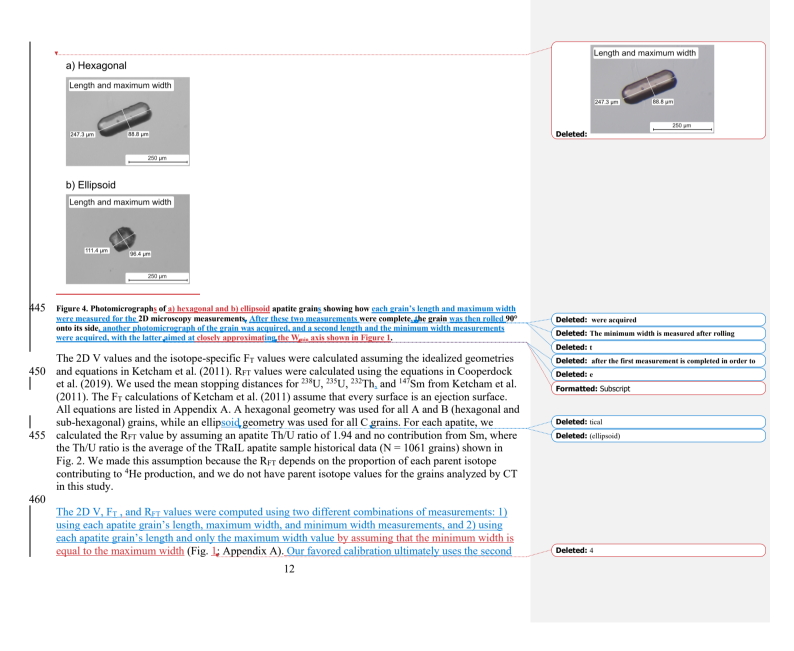
- 400 Leica LAS X or Leica LAS 4.12 software, and assigned a GEM value (Fig. 3). The calibration of the software was checked before, during, and after the measurements using a micrometer. The measurement procedure consisted of first identifying the long axis of the apatite grain parallel to the c-axis, then identifying the apatite's maximum width that is perpendicular to the grain length, acquiring a photograph using the Leica digital camera, and then measuring the length and maximum width using
- 405 the Leica software (Fig. 4), This was followed by rolling the grain 90° to identify the apatite grain's minimum width, acquiring another grain photograph, and again measuring the long and short axes using the Leica software to obtain a second length measurement and the grain's "minimum width". Determining and measuring the apatite's minimum width accurately is challenging owing to the difficulty of rotating and stabilizing the grain for a photograph while the grain is balanced on an axis
- 410 without a flat surface, For rounded grains (GEM C, ellipsoid idealized geometry), the length and widths can be difficult to identify. Our microscopy measurement method is similar to that used in many labs, although a common practice is to acquire only one grain image and thus only a single width measurement (e.g., Cooperdock et al., 2019).
- 415 We find a typical <u>2D</u> measurement uncertainty of 2.8 µm at 1s standard deviation. This was determined based on repeat measurements by 3 individuals of 258 apatite grains using the same images and software for each grain. Each individual measured both lengths and the maximum width of each grain, for a total of 774 measurements per person. The 1s sample standard deviation for each grain dimension was calculated, with an average standard deviation of 2.8µm.

Deleted: The final corrections and uncertainties are most appropriate for grains with characteristics like those used in this calibration study, with geometries like those in Figure 3, axial ratios < 1.7, maximum widths of 50-160 µm, and with microscopy measurements and 2D calculations done as described below.

Deleted: concentration uncertainties or

(Deleted: a GEM value assigned
(Deleted: initial
(Deleted: measuring
	Deleted: and short axes of each grain to obtain the grain's length and "maximum width" (Figure 4).", followed by rolling the grain 90° and again measuring the long and short axes to obtain a second length measurement and the grain's "minimum width". but usually
(Deleted: is identified first and the length
Ć	Deleted: assumed to be
	Deleted: at
Ĉ	Deleted: i
Ĉ	Deleted:
Ĉ	Deleted: its narrowest axis
Ĉ	Deleted: especially

De	leted	crystals



approach owing to the difficulty of measuring the minimum width accurately, as discussed further 475 below (Sect. 5.1, 6.1).

4.3 Nano-computed tomography and 3D calculation methods

After 2D measurements, apatite grains were mounted for CT. Crystals were mounted in an ~600 x 600 μm area on a thin, 2000 μm wide plastic disc that was hole-punched from a plastic sheet protector and 480 then covered with double sided tape (Fig. 5). Each plastic disc was constructed with a 0.025 mm diameter wire running down the center to act as a point of orientation to aid in the identification of

grains post-scan. It was later discovered that the high-density wire created challenges for data reduction, so this approach is not recommended for future studies. Each plastic disc held 4-10 grains and 5-6 discs were stacked vertically to create a mount (Fig. 5). Mounts were secured by a thin layer of parafilm, 485 attached to a 1-2 mm thick rubber cylinder for stabilization, and then glued to the top of a flat-head pin

(Fig. 5).

Each mount was scanned on a Zeiss Xradia 520 Versa X-ray Microscope in the University of Colorado Boulder Materials Instrumentation and Multimodal Imaging Core (MIMIC) Facility. Scanning

490 parameters were optimized to reduce noise and scanning artifacts during test scans of the first mount. Scanning parameters were kept constant for subsequent mounts. All mounts were scanned with the 20X objective at relatively low power and voltages with small distances between the mount, source, and detector, which allowed for high resolution (0.64 µm). Table B1 reports the scan parameters.



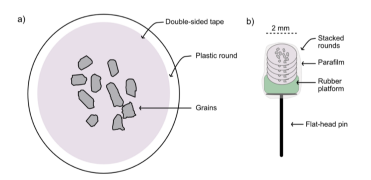


Figure 5. Schematic showing (a) an individual plastic round and (b) a final grain mount for CT analysis. The wire is not shown because it should not be included in future studies. Grains are placed onto a ~2 mm wide sturdy plastic disc (hole punched from a plastic sheet protector) covered with double-sided tape. Each plastic round can hold between 4-10 grains. Rounds are stacked on 500 top of each other and placed on a rubber platform cut from old test tube stoppers, which is glued to a flathead pin and covered with Parafilm.

13

Deleted: However, given Deleted: see
Deleted: see
Deleted: 6.2.
Deleted: Tabl
Deleted: , we ultimately used the grain's length and maximum width to calculate the 2D values Results using both the minimum and maximum width are included in Appendix E.
Formatted: Font: 12 pt

Deleted: of rubber Deleted: head

Raw CT data were imported into Blob3D (Ketcham, 2005; freely distributed software) to calculate the dimensions, V, surface area, and isotope-specific F_T values for each grain. First, the grains were segmented, or separated, from the matrix, noise, and other grains, such that each grain was a separate

- 515 'blob', Segmentation was done with Dragonfly software, Version 2020.2 for Windows (Dragonfly, 2020) due to the complex nature of the artifacts arising from the use of the wire. After segmentation, the 3D V, surface area, and F_T values were calculated by Blob3D. Blob3D calculates grain dimensions by first identifying the shortest caliper dimensions (Box C), then measuring the shortest dimension orthogonal to it (Box B), and finally measuring the dimension orthogonal to both (Box A) (Ketcham,
- 520 2005; Cooperdock et al., 2019). These dimensions generally correspond to the minimum width (Box C), the maximum width (Box B) and the length (Box A) of a regularly shaped apatite (Cooperdock et al., 2019). Blob3D calculates V by counting the number of voxels (3D-pixels) in the segmented object and multiplying that number by the volume of each voxel. Blob3D calculates surface area by summing the faces of the isosurface surrounding the grain voxels and then smoothing it to reduce the effects of
- 525 pixelation caused by the cubic voxels (Ketcham, 2005; Cooperdock et al., 2019). Blob3D calculates the ²³⁸U, ²³⁵U, ²³²Th, and ¹⁴⁷Sm F_T corrections using a Monte Carlo approach that randomizes the starting location of an alpha particle within the selected volume of an object. The direction of ejection of the alpha particle is calculated via uniform sampling (Ketcham and Ryan, 2004). Blob3D uses stopping distances as reported in Ketcham et al_x(2011) and assumes that ejection occurred across all surfaces.
- 530 Like for 2D R_{FT} values, we calculated 3D R_{FT} values using the equations of Cooperdock et al. (2019) and assuming a Th/U ratio of 1.94 based on TRaIL apatite sample historical data.

In order to confirm our assumption that the CT measurements are representative of the "real" grain measurements, we assumed a \pm 1% uncertainty in our CT measurements, based on preventative

maintenance measurements performed by the MIMIC lab and technicians, and performed simulations in Blob3D by varying the voxel size similar to those done in Cooperdock et al. (2019). Like Cooperdock et al. (2019), we find that uncertainties in the CT data translate to negligible differences in the relevant values output by Blob3D and are vanishingly small compared to the uncertainties in the 2D measurements.
 540

Some <u>apatite grains</u> were removed from the final dataset owing to issues during CT scanning or subsequent data processing. Due to the use of the 20X objective for high resolution, many of the original 400 grains were lost because the edges of grains were 'cut off' during scanning. Additionally, the high-density wire in the apatite mounts introduced challenges for data reduction, like 3D models that had large holes or complex surface artifacts. The final dataset after removal of the grains with

545 that had large holes or complex surface artifacts. <u>The final dataset after removal of the grains with problematic analytical results</u> consists of 26<u>4</u> apatite grains with high-quality CT data.

4.4 Statistical comparison of 2D and 3D values

The first step in our 2D-microscopy vs. 3D-CT data comparison was to generate scatter plots of 3D vs. 50 2D data for all 264 apatite grains in our dataset. Figure 6 shows these plots for length, maximum width, Deleted: that had its V, surface area, and F_T values calculated

Deleted: final 3D parameters

Formatted: Subscript

Deleted:

Formatted: Font. (Delault) + body (Times New Roman)	_
Formatted: Font: (Default) +Body (Times New Roman)	
Formatted: Font: (Default) +Body (Times New Roman), Font color: Text 1	
Formatted: Font: (Default) +Body (Times New Roman), 1 pt, Not Bold, Font color: Text 1	2
Formatted: Font color: Text 1	
Deleted: W	
Deleted: ou	
Deleted: any	
Deleted: present	
Formatted: Font color: Text 1	
Formatted: Font: (Default) +Body (Times New Roman), Font color: Text 1	
Formatted: Font color: Text 1	
Deleted: data points	
Deleted: due	
Deleted: This	
Deleted: 7	

Deleted: 7

and minimum width. Figure 7 includes these plots for V, isotope-specific F_T, and R_{FT} values, In Figure 7, we show only the isotope-specific ²³⁸U F_T value for illustrative purposes because ²³⁸U dominates the

565 ⁴He production budget, <u>However</u>, we plotted and regressed the data for the ²³⁵U, ²³²Th, and ¹⁴⁷Sm isotope-specific F_T values in the same manner <u>as for the ²³⁸U F_T value and include those plots in Figure</u>, C1, We did not examine surface area separately because although it is used together with volume to determine the F_T value, it is not alone used to calculate any other geometric parameter (unlike volume, which is used to calculate concentrations).

570

Our next step was to carry out regressions of the 3D vs. 2D data for V, isotope-specific F_T, and R_{FT}

values. On the 3D versus 2D plots, if the data fall on the 1:1 line (bold black line), then no correction for systematic error is needed for the 2D data because the 2D data are in agreement with the 3D data. If the data fall off the 1:1 line, then the correction desired for the 2D data can be viewed as the offset of

- 575 the data and its linear regression line from the 1:1 line. To determine corrections for systematic error, ordinary least squares linear regression with the intercept fixed at the origin was used. We explored several regression approaches, but ultimately chose an unweighted approach because the scatter of the 2D data that we wish to characterize includes both the uncertainty on the grain length and width measurements and other factors such as surface roughness and deviation from the assumed idealized
- 580 grain geometry. We also explored fixing versus not fixing the y-intercept at (0,0). Here we present only the results of regressions with the y-intercept fixed at 0, because the unconstrained regressions generally yield intercepts within uncertainty of 0 and we would expect that if 2D measurement of any parameter was 0, then the 3D value would also be 0.
- 585 We <u>ultimately</u> excluded from the regressions the <u>apatite</u> (N = 27) with 3D F_T values <0.5, which are grains smaller than those typically analyzed by (U-Th)/He. <u>This was done to avoid biasing the corrections and uncertainties with data for grains that are not representative of regularly dated apatite</u>. This exclusion resulted in the elimination of all "small"-sized grains with <50 µm maximum width from the regressions. These small grains are characterized by greater differences between 2D and 3D values and higher scatter than the medium- and large- sized grains in our dataset, as shown by the grey points</p>
 - in Figure 7, The final regressed dataset has 237 apatite grains.

To evaluate if different groups of grains have statistically different slopes (and thus should have different corrections applied to them) we used Tukey's test (Table C1). Separate linear regressions were

- done for grains that use different geometric assumptions, so hexagonal apatite (A and B grains in Figure 3) were regressed separately from ellipsoid apatite (C grains in Fig_{*}3). The slopes for the linear regressions of these two groups are statistically distinguishable, justifying their separation by geometry. Linear regressions were also done by grouping by surface roughness (1 vs. 2 on the GEM, Fig_{*}3) and size (medium, large). The linear regression slopes for these different categories are each statistically
- 600 indistinguishable, indicating it is reasonable to only group the data by geometry for all parameters (Table C1).

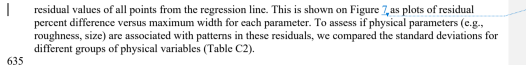
The uncertainty for each 2D geometric parameter is the scatter of the points about the regression line. To determine the uncertainty of each 2D parameter, we calculated the 1s standard deviation of the

~	
•	Deleted: volume
•	Deleted: (Fig. 6)
•	Deleted: W
•	Deleted: , but we
(Deleted: treated
Ć	Deleted: ²
Ć	Formatted: Superscript
C	Deleted: (
C	Deleted: .
r	Deleted:)

(Deleted: plots and
(Deleted: grains
(Deleted: 8
	Deleted: 2D and
	Deleted: Despite initially including these small grains in our analysis, we ultimately excluded them because we wanted
1	Deleted: for the routinely dated apatite
	Deleted: not
	Deleted: typically analyzed
	Deleted: Overall,
1	Deleted: t
1	Deleted: 6
1	Deleted: , and we wanted to avoid biasing the corrections and uncertainties for the routinely dated apatite with data for atypically analyzed grains.

Deleted: ure

Deleted: ure



The correlation of isotope-specific F_T uncertainties was also evaluated because we expect them to be highly correlated (Martin et al., 2023). The correlation coefficient between each isotope-specific F_T was

Deleted: 6

Deleted:

Deleted: (Martin et al., in review)

Deleted: R

Deleted:	grain
Deleted:	we find that we can
Deleted:	reliably and
Deleted:	Α
Deleted:	В
Deleted:	a result of
Deleted:	tilized
Deleted:	(Box C)
Deleted:	much
Deleted:	dimension
Deleted:	C
Deleted:	We find that o
Deleted:	made by Blob3D
Fig. 4) we	When initially measuring grains (as described in Sect 4.2; measured both the 'maximum width' and the 'minimum the length We utilized a verify of grain existence in

width' and the length. We utilized a variety of grain orientations in addition to the method described in Sect. 4. 2 in order to best estimate the minimum width (e.g., standing the grain upright on the caxis or holding it with tweezers). We then compared these values to the equivalent dimensions output by Blob3D (as described in Sect. 4.3). As shown in Figure E1a, grain length can be reliably measured in 2D and matches quite well to the 3D length. Figure E1b shows that the maximum width can also be reliably and accurately measured. ...These results are discussed more in Appendix E. These results are discussed more in Appendix E. Cooperdock et al. (2019) only measured the length and maximum width but our results are broadly similar with 3D/2D values and absolute differences of 0.98 and 4% for length and 1.03 and 16% for maximum width.

Deleted: These results are discussed more in Appendix E. (2019) only measured the length and maximum width but our results are broadly similar with 3D/2D values and absolute differences of 0.98 and 4% for length and 1.03 and 16% for maximum width.

Formatted: Normal

Deleted: 7

540 <u>5.1 Comparison of grain dimensions from 2D microscopy and 3D CT data</u>

5 Results: Corrections and Uncertainties

calculated using Pearson's L

For apatite dimension data, the 3D versus 2D scatter plots illustrate that the 2D values accurately measure the length (Box A) and the maximum width (Box B), with average 3D/2D values of 1.0 and 0.99 and average differences of 5%, respectively (Fig 6a-b). Outliers are due to oddly shaped or fragmented grains, which can be inaccurately measured by the procedure used by Blob3D (Cooperdock et al., 2019). However, we find that the third dimension, the minimum width, is more difficult to

- 645 et al., 2019). However, we find that the third dimension, the minimum width, is more difficult to measure accurately via microscopy (Fig. 6c). Our 2D minimum width measurements consistently underestimate the 3D Box C measurements with a large amount of scatter; the average 3D/2D value is 1.09 with an average absolute difference of 13%. This high uncertainty on minimum width is attributable to the practical challenges associated with photographing an apatite crystal in the proper
- 650 <u>orientation for minimum width measurement (Sect. 4.2).</u>

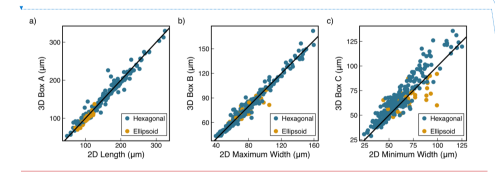


 Figure 6. Scatter plots of 3D vs. 2D data (N = 264) for grain dimension measurements. a) 3D Box A vs. 2D length measurement, b)

 555
 3D Box B vs. 2D maximum width measurement, and c) 3D Box C vs. 2D minimum width measurement. Note that the 2D minimum width data have greater scatter and systematically underestimate the corresponding 3D Box C measurement.

5.2 Corrections for systematic error

- 595 The 3D versus 2D scatter plots for V, FT, and RFT (Figure 7a, c) using the maximum width for both width values for 2D calculations all show data that systematically plot below the 1:1 line (bold black line), indicating that for all parameters the 2D values overestimate the "true" 3D values. The 2D data can be corrected for their systematic overestimation of the 3D data by multiplying the 2D data by the slope of the 3D vs. 2D data, so that the 2D data are centered around the 1:1 line, thereby "correcting" them. As noted in Sect. 4.4, regressions of the 3D vs. 2D data are separated by geometry because the
- regressions of hexagonal and ellipsoidal grains yield statistically distinguishable slopes.

The corrections for systematic error for apatite V, F_T, and R_{FT} are summarized in Table 2. For all parameters, the magnitude of the correction is smaller for hexagonal grains than for ellipsoid grains. For

- example, for V, the slope of the regression line is 0.83 for hexagonal grains and 0.74 for ellipsoid grains. This means that the volumes estimated by microscopy measurements typically overestimate the true grain volume by 17% for hexagonal grains, and by 26% for ellipsoid grains. For ²³⁸F_T, the corrections are substantially smaller, with values of 0.97 and 0.92 for hexagonal and ellipsoid grains. For R_{FT}, the corrections are 0.93 and 0.85 for hexagonal and ellipsoid grains.
- 710 Figure D1a-c, includes 3D versus 2D scatter plots for V, F_T, and R_{FT} using both the maximum and minimum width values for 2D calculations, with the associated corrections for systematic error summarized in Table D1. In this case, for hexagonal grains, all data systematically plot above the 1:1 line (bold black line), indicating that the 2D values consistently underestimate the "true" 3D values
- 715 (Figure D1a-c). The corrections for systematic error are systematically Jarger for all parameters than the corrections using only the maximum width (Table 2). For example, for hexagonal grains, V, F_T, and R_{FT} are underestimated by 27%, 8%, and 15%, respectively when using both widths, compared with an overestimation of 17%, 3%, and 7% when using only the maximum width. For ellipsoid grains, using both widths causes 2D values to overestimate the 3D values (the 2D data plot below the 1:1 line in
- 720 Figure D1a-c), however the magnitude of these corrections are the same or slightly smaller than when using only the maximum width for 2D calculations (Table 2, D1).

5.3 Uncertainties

The uncertainties for V, F_T, and R_{FT} are derived from the scatterplots of the percent difference in the residuals versus maximum width where the bold black line represents no difference between the 2D and

- 3D data (Figure 7d-f, for the analysis using the maximum width only for 2D calculations). The uncertainties are grouped by geometry for all parameters, because the residuals are derived from the regression lines, which group data in this way. The standard deviation of the percent difference in the residuals of each group is the uncertainty on each parameter, reported in Table 2 at 1s. A single uncertainty is reported for ellipsoid apatite grains for all parameters due to the relatively small number
- 730 of ellipsoid grains in the dataset (N = 36). However, for hexagonal grains, the data population (N= 201) is large enough that we explored surface roughness and grain size as potential grouping variables. We did not find a consistent, substantial relationship between surface roughness and uncertainty in the data

Deleted: 1		

Deleted: 6 Deleted: A

Deleted: C

Deleted: above

Dele	eted: A
Dele	eted: C
Forr	natted: Font: Not Bold, Font color: Auto, Subscript
Forr	natted: Font: Not Bold, Font color: Auto, Subscript
	e ted: ¶ nown in Figure E2,
Dele	eted: are now underestimated by
Dele	eted: measurements
Dele	eted: A
Dele	eted: C
Dele	eted: t
Dele	eted: magnitude of the
Dele	eted: and uncertainties is
Dele	eted: hexagonal grains when the average width is used.
Dele	eted: A
Dele	eted: C
Dele	eted: t
Dele	eted: is
sligh	ted: average width but the uncertainties are the same or tly larger when using the average width when compared to calculated using the
Dele	eted: 2
Dele	eted: in Figure 6D-F,
Dele	eted: .
Dele	eted: D
Dele	eted: F
Dele	ted: the

Deleted: which are

(Table C2). However, for grain size, the ${}^{238}F_{T}$ uncertainty for medium-sized (maximum width 50-100 μ m) hexagonal apatite is greater than for large-sized (maximum width > 100 μ m) hexagonal apatite. As described below, this pattern is sensible, so we report two uncertainties for the isotope-specific F_{T} values of hexagonal grains based on size.

For all parameters, the uncertainty for hexagonal grains is smaller than the uncertainty for ellipsoid
 grains (Table 2). For V, the uncertainty is 20% for hexagonal grains and 23% for ellipsoid grains of all sizes. For ²³⁸F_T, the uncertainties are 3% and 2% for medium and large hexagonal grains, respectively, and 5% for all ellipsoid crystals. For R_{FT}, the uncertainty is 6% for hexagonal grains and 10% for all ellipsoid grains of all sizes.

775 As anticipated, the isotope-specific F_T uncertainties are highly correlated, yielding <u>correlation</u> <u>coefficients</u> of 0.972-0.999. For this reason, below we assume fully correlated uncertainties of 1 for F_T uncertainty propagation into the corrected date.

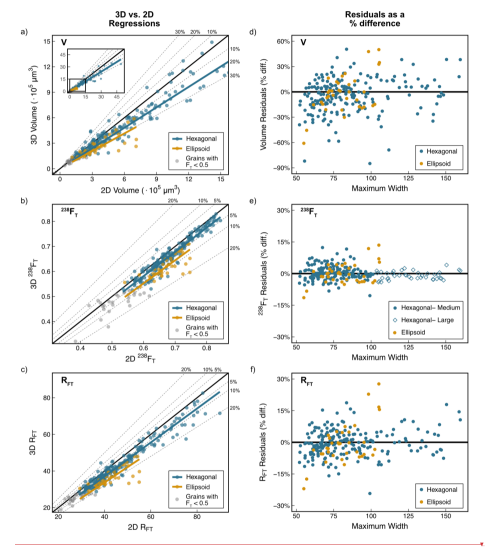
Figure D1d-f includes the relevant scatterplots for determining uncertainties on V, F_T, and R_{FT} using
 both the maximum and minimum width values for the 2D calculations, with the derived uncertainty values listed in Table D1. For hexagonal grains, the uncertainties are consistently larger for all parameters when using both widths rather than only the maximum width for 2D calculations. For ellipsoid grains, the uncertainties are the same or larger when using both widths instead of only the maximum width (Table 2, D1).

785

Deleted: values

Deleted: D Deleted: F Formatted: Font: Not Bold, Font color: Auto, Subscript Formatted: Font: Not Bold, Font color: Auto, Subscript Deleted: f Deleted: As shown in Table E1, the magnitude of the correction and ... Deleted: is Deleted: the magnitude of the correction is slightly smaller when using the average width but Deleted: slightly Deleted: the average

Deleted: when compared to those calculated using



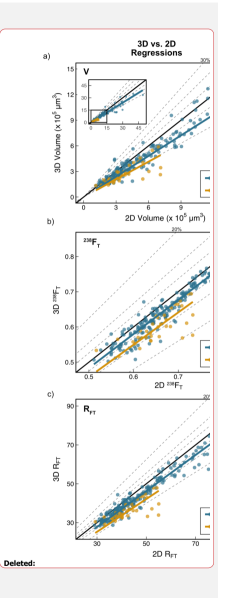




Figure 7, Plots illustrating how the corrections for systematic error and how uncertainties were determined for V, FT, and RFT. Scatter plots of 3D vs. 2D data (N=264) with regression lines and data distinguished by geometry for (a) V, (b) ²³⁸F_T, and (c) R_{FT}. 2D data in these plots were calculated using the maximum width for both width values (see Appendix A). Grains with $F_T < 0.5$ were excluded from the regressions but are included on the plots in light grey. A total of 237 apatite grains are in the regressed dataset. The bold black line is the 1:1 line and the dashed lines mark the percent difference from the 1:1 line. Note that for all

regressions, the regression line falls below the 1:1 line, indicating that the 2D-microscopy data overestimate the 3D-CT data. The 2D data can be corrected for systematic error by multiplying the 2D data by the slope of the regression. Plots of the difference of

805

2D tail can be considered and a systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by multiplying in 2D tails of methods in the systematic trib by methods in the systemati 810

width) vs. large (> 100 µm maximum width) size. The bold black line is 0% difference. Note the larger y-axis scale for V as compared with ²³⁸F_T and R_{FT}, reflecting the greater uncertainty of V. The standard deviation of the % difference in the residuals of each group is the uncertainty on the parameter.

Table 2. Corrections and uncertainties (1s) for all geometric parameters.

Volume

Geometry	Correction ^a	% Uncert. ^b (1s) for apatite grains of all sizes
Volume		
Hex.	0.83	20%
Ellip.	0.74	23%

Isotope-specific F_T values

%	Uncert.	(1s) for	%	Uncert.	(1s)
---	---------	----------	---	---------	------

		medium-sized ^c	for large-sized ^d
Geometry	Correction	apatite grains	apatite grains
²³⁸ F _T			
Hex.	0.97	3%	2%
Ellip.	0.92	5%	5%
²³⁵ F _T			
Hex.	0.96	4%	2%
Ellip.	0.91	6%	6%
²³² F _T			
Hex.	0.96	4%	2%
Ellip.	0.91	6%	6%
¹⁴⁷ F _T			
Hex.	0.99	1%	1%
Ellip.	0.97	1%	1%

 \mathbf{R}_{FT}

••••		
		% Uncert. (1s) for apatite grains of
Geometry	Correction	all sizes
R _{FT}		
Hex.	0.93	6%
Ellip.	0.85	10%

Deleted: 6	
Deleted: These p	
Deleted: e	
Deleted: 3	
Deleted: 7	
Deleted: 3D/2D	

Deleted:

^a The correction value is the slope of the 3D vs. 2D regression line for each parameter in Figures 74-74

^b The uncertainty is the scatter of the 2D data about each regression line in Fig. 7455 calculated as the 1s standard deviation of the % difference of each 2D value from the regression line (Fig. 7450).

^c "Medium-sized" apatite have maximum widths of 50-100 μm.

^d "Large-sized" apatite have maximum widths of >100 μm.

6 Discussion

225

6.1. <u>Measurements and grain characteristics that influence the accuracy and precision of 2D</u> geometric data

The goal of this study was to develop a simple method for correcting for systematic error and for assigning uncertainties to geometric parameters estimated from microscopy measurements for the full spectrum of apatite <u>grains</u> that are regularly analyzed by (U-Th)/He. Thus, the corrections for systematic error are intended to improve the accuracy of the V, F_T, and R_{FT} values derived from 2D

- 835 data. The uncertainties are aimed at appropriately representing the reproducibility or precision of these geometric parameters. Accomplishing this required determining the <u>measurements and</u> grain characteristics that most affect the accuracy and precision of the 2D data.
- Whether only the length and maximum width, or the length, maximum width, and minimum width, are used for calculating the 2D geometric parameters influences both the magnitude of the correction for systematic error and the uncertainties (Sect. 5). We recommend using the maximum width only rather than both the maximum and minimum widths for 2D geometric parameter calculations for several reasons. First, the length and maximum width are the two most accurately and reproducibly measured dimensions; it is difficult to efficiently and reliably measure the apatite's minimum width (Fig. 6c).
- 845 Second, no excess correction or uncertainty is introduced by measuring and using only the maximum width rather than both widths. In fact, for most apatite grains, the corrections and uncertainties are higher when using both width measurements (see Sect. 5), due to the inaccuracy and scatter of the 2D minimum width data. Third, it is common practice in many labs to acquire only one grain image and measure only an apatite's maximum width (Cooperdock et al., 2019), such that this set of corrections
- 850 and uncertainties may be more widely useful. This may be especially true for retroactive application to published data. Finally, time is saved by not acquiring a second set of measurements at no detriment to the data quality. The rest of our discussion below is focused entirely on these outcomes that use only the length and maximum width in the 2D calculations.
- We find that the <u>first-order grain morphology is</u> the <u>grain characteristic that most strongly influences the</u> magnitude of the systematic error <u>on 2D geometric data</u>. For example, whether apatite grains are hexagonal or sub-hexagonal (A or B on the GEM) <u>versus</u> ellipsoid (C on the GEM), dictates the choice of a hexagonal or ellipsoid idealized geometry. This in turn determines the magnitude of the correction

Deleted: 6	
Deleted: A	
Deleted: 6C	
Deleted: 6	
Deleted: A	
Deleted: C	
Deleted: 6	
Deleted: D	
Deleted: F	
Deleted: G	

	Deleted: crystals
Å	Deleted: ,
М	Deleted: It is important to note that part of the reason for this systematic overestimation is because we
7 χ	Deleted: ed
/	Deleted: using the maximum width only
Ζ,	Deleted: suggest
ζ,	Deleted: two main
ζ,	Deleted: : 1): 1)
Δ	Deleted: liably
λ	Deleted: there is no
(Deleted: e
	Deleted: way to
Y	Deleted: C
-(Deleted: only
$ \rightarrow $	Deleted: , 2)
Λ	Deleted: and 2) because no excess correction or uncertainty is introduced by measuring the maximum width only.
	Deleted: However, we recognize that some labs may want to continue measuring and using the minimum width to calculate V, F _T , and R _{FT} , so we present a set of corrections and uncertainties that can be used in this case (Fig. E2, Table E1).
1	Deleted: We find that
1	Deleted:
(Deleted: is influenced primarily by the first-order grain morphology
(Deleted: vs

Deleted:

required to make the geometric parameters calculated from the microscopy data accurate (e.g., for $^{238}F_T$ a 0.97 correction for hexagonal grains vs. a 0.92 correction for ellipsoid grains).

- 900 Our results show that the uncertainty in the 2D geometric parameters is controlled primarily by the grain geometry, and for F_T, secondarily by the grain size. Uncertainties on hexagonal grains are consistently smaller than those for ellipsoid grains (Table 2). For example, for V, uncertainties are 20% and 23% for hexagonal and ellipsoid grains, respectively. For R_{FT}, the uncertainties on hexagonal grains (6%) are again smaller than for ellipsoid grains (10%). For ²³⁸F_T, grain size exerts additional influence
- 905 on the uncertainty of hexagonal grains, with uncertainties of 3% and 2% for grains with maximum widths of 50-100 μm and > 100 μm, respectively, compared with an uncertainty of 5% for ellipsoid grains of all sizes. The influence of size on the F_T uncertainty is not surprising given that the effect of the uncertainty in grain measurements is proportionately larger for smaller grains. This pattern is consistent with early work that estimated F_T uncertainty increased with decreasing grain size (Ehlers and Farley, 2003).

6.2 Overestimation of the 3D geometric parameter values by the 2D microscopy method

6.2.1. Overview

In this study, all values calculated from the 2D microscopy measurements overestimate the real 3D values (when using length and maximum width for 2D geometric parameter calculations, as discussed

- 915 in the previous section). This overestimation is true regardless of grain size, morphology, and other grain characteristics. Compared with past work (Herman et al., 2007; Evans et al., 2008; Glotzbach et al., 2019; Cooperdock et al., 2019), in this study we analyzed more apatite (237 compared with 4-109) and at a higher CT resolution (0.64 µm compared with 1.2 -6.3 µm). We also deliberately included the full variety of grain morphologies across a range of grain sizes from samples of variable age and
- 920 lithology, so we have confidence that the results are applicable to the spectrum of routinely analyzed apatite.

As explained in Sect. 4.4, the corrections and uncertainties discussed above and reported in Table 2 are calculated from the regressions and are computed in this way because the objective of our work is to

- 925 systematically correct real 2D data and routinely apply the associated uncertainty to them. However, previous studies, which did not have these same goals in mind, reported the average 3D/2D value and its 1s uncertainty as a measure of systematic error, and reported the average absolute percent difference between the 2D and 3D data and its 1s uncertainty as a measure of the uncertainty of each parameter. To directly compare our results to this past work, in Table 3 we also report our results in this way. This
- 930 table directly follows the structure of Table 3 in Cooperdock et al. (2019). In our Table 3, we report values for our entire dataset, as well as subdivided by hexagonal and ellipsoid geometry. However, for simplicity, we use only the average values for our whole dataset in the discussion below.

We place our results in the context of those of Cooperdock et al. (2019) and Glotzbach et al. (2019) 935 because these two studies directly compared 2D microscopy with 3D CT values for a moderate to large

Deleted: se	
Deleted: (6%)	
Deleted: (± 2.7 μm)	
Deleted: (± 2.7 μm) Deleted: de	

Deleted: ¶ ¶

Deleted: crystals

	suite of apatite crystals. Cooperdock et al, (2019) characterized 109 hexagonal to sub-hexagonal apatite	
45	grains (A1 and B1 in our GEM) by CT (5 µm resolution) and calculated 2D parameters using the length	
	and maximum width only. Glotzbach et al. (2019) analyzed 24 apatite crystals (1.2 µm CT resolution)	
	with a wider range of characteristics (rounded through euhedral morphologies) and calculated 2D	
	parameters using measurements of the length, maximum width, and minimum width. Although Evans et	
	al. (2008) also carried out a study of this kind and was the first to do this type of comparison, that work	

950 included only four apatite crystals (3.8 μm CT resolution). Herman et al. (2007) used CT to derive geometric parameter data for 11 detrital apatite grains (6.3 μm CT resolution) but did not compare the results with 2D microcopy estimates for the same grains.

955 Table 3. 2D microscopy and 3D CT data comparison for this and previous studies^a

	leted:	
U De	ieteu:	

Deleted: , while

Deleted:

This Study: 227 an	atite grain	CT recolu	tion: 0.64	Im	
This Study: 237 apatite grains; CT resolution: 0.64 µm					
	avg.		abs. avg. %		
	3D/2D ^b	1s	diff. ^c	1s	
All data: 237 grain	s				
Volume	0.85	0.17	19	13	
²³⁸ F _T	0.96	0.04	4	4	
R _{FT}	0.92	0.07	8	6	
Length/Box A	1	0.07	5	6	
W _{max} /Box B	0.99	0.06	5	4	
W _{min} /Box C	1.09	0.14	13	10	
Hexagonal apatite	: 201 grain	s			
Volume	0.87	0.17	18	12	
²³⁸ F _T	0.97	0.03	4	3	
R _{FT}	0.93	0.06	7	5	
Length/Box A	1.01	0.07	5	6	
W _{max} /Box B	1	0.06	4	4	
W _{min} /Box C	1.11	0.12	13	10	
Ellipsoid apatite: 3	6 grains				
Volume	0.75	0.17	26	15	
²³⁸ F _T	0.92	0.05	8	4	
R _{FT}	0.86	0.08	15	8	
Length/Box A	0.98	0.06	6	3	
W _{max} /Box B	0.97	0.07	6	5	
W _{min} /Box C	0.97	0.16	12	11	

Previous Studies

	avg.		abs. avg. %	
	3D/2D	1s	diff.	1s
Cooperdock et al.	2019): 108	apatite g	grains; CT reso	olution: 4-5 µm
Volume	0.82	0.22	23	16
$^{238}F_{T}$	1.01	0.02	2	2
R _{FT}	1.02	0.07	5	5
Length/Box A	0.98	0.1	4	6
W _{max} /Box B	1.03	0.07	16	8
W _{min} /Box C	N/A ^d	N/A	N/A	N/A
Glotzbach et al. (2	019): 24 aj	oatite gra	ins; CT resolu	tion: 1.2 μm
Volume	1.04	0.2	15	13
²³⁸ F _T	0.99	0.02	2	2
R _{sv} ^e	0.93	0.06	8	5

24

This Study: 237 apatite grains; CT resolution: 0			
	avg.		abs. avg. 9
	3D/2D ^b	1s	diff. ^c
All data: 237 g	grains		
Volume	0.85	0.17	19
$^{238}F_{T}$	0.96	0.04	4
R _{FT}	0.92	0.07	8
Hexagonal apa	atite: 201 gr	ains	
Volume	0.87	0.17	18
$^{238}F_{T}$	0.97	0.03	4
R _{FT}	0.93	0.06	7
Ellipsoid apati	te: 36 grain	s	
Volume	0.75	0.17	26
$^{238}F_{T}$	0.92	0.05	8
R _{FT}	0.86	0.08	15
Previous Studi	es		
	avg.		abs.avg. 9
	3D/2D	1s	diff.
Cooperdock et	, ,		
Volume	0.82	0.22	23
$^{238}F_{T}$	1.01	0.02	2
R _{FT}	1.02	0.07	5
Glotzbach et a	ıl. (2019): 24	4 apatite	grains; CT r
Volume	1.04	0.2	15
$^{238}F_{T}$	0.99	0.02	2
R _{sv} ^d	0.93	0.06	8

Deleted:

^a Directly follows the structure of Table 3 reported in Cooperdock et al. (2019) to facilitate comparison with previous studies. ^b avg. 3D/2D is the average of all 3D/2D values in each study ^c abs. avg. % diff. Is the average absolute percent difference between the 2D and 3D data. We used the formula $\left(\frac{|2D-3D|}{2D}\right) \times 100$ to calculate the percent difference for consistency with Cooperdock et al. (2019). 965 ^d N/A is not available Glotzbach et al. (2019) reports R_{SV} rather than R_{FT}. Deleted: d 6.2.2. Volume Of the geometric parameters evaluated in this study, V shows the greatest overestimate of 2D relative to 3D values (2D value corrections of 0.83 and 0.74 depending on geometry) and the greatest data scatter 970 (20% and 23%) (based on the data regressions, Table 2). If we instead report our outcomes as the average 3D/2D value and the average absolute % difference, we obtain values of 0.85 and 19% for all grains (Table 3). This result is generally consistent with those of previous work. Cooperdock et al., Deleted: n (2019) also found a V overestimate with an average 3D/2D value of 0.82 and an average difference of 23%. Glotzbach et al. (2019) found no systematic over- or underestimate in volume (avg 3D/2D =975 1.04), partly attributable to their use of all three dimensions (3D-He method) in their 2D parameter calculations, and they report a similar magnitude of variation (15%). Deleted: but 6.2.3. F_T For F_T, our 2D values overestimate the 3D values. The isotope-specific ²³⁸F_T has a 2D correction value of 0.97 for hexagonal grains and 0.92 for ellipsoid grains, with uncertainties of 2-5% depending on 980 geometry and size (based on the regressions, Table 2). The corrections and uncertainties for the other isotope-specific F_T values vary from 0.99 to 0.91 and 1-6%, respectively, (again depending on grain geometry and size, Table 2), but we focus on the ²³⁸F_T value here because it dominates the ⁴He Deleted: again production. Our average 3D/2D value for ²³⁸F_T is 0.96, with an average difference of 4% (Table 3). This outcome is similar to that of Glotzbach et al. (2019) (avg. 3D/2D = 0.99; avg. abs. diff. = 2%). In 985 contrast, Cooperdock et al. (2019), report 2D values that slightly underestimate the 3D F_T values (average 3D/2D = 1.01), but with a comparable magnitude of scatter (2%). This may be due, in part, to their grain selection, which focused mainly on high quality, hexagonal apatite grains. 6.2.4. R_{FT} For R_{FT}, we found that 2D measurements were systematically larger than 3D measurements (2D 990 correction values of 0.93 and 0.85), with uncertainties of 6-10% depending on geometry (based on the regressions, Table 2). Our average 3D/2D value for R_{FT} is 0.92, with an average difference of 8% (Table 3). Glotzbach et al. (2019) reports R_{SV} (the equivalent sphere with the same surface area to volume ratio as the grain) rather than R_{FT}, but these values typically have negligible difference. Their Deleted: little to dataset yields R_{SV} outcomes nearly identical to our R_{FT} results (avg. 3D/2D = 0.93; avg. abs. diff. = 995 8%). In contrast, Cooperdock et al. (2019) found an average 3D/2D value of 1.02 and an average

difference of 5% (Table 3). Their underestimation of R_{FT} by 2D measurements is expected given the systematic underestimation they report for F_T .

6.3 Implications: How much do the corrections and geometric uncertainties matter?

6.3.1 Overview

- To determine how much the corrections and geometric uncertainties (Table 2) affect the values and uncertainties on real (U-Th)/He dates and other key parameters, we apply our corrections and uncertainties to the V, F_T, and R_{FT} values of a subset of representative apatite grains from three samples (N = 24) that were used in this study and that were previously dated in the CU TRaIL (Table F1). This
- apatite suite includes both hexagonal and ellipsoid grains with a range of sizes. We then use the order of the corrected V and isotope-specific F_T values to calculate the parameters derived from them—mass, eU,
- and the corrected (U-Th)/He date—and propagate the geometric uncertainties on V and F_T into the uncertainties of the derived values. Below, we then compare the "new" values and uncertainties on all parameters with their "original" uncorrected counterparts (Sect. 6.3.2-6.3.5), generate corrected apatite (U-Th)/He (AHe) date vs. eU plots using both the new and original values (Fig. 8), and consider the
- 015 broader implications of these outcomes for interpretation of AHe data (Sect. 6.3.6).

Table 4 summarizes the average new/original values for this example dataset, as well as how much the uncertainty on each parameter increases owing to the inclusion of geometric uncertainties (which have traditionally been excluded from the uncertainties reported on these parameters). For uncertainty

- p20 propagation into the corrected (U-Th)/He date, we use HeCalc (Martin et al., 2023) and assume fullycorrelated (r = 1) isotope-specific F_T uncertainties. In Table 4 and the discussion below all uncertainties are reported at 1s. Standard practice in the CU TRaIL over the last several years has been to report 15% 1s uncertainties on eU based on estimates by Baughman et al. (2017). However, how eU uncertainties are reported varies widely across the community and it is common for no uncertainty to be reported on
- 025 eU data, therefore for comparative purposes, no uncertainty is shown on eU_{orig} in Fig. 84-c, and none is reported in Table F1.

6.3.2 Mass and eU

To calculate eU, absolute quantities of U, Th, and Sm must be converted to concentrations using the apatite grain mass, which is computed from V assuming an apatite density (here we use 3.20 g/cm³).
O30 Absolute amounts of parent isotope carry an analytical uncertainty, but conventionally the grain mass reported by labs has had no uncertainty attached to it because the geometric uncertainty on V (and therefore on mass) was not well constrained. By applying a correction factor to V based on grain geometry (0.83 or 0.74) and calculating mass using the corrected V, the mass_{new} decreases by the same correction factor as volume. The mass then inherits the same percent uncertainty as volume (20 or 23%,

035 1s, depending on geometry).

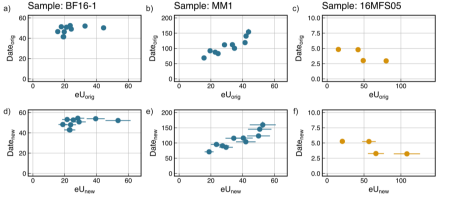
For eU, the smaller mass_{new} values (relative to mass_{orig}) are translated into larger eU_{new} values (relative to eU_{orig}). In our example dataset (Table 4), the average eU_{new}/eU_{orig} is 1.2 for hexagonal grains and 1.4



ples his	(Deleted: D
U,		
all atite		
e	(Deleted: 7
the ave		
ly-	(Deleted: (Martin et al., in review)
nties 15%		
ies l on		
e is	(Deleted: 7
	$\langle \langle \langle \langle \langle \rangle \rangle \rangle \rangle$	Deleted: A
	\searrow	Deleted: C
	X	Deleted: D

for ellipsoid grains. We propagated the analytical uncertainties on the parent isotopes only, as well as the <u>parent isotope</u> and geometric uncertainties, into the eU_{new} values. Propagating <u>parent isotope</u> uncertainties only yields average eU uncertainty values of 3% for hexagonal and ellipsoid grains in this dataset (with a range from 1 to 6%). Including both analytical and geometric uncertainties yields average uncertainties of 15% and 16% for hexagonal and ellipsoid grains (varying from 14-17%).





Deleted: analytical Deleted: analytical Deleted:

Figure & Date-eU plots for three samples previously dated in the CU TRaIL showing the effects of corrections and uncertainty estimates on typical AHe data. (apc) are date_{arig} vs. eU_{orig} plots, while (daf) are date_{aren} vs. eU_{new} plots. When uncertainty bars are not visible they are on the order of the symbol size, except for the top row where no eU uncertainty is plotted. <u>An idealized</u> <u>hexagonal geometry was used for 2D geometric parameter calculations for the igneous apatite in samples BF16-1 and MM1 (blue circles), while an idealized ellipsoid geometry was used for the detrital apatite in sample 16MFS05 (xellow circles).</u>

6.3.3 Combined F_T values

The combined F_T values are calculated using both the isotope-specific F_T values and the amount of the parent isotopes, because the proportion of the parent isotopes dictates the proportion of the ⁴He atoms that travel different mean stopping distances. The combined F_T values are not used for any additional calculations except R_{FT}, but are typically reported in data tables (e.g., Flowers et al., 2022a). For our example dataset, we apply the correction factors in Table 2 based on grain geometry and size to the isotope-specific F_T values, and then use these corrected values to calculate the combined F_{T,new} value.
 65 F_{T,new} is always smaller than F_{T,orig} (F_{T,new} / F_{T,orig} = 0.97 and 0.92 for hexagonal and ellipsoid grains;

Table 4).

 F_T values have not typically been reported with an uncertainty, because until now the geometric uncertainty on F_T has been poorly quantified. For comparative purposes, we propagated uncertainties

b70 into the combined F_T value using the <u>parent isotope</u> uncertainties only, as well as using both <u>parent</u> <u>isotope</u> and geometric uncertainties. For the example dataset, inclusion of analytical uncertainties only yields average uncertainties on the combined F_T of 1% (1s, with a range from 0-3%) for both grain

(Deleted: 7
(Deleted:)
1	Deleted: (
Y	Deleted:)
Y	Deleted: (
)(Deleted: orange

(Deleted: W
(Deleted: analytical
(Deleted: analytical

geometries. The propagation of both <u>parent isotope</u> and geometric uncertainties generates average values of 2% for hexagonal grains (varying from 1-3%) and 4% for all ellipsoid grains (Table 4).
 Variability in the uncertainties for the combined F_T is due to variability in the total <u>parent isotope</u> uncertainty.

p90 Table 4. The average percent difference between the original and new values for example dataset of Table **E**1,

	_	% Analy	ytical uncertaint	y only ^c , 1s	% Analytica	l + Geometric U	ncertainty ^d , 1s	
Parameter and Geometry ^a	Avg. New/Orig. ^b	Avg.	Min (%)	Max (%)	Avg.	Min (%)	Max (%)	Avg. % uncert. increase ^e , 1s
Mass								
Hex.	0.83	NA	NA	NA	20%	20%	20%	NA
Ellip.	0.74	NA	NA	NA	23%	23%	23%	NA
eU								
Hex.	1.20	3%	1%	6%	15%	14%	16%	12%
Ellip.	1.40	3%	2%	3%	16%	16%	17%	13%
Combined F_{T}								
Hex.	0.97	1%	0%	3%	2%	1%	3%	1%
Ellip.	0.92	1%	1%	1%	4%	4%	4%	3%
Corr. Date								
Hex.	1.04	2%	1%	6%	3%	2%	7%	1%
Ellip.	1.09	4%	2%	6%	7%	6%	8%	3%
R _{FT}								
Hex.	0.93	NA	NA	NA	6%	6%	6%	NA
Ellip.	0.85	NA	NA	NA	10%	10%	10%	NA

NA indicates "Not Applicable", for example, mass doesn't have any analytical uncertainty on the parent isotopes.

^a There are N = 20 hexagonal and N = 4 ellipsoid grains.

^b The average of the new parameter (calculated using the new values) divided by the average of the original values (calculated using the original values) for the example data in Table [1]. Values under 1 indicate that the original value is larger than the new. Values over 1 indicate that the original value is smaller than the new.

^c The average of the percent analytical (i.e., parent isotope) uncertainties only for the example data in Table **E**1.

^d The average of the percent analytical + geometric uncertainties for the example data in Table **L**1.

^e The average percent increase is the difference between the analytical only and analytical + geometric uncertainties.

100 6.3.4 Corrected (U-Th)/He dates

The most rigorous means of calculating F_T -corrected (U-Th)/He dates is by incorporating the isotopespecific F_T corrections into the age equation and calculating the corrected date iteratively (Ketcham et al., 2011). For our example dataset, we used the corrected isotope-specific F_T values (as described above) to calculate the F_T -corrected AHe date_{new}. For the AHe dates, the smaller $F_{T,new}$ values (relative

105 to $F_{T,orig}$) are translated into larger corrections for alpha-ejection. Thus, the date_{new} values are always older than the date_{orig} values (avg. date_{new} / date_{orig} = 1.04 and 1.09 for hexagonal and ellipsoid grains).

28

Deleted: analytical

Deleted: analytical

Deleted: D				
	Parameter and	-	Avg. analyt.	
	Geometry ^a	Avg. New/Orig. ^b	uncert. only ^c	
	Mass	New/Ong.	(%), 1s	N
	Hex.	0.83	NA	
	Ellip.	0.74	NA	
	eU			
	Hex.	1.20	3%	
	Ellip.	1.40	3%	
	Combined F_T			
	Hex.	0.97	1%	
	Ellip.	0.92	1%	
	Corr. Date Hex.	1.04	2%	
	Ellip.	1.04	4%	
	R _{FT}			
	Hex.	0.93	NA	
	Ellip.	0.85	NA	
Deleted:				

Deleted: D
Deleted: D
Deleted: D
Deleted: D

Deleted: D

- We calculated the uncertainty on the corrected (U-Th)/He dates in two ways for comparative purposes: first by propagating the analytical uncertainties on the parent and daughter only, and next by additionally including the geometric uncertainties on the isotope-specific F_{T,new} values and assuming fully-correlated F_{T,new} uncertainties (Table 4). For this dataset, we find that propagating only analytical uncertainties yields average uncertainties of 2% and 4% for hexagonal and ellipsoid grains (varying
- 120 from 1-6% and 2-6%, respectively). Including both analytical and geometric uncertainties yields average uncertainties of 3% and 7% for the two geometries (with 2-7% and 6-8% variability). The difference in the uncertainty on the date varies so widely because it is dependent on a variety of grain-specific factors—the absolute amounts of U, Th, Sm, and He, as well as grain geometry and size.

6.3.5 R_{FT}

125 We applied the correction factors based on grain geometry in Table 2 to R_{FT} values from the example dataset. The R_{FT,new} values are always smaller than R_{FT,orig} values (R_{FT,new} / R_{FT,orig} = 0.93 and 0.85 for hexagonal and ellipsoid grains) (Table 4). The uncertainty on R_{FT} is 6% (1s) for hexagonal grains and 10% (1s) for ellipsoid grains. This parameter is not used in the calculation of (U-Th)/He dates, but the uncertainty should be used during thermal history modeling when possible.

130 6.3.6 Summary

This exercise in which we both 1) correct real AHe data for systematic error associated with the 2D microscopy approach for determining geometric parameters, and 2) propagate geometric uncertainties into the uncertainties on eU and corrected AHe dates reveals a substantial influence of both on some aspects of the results. The most striking outcome is the impact on eU. For example, the U_{new} values of

- 135 the example dataset increase by 20-40%, resulting in a noticeable shift of data to the right on the dateeU plots (compare Fig. & c, with Fig. & d-1). Moreover, the eU uncertainties when both analytical and geometric uncertainties are included are as much as 17% at 1s, indicating the importance of appropriately reporting and representing eU uncertainties. The influence of systematic error and uncertainties are less substantial for the corrected AHe date than for eU, but are still important. For
- 140 ellipsoid grains, the AHe date_{new} values are as much as 9% older than the date_{orig} values, with typical uncertainties that increase by as much as 3% when geometric uncertainties are propagated in addition to analytical uncertainties. For hexagonal grains, the corrections and uncertainties are less than for ellipsoid grains, but non-negligible. Including the geometric uncertainty on the corrected AHe dates may help account for overdispersion in some (U-Th)/He datasets. Properly correcting for systematic
- 145 error and propagating uncertainties associated with the geometric parameters is an important step for rigorously presenting and interpreting apatite (U-Th)/He data.

6.4 The Geometric Correction Method: A practical workflow

The Geometric Correction Method described here and shown in Fig. 9 can be easily integrated into existing (U-Th)/He dating workflows with no additional time, cost, or equipment. This method is most appropriate for grain characteristics like those in this calibration study, with 2D microscopy F_T values > 0.5, length/maximum width ratios of 0.8-3.6 and maximum width/minimum width ratios of 1-1.7, This

29

Del	eted:	3

Deleted: 7	
Deleted: A	
Deleted: C	
Deleted: 7	
Deleted: D	
Deleted: F	
Deleted:	

Deleted: We contend that p

Λ	Deleted: 8
λ	Deleted: assumes that
λ	Deleted: are
/	Deleted: and axial ratios < 1.7
(Deleted: It

- method also assumes that grain measurements are made in the same manner as this study (Fig. 4) and that 2D V, F_{T} , and R_{FT} values are calculated using the equations of Ketcham et al. (2011) and Cooperdock et al. (2019). All equations required for the calculations below are in Appendix A. The corrections for systematic error and the uncertainties reported here are only those associated with grain
- 170 geometry. For F_T , additional inaccuracy and uncertainty may be introduced by parent isotope zonation (e.g., Farley et al., 1996), grain abrasion (e.g., Rahl et al., 2003), and grain breakage (e.g., He and Reiners, 2022), which have potential to be accounted for separately. For mass and the derived eU concentration, additional uncertainty may be associated with the assumed mineral density.
- 175 Step 1. Select grain geometry and GEM category. Choose apatite grains for analysis. Decide whether the grain is hexagonal or ellipsoid, which is all that is strictly required to correct the 2D values and assign uncertainty. However, we strongly encourage assigning a GEM category (Fig. 3) and making other descriptive notes, which can be helpful for data interpretation.
- 180 Step 2. Measure the grain. Measure the grain using the procedure outlined in Sect. 4.2 and Figure 4.
 - Measure the grain length parallel to the c-axis. Only a single length is required; however, if the grain has an extremely angled or uneven end then two lengths may be measured and their average reported to better capture the average length.
 - Measure the apatite grain's maximum width, which is perpendicular to the grain length,
- 185

Step 3. Calculate the 2D values. Calculate 2D microscopy V and isotope-specific F_T values using the hexagonal or ellipsoid equations of Ketcham et al. (2011) depending on grain geometry. Calculate R_{FT} using the equations of Cooperdock et al. (2019). Note that parent isotope data must first be acquired for the F_T and R_{FT} values to be computed.

190

Step 4. Correct the 2D values. Multiply the 2D microscopy V, isotope-specific F_T, and R_{FT} values by the correction factor according to the grain geometry to produce the Vnew, FT.new, and RFT.new values (Table 2). Typically, combined F_T values are reported by labs, but the isotope-specific F_T values are required for the most accurate and rigorous calculation of corrected (U-Th)/He dates (Ketcham et al., 195 2011)

Step 5. Assign uncertainty. Attach the uncertainty value to each parameter according to the grain geometry (for V_{new}, F_{T, new}, R_{FT, new}) and maximum width (for F_{T, new}) (Table 2).

Step 6. Calculate derived parameters and propagate uncertainties. 200

- Calculate mass and eU using the V_{new} values. Uncertainty on V should be propagated into the uncertainty on these derived parameters.
- Calculate corrected (U-Th)/He dates using the isotope-specific F_{T.new} values. Uncertainty on F_T should be propagated into the final uncertainty on the corrected He date. This uncertainty
- 205 propagation can be easily accomplished, for example, by using the open access Python program HeCalc for (U-Th)/He data reduction (Martin et al., 2023),

(_ ·		
De	eted:	crystal

Mo wid	ved down [2]: <#>Measure the apatite grain's maximum th.¶
Del	eted: <#>,
Del	eted: <#>however
Move	d (insertion) [2]

Deleted:

Deleted: (Martin et al., in review2023)

Consider the following example: an apatite grain selected for analysis has a maximum width of 98 µm and a GEM value of B1. The ²³⁸F_{T,orig} of this grain is 0.67 (see Appendix A and the footnotes of Table E1 for the details of this calculation). The analyst uses Table 2 to select the correction for hexagonal grains (0.97) and performs the following calculation:

220 $F_{T, new} = F_{T, orig} \times correction = 0.67 \times 0.97 = 0.65$

The analyst then selects the proper uncertainty from Table 2 based on grain geometry and maximum width. This hexagonal grain is considered medium-sized because it is 98 μ m wide, so it has a geometric, uncertainty of 3%. The final ²³⁸F_{T, new} = 0.65 ± 3%, if the analytical uncertainty on the absolute amount of ²³⁸U is not also propagated into the ²³⁸F_T values. This procedure is repeated for each isotope-specific

 $\overline{F_{T, \text{ orig}}}$. The isotope-specific $\overline{F_{T, new}}$ values are used in the calculation of the corrected date and both the uncertainty on each isotope-specific $\overline{F_T}$ and the analytical uncertainty <u>on the parent and daughter</u> isotopes is propagated into the uncertainty on the corrected (U-Th)/He date.

7 Conclusions

225

- 230 Uncertainties on the geometric parameters and the data derived from them V, F_T, R_{FT}, eU, and corrected (U-Th)/He dates have not traditionally been included in the reported uncertainties on (U-Th)/He datasets. Nor have such data been corrected for systematic error that might arise from the 2D microscopy approach for determining these values. Although both uncertainties and corrections are important for accurate interpretation of (U-Th)/He datasets, the lack of well-quantified values that can
- 235 easily be determined and applied to routinely generated data has hindered progress in this area.

In this paper we present the only no-cost, easy-to-implement, and backwards-compatible solution to this problem. The Geometric Correction Method is a simple and effective set of corrections and uncertainties derived for V, F_T, and R_{FT} values that can be easily incorporated into existing workflows

- (Fig. 9). This approach corrects these parameters for systematic overestimation and provides an uncertainty that can be propagated into the uncertainty on derived parameters (eU, corrected date). It also can be easily applied to previously published data. These corrections and uncertainties are most appropriate for apatite grains like those in this calibration study, with F_T >0.5, length/maximum width ratios of 0.8-3.6 and maximum width/minimum width ratios of 1-1.7, with grain measurements and propriate or englated as in this user.
- 245 parameter calculations performed as in this work.
- We also present the Grain Evaluation Matrix (GEM), which is a simple, clear, and consistent <u>tool</u> to systematically characterize apatite grain quality (Fig. 3). Although use of the GEM is not required to apply the Geometric Correction Method, assigning GEM values during grain selection can assist in
- 250 quickly assessing a sample's overall quality and can help identify potential causes of outlier analyses. The GEM is also an effective teaching tool for those who are new to picking apatite grains, so that the wide spectrum of possible apatite morphologies is clearly communicated.

	Deleted:	D
- N	Deleteu.	$\boldsymbol{\nu}$

Deleted: n

-(Deleted: 8
-(Deleted: applicable
-(Deleted: to
~(Deleted: crystals
~(Deleted: and axial ratio < 1.7

Deleted: method

The corrections and uncertainties in this study were derived from the regression of 2D and 3D measurements of 237 apatite grains displaying a wide variety of morphologies commonly dated for (U-Th)/He thermochronology. The derived corrections and uncertainties were then applied to a set of real data analyzed in the CU TRaIL to determine their impact. The primary outcomes are:

- <u>1.</u> There is both uncertainty and systematic error associated with the microscopy approach to calculating V, F_T , and R_{FT} for apatite.
- For simplicity, consistency, and efficiency we recommend measuring and using only the apatite length and maximum width for 2D geometric parameter calculations. For most apatite grains, this method yields lower correction magnitudes and uncertainties than using the length, maximum width, and minimum width measurements because of the underestimation and scatter of 2D minimum width values.
 - Using only the length and maximum width measurements, the true values of V, F_T, and R_{FT} for apatite are all overestimated by the 2D microscopy measurements.

275

285

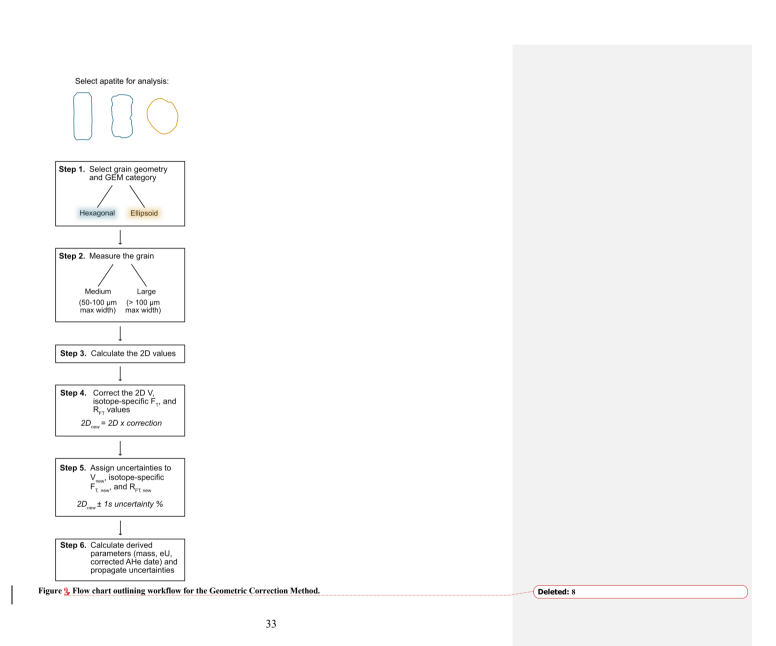
- All corrections for systematic error and all uncertainties are larger for ellipsoid grains than for hexagonal grains. For both, V has the largest magnitude of overestimation and uncertainty, followed by R_{FT}, and then F_T.
- 5. For a subset of real <u>AHe</u> data (N = 24 analyses), the correction factor for eU typically increases the eU by ~20% with associated 1s uncertainties of 15-16% when both analytical and geometric uncertainties are included. This has important implications for how data are treated during interpretation, and thermal history modeling.
 - 6. For the real dataset, the correction factor for the corrected (U-Th)/He date generally increases the date by 4-9% with associated 1s uncertainties of 3-7% if both analytical and geometric uncertainties are included. Propagating the geometric uncertainty into the corrected date may help account for overdispersion in some (U-Th)/He datasets.

The geometric corrections and geometric uncertainties are substantial enough that they should be routinely included when reporting eU and corrected (U-Th)/He dates to enhance rigorous data

290 interpretation. Ongoing work is using this same approach to quantify appropriate corrections and uncertainties for zircon geometric parameters in (U-Th)/He datasets (Baker et al., 2020).

Deleted:	
Deleted:	Therefore
Deleted:	and
Deleted:	only
Deleted:	and applying the corrections.
Deleted:	this
Deleted:	approach,
Deleted:	Т

Deleted: s
Deleted: during



Appendix A: Equations required to use the Geometric Correction Method

All equations necessary to use the corrections and uncertainties are listed below.

Equations for a hexagonal (GEM = A or B) grain from Ketcham et al. (2011), modified to reflect the use of only a maximum width (W_{max}; assuming that the minimum width = maximum width) because 310 only a maximum width is used in our preferred Geometric Correction Method, and where we use L to denote grain length instead of H.

$$\Delta V = \frac{1}{6\sqrt{3}} \left(W_{max} - \frac{\sqrt{3}}{2} W_{max} \right)^3$$
(A1)

$$V = LW_{max} \left(W_{max} - \frac{W_{max}}{2\sqrt{3}} \right) - N_p \left(\frac{\sqrt{3}}{8} W_{max}^3 - \Delta V \right)$$
(A2)

$$SA = 2L \left(W_{max} + \frac{W_{max}}{\sqrt{3}} \right) + 2W_{max} \left(W_{max} - \frac{W_{max}}{2\sqrt{3}} \right) - N_p \left(\frac{\sqrt{3}}{4} W_{max}^2 + \left(2 - \sqrt{2} \right) W_{max}^2 + \frac{\sqrt{2} - 1}{2\sqrt{3}} W_{max}^2 \right)$$
(A3)

$$R_{SV} = \frac{3V}{SA}$$
(A4)

$$F_T = 1 - \frac{3}{4} \frac{S}{R_{SV}} + \left[(0.2093 - 0.0465N_P) \left(W_{max} + \frac{W_{max}}{\sqrt{3}} \right) + \left(0.1062 + \frac{0.2234S}{S + 6(W_{max}\sqrt{3} - W_{max})} \right) \times \left(L - \frac{N_P \frac{W_{max}\sqrt{3} + W_{max}}{4}}{\sqrt{3}} \right) \right] \frac{S^2}{V}$$
(A5)

where S is the stopping distance of an alpha particle for a given parent isotope (18.81, 21.80, 22.25, and 5.93 μ m for ²³⁸U, ²³⁵U, ²³²Th, ¹⁴⁷Sm respectively), R_{SV} is the SV-equivalent spherical radius, and N_p is 325 the number of pyramidal terminations. Equation A5 is used to calculate each isotope-specific F_T value, each with different stopping distance (S).

330 Equations for an ellipsoid grain (GEM = C), from Ketcham et al. (2011):



	Deleted: abc
(Deleted: →
1	Deleted: 1
	Deleted: $S = 4\pi \left(\frac{a^{p_b p_+ b^p c^p + c^p a^p}}{3}\right)^{1/p}$ with $p = 1.6075$ $\Rightarrow \Rightarrow \Rightarrow \Rightarrow \Rightarrow \Rightarrow (A2)$
(Formatted: Font: Times New Roman, Not Italic

Formatted: Superscript

Formatted: Subscript



$$F_T = 1 - \frac{3}{4} \frac{S}{R_{SV_{+}}} + \left[\frac{1}{16} + 0.1686 \left(1 - \frac{W_{max}}{R_{SV}}\right)^2\right] \left(\frac{S}{R_{SV_{+}}}\right)^3$$

where S is the stopping distance of an alpha particle for a given parent isotope (18.81, 21.80, 22.25, and 5.93 μ m for ²³⁸U, ²³⁵U, ²³²Th, ¹⁴⁷Sm respectively) and R_{SV} is the SV-equivalent spherical radius. Equation <u>A9</u> is used to calculate each isotope-specific F_T value, each with a different stopping distance, 345

Age equation, from Ketcham et al. (2011): 350

$${}^{4}He = 8F_{T,238}{}^{238}U(e^{\lambda_{238}t} - 1) + 7F_{T,235}{}^{235}U(e^{\lambda_{235}t} - 1)$$

$$+6F_{T,232}{}^{232}Th(e^{\lambda_{232}t}-1)+F_{T,147}{}^{147}Sm(e^{\lambda_{238}t}-1)$$
(A10)

355 Equation for combined F_T and R_{FT} from Cooperdock et al. (2019):

	$\frac{S}{R} = 1.681 - 2.428F_T + 1.153F_T^2 - 0.406F_T^3$	(A <u>11</u>)
360	$A_{238} = (1.04 + 0.247 [Th/U_{]})^{-1}$	(A1 <u>2</u>)
	$A_{232} = (1 + 4.21/[Th/U])^{-1}$	(A1 <u>3)</u>
l	$\overline{F_T} = A_{238}F_{T,238} + A_{232}F_{T,232} + (1 - A_{238} - A_{232})F_{T,235}$	(A1 <u>4</u>)
365	$\overline{S} = A_{238}S_{238} + A_{232}S_{232} + (1 - A_{238} - A_{232})S_{235},$	(A1 <u>5</u>)

$$\beta 65 \quad S = A_{238}S_{238} + A_{232}S_{232} + (1 - A_{238} - A_{232})S_{235},$$

where S_{238} , S_{232} , S_{235} are the weighted mean stopping distances for each decay chain (18.81, 21.80, and $22.25 \,\mu\text{m}$, respectively, for apatite).

$$370 \quad R_{FT} = \overline{S} / \left(\frac{S}{R}\right)$$

Equation for eU from Cooperdock et al. (2019):

$$\begin{bmatrix} eU = [U] + 0.238[Th] + 0.0012[Sm] (or \ 0.0083[^{147}Sm]) \\ 375 \end{bmatrix}$$

380

/	Deleted: s
	Deleted: a
	Deleted: s
11	Deleted: s
V/i	Deleted: →
(W)	Deleted: 3
\mathbb{N}/\mathbb{N}	Deleted: W
	Deleted: and
	Formatted: Superscript
	Deleted: This e
	Deleted: nce
	Deleted: s
	Deleted: 1
	Deleted: Equations for a hexagonal (GEM = A or B) grain from Ketcham et al. (2011), modified to reflect the use of only a maximum width (W _{max}) because only a maximum width is used in our preferred Geometric Correction Method, and where we use L to denote grain length instead of H. Because only a maximum width is used in the Geometric Correction Method, ΔV is always equal to Equation (4). ¶ $\Delta V = \frac{1}{6\sqrt{3}} \left(W - \frac{\sqrt{3}}{2} W \right)^3 \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow (A4)$ ¶ $V = LW \left(W - \frac{w}{2\sqrt{3}} \right) - N_p \left(\frac{\sqrt{3}}{8} W^3 - \Delta V \right)^{+} \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow \rightarrow (A5)$ ¶ $S = 2L \left(W + \frac{w}{\sqrt{3}} \right) + 2W \left(W - \frac{w}{2\sqrt{3}} \right) - N_p \left(\frac{\sqrt{3}}{4} W^2 + (2 - \sqrt{2}) W^2 + \frac{\sqrt{2} - 1}{2\sqrt{3}} W^2 \right)^{+} \rightarrow \rightarrow (A6)$ ¶ $F_T = 1 - \frac{3}{4R_s} + \left[(0.2093 - 0.0465N_p) \left(W + \frac{w}{\sqrt{3}} \right) + \left(0.1062 + \frac{0.2234s}{3+6(W(\sqrt{3}-W))} \times \left(L - N_p - \frac{\frac{W}{2} + W}{4} \right) \right] \frac{5^2}{V} \cdot (A7)$ ¶ (
	Deleted: 1
	(Deleted: 8
	Deleted: (<i>B</i> = 1.31)
	(Deleted: 9
	Deleted: 0
	Deleted: 1
111	Deleted: 2
	Deleted: 3
	Deleted: s
	Deleted: 4
1	Deleted: 5

(A9)

(A1<u>6</u>)

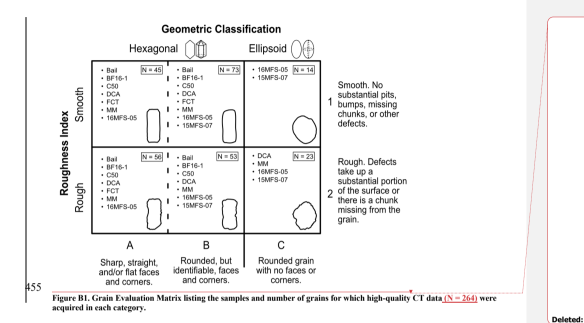
(A1<u>7</u>)

Appendix B: Additional sample information

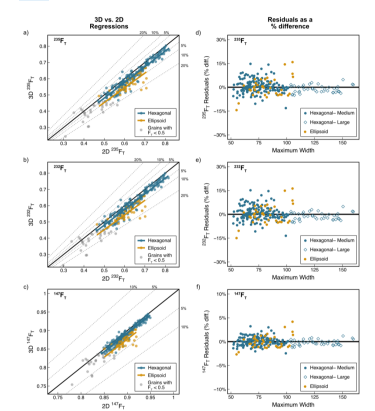
Table B1. Apatite CT scan parameters

Mount	1	2	3	4	5	6	7	8	9
Objective	20X								
Pixel Size (µm)	0.64	0.63	0.63	0.63	0.63	0.64	0.63	0.63	0.63
X-Ray Power (W)	3	3	3	3	3	3	3	3	3
X-Ray Voltage (kV) Number of	40	40	40	40	40	40	40	40	40
Projections	3201	3201	3201	3201	3201	3201	3201	3201	3201
Binning	2	2	2	2	2	2	2	2	2
Filter	Air								
Height (pixels)	1024	993	993	993	993	993	993	993	993
Width (pixels)	1024	993	993	993	993	993	993	993	993
Sample Theta (°) Detector To Sample	-180	-180	-180	-180	-180	-180	-180	-180	-180
Distance (mm)	5.01	5.17	4.95	4.97	4.99	5.42	5.07	5.08	5.02
Source To Sample Distance (mm)	-4.44	-4.51	-4.33	-4.33	-4.34	-4.75	-4.33	-4.32	-4.38
Exposure (s)	2.1	2.5	2.3	2.3	2.0	2.7	2.3	2.5	2.5
Total Scan Time (h)	3.0	3.4	3.2	3.2	2.9	3.6	3.2	3.4	3.4

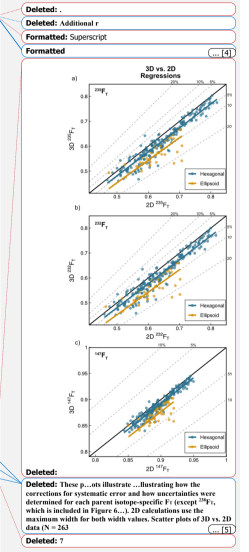
Deleted: ¶

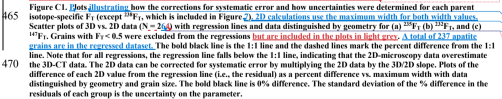


Mount	1	2	
Objective	20X	20X	
Pixel Size (µm)	0.64	0.63	
X-Ray Power (W)	3	3	
X-Ray Voltage (kV) Number of	40	40	
Projections	3201	3201	
Binning	2	2	
Filter	Air	Air	
Height (µm)	1024	993	
Width (µm)	1024	993	
Sample X (µm)	-676	82	
Sample Y (µm)	-12412	-12602	
Sample Z (µm)	87	106	
Sample Theta (°) Detector To Sample	-180	-180	
Distance (mm)	5.01	5.17	
Source To Sample Distance (mm)	-4.44	-4.51	
Exposure (s)	2.1	2.5	
Total Scan Time (h)	3.0	3.4	









	Difference in		Adjuste
Grouping & Pairs	Slopes	95% Cl ^b	P-value
Volume			
GEM: Geometric Classi	fication		
B-A	< 0.001	[-0.001, 0.001]	0.922
C-A	0.153	[0.153, 0.153]	< 0.001
C-B	0.153	[0.153, 0.153]	< 0.001
Size			
Medium-Large	0.011	[-0.007, 0.029]	0.213
GEM: Roughness			
1-2	0.010	[-0.004, 0.024]	0.157
²³⁸ FT			
GEM: Geometric			
B-A	< 0.001	[001, 0.001]	0.922
C-A	0.153	[0.153, 0.153]	< 0.001
C-B	0.153	[0.153, 0.153]	< 0.001
Size			
Medium-Large	0.011	[-0.007, 0.029]	0.231
Roughness			
1-2	0.010	[001, 0.001]	0.157
R _{FT}			
GEM: Geometric			
B-A	0	[001, 0.001]	1
C-A	0.055	[0.055, 0.055]	< 0.001
C-B	0.055	[0.055, 0.055]	< 0.001
Size		- ,,	
Medium-Large	0.004	[001, 0.001]	0.213
Roughness		, 1	
1-2	0.004	[-0.001, 0.009]	0.157

Table C1. Results of Tukey's Highly Significant Difference^a test to determine if different groups of grains have statistically different slopes.

		Difference in
	Grouping & Pairs	Slopes ^b
	Volume	
	GEM: Geometric Classifi	cation
	B-A	< 0.001
	C-A	0.153
	C-B	0.153
	Size	
	Medium-Large	0.011
	GEM: Roughness	
	1-2	0.010
	²³⁸ FT	
	GEM: Geometric	
	B-A	< 0.001
	C-A	0.153
	C-B	0.153
	Size	
	Medium-Large	0.011
	Roughness	
	1-2	0.010
	R _{FT}	
	GEM: Geometric	
	B-A	0
	C-A	0.055
	C-B	0.055
	Size	
	Medium-Large	0.004
	Roughness	
Deleted:	1-2	0.004

^a Tukey's Highly Significant Difference tests if slopes are significantly different from each other or not and takes into account the uncertainties on the slopes. Where the null hypothesis, H_0 , is $\beta_1 = \beta_2$ and the alternative hypothesis, H_1 , is $\beta_1 \neq \beta_2$.

^b/_vThe 95% confidence interval (CI) of the difference in slopes.

\$A p-value < 0.05 indicates that H₀ can be rejected, i.e., there is a significant difference between the slopes of the pair. If the p-value is > 0.05, this indicates that there is no significant difference between the means of the pair. Bolded pairs of slopes are those with p-values <0.05 and therefore are treated as separate groups.</p>

Deleted: ^b

Deleted: d

Deleted: 1

_			
Table C2. Uncertaint	v values (1s) for diff	erent groupings of	physical variables.

Geometry	Size ^a	Roughness	Ν	Uncertainty
Volume				
Hex.	Medium & Large	1&2	201	20%
Hex.	Medium	1&2	161	20%
Hex.	Medium	1	86	19%
Hex.	Medium	2	75	21%
Hex.	Large	1&2	40	23%
Hex.	Large	1	18	15%
Hex.	Large	2	22	28%
Ellip.	Medium & Large	1&2	36	23%
²³⁸ F _T				
Hex.	Medium & Large	1&2	201	3%
Hex.	Medium	1&2	161	3%
Hex.	Medium	1	86	3%
Hex.	Medium	2	75	4%
Hex.	Large	1&2	40	2%
Hex.	Large	1	18	1%
Hex.	Large	2	22	2%
Ellip.	Medium & Large	1&2	36	5%
R _{FT}				
Hex.	Medium & Large	1&2	201	6%
Hex.	Medium	1&2	161	6%
Hex.	Medium	1	86	6%
Hex.	Medium	2	75	6%
Hex.	Large	1 & 2	40	7%
Hex.	Large	1	18	5%
Hex.	Large	2	22	8%
Ellip.	Medium & Large	1&2	36	10%

^{510 &}lt;sup>a</sup> Groups in **bold** are the groups for which uncertainties are reported (i.e., geometry only for V and RFT; geometry and grain size for FT).

Appendix **D**: In the case of **2D** calculations using the minimum and maximum width

We recommend using the maximum width only for apatite 2D calculations for the reasons discussed in Section 6.1. However, for completeness, in this Appendix we present a set of corrections and

515 <u>uncertainties based on our dataset that can be used if both maximum and minimum width measurements</u> are acquired and used to calculate the 2D parameters (Fig. D2, Table D1).

Deleted: E

Deleted: E

Deleted: ¶

		Deleted: measuring
	V X	Deleted: suggest
)		Deleted: we recognize that some labs may want to continue measuring and using the minimum width to calculate V, F_T , and R_{FT} , so
	(Deleted: T
<u>s</u>		Deleted: in this case
	(Deleted: .
	\searrow	Deleted: E

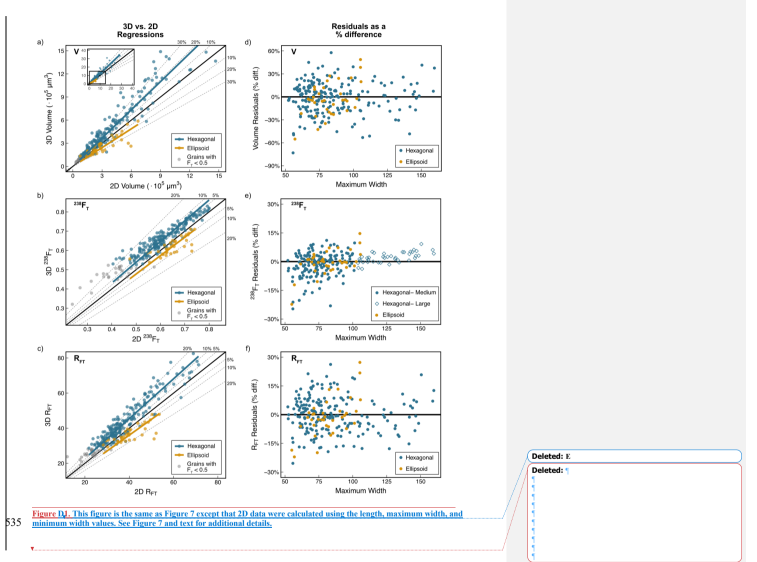


Table D1. Corrections and uncertainties (1s) for all geometric parameters where 2D values are calculated using the length, maximum width, and minimum width.

Formatted: Font: 9 pt, Bold

Volume		
		% Uncert. ^b (1s) for apatite grains
Geometry	Correction ^a	of all sizes
Volume		
Hex.	1.27	21%
Ellip.	0.86	28%

Isotope-specific F_T values

	% Uncert. (1s) for	% Uncert. (1s)
	medium-sized ^c	for large-sized ^d
Correction	apatite grains	apatite grains
1.08	6%	3%
0.96	6%	6%
1.08	8%	4%
0.95	7%	7%
1.08	8%	4%
0.95	7%	7%
1.02	2%	1%
0.98	1%	1%
	1.08 0.96 1.08 0.95 1.08 0.95 1.02	medium-sized ^c apatite grains 1.08 6% 0.96 6% 1.08 8% 0.95 7% 1.08 8% 0.95 7% 1.08 8% 0.95 7% 1.08 8% 0.95 7% 1.08 8% 0.95 2%

R_{FT}

		% Uncert. (1s) for apatite grains of
Geometry	Correction	all sizes
R _{FT}		
Hex.	1.15	9%
Ellip.	0.91	10%

* The correction value is the slope of the 3D vs. 2D regression line for each parameter in Figures D1a-c.

 ^b The uncertainty is the scatter of the 2D data about each regression line in Fig. D1_{4/C}, calculated as the 1s standard deviation of the % difference of each 2D value from the regression line (Fig. D1_{4/C}).

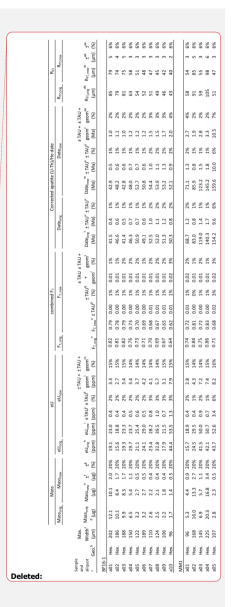
c "Medium-sized" apatite have maximum widths of 50-100 μm.

^d "Large-sized" apatite have maximum widths of >100 μm.

Deleted: Table E1.	
Deleted: 6A	
Deleted: 6C	
Deleted: A	
Deleted: C	
Deleted: D	
Deleted: F	

565	Appendix E: Application of geometric parameter corrections and uncertainties to a real dataset.	(Deleted: ¶
	ייים או איז	(Deleted: D
	Table <u>[1]</u> . Results of applying geometric corrections and uncertainties (1s) to apatite (U-Th)/He data from <u>a suite of</u> samples previously dated in the CU TRaIL.	(Deleted: D1

1			*+ (70)	8		89	6%	6%	6%	6%	6%	6%	6%	6%	6%		88	6%	6%	6%	6%	6%	6%	6%	6%	6%		10%	%01	10%	
	new		R+ (41)			۰ N	4	ŝ	m	m	m	e	m	m	2		~	ŝ	е	9	m	4	4	4	4	s		4	4	4	
Rrt	R		R _{FT, new} ^{ac}	(und)		79	74	75	58	51	48	47	45	42	40		54	85	55	98	47	70	68	72	60	76		37	36	36	
	RFT,orig		RFT,orig ^{ab} F	(mm)		58 1	79	81	63	54	52	51	49	46	43		82	91	59	105	51	75	74	77	64	82		43	43	42	
j		+ NV.	geom ³³ F	R		2%	2%	2%	2%	2%	2%	3%	3%	3%	4%		4%	2%	2%	2%	7%	5%	4%	4%	5%	4%		6%	6%	8%	
		FAU+±1	geom ² ge	(PIM)		1.0	1.1	1.0	1.2	1.2	1.2	1.5	1.5	1.7	2.0		2.7	1.9	2.8	3.3	10.5	4.4	4.0	4.1	6.1	4.0		0.2	0.2	0.4	
date	ite _{new}	+	TAU' B			1%	1%	1%	1%	1%	1%	2%	2%	2%	2%		2%	1%	1%	1%		4%	3%	3%	5%	4%		3%	2%	6%	
J-Th)/He	D		± TAU ^X ±			0.5	0.6	9.0	0.7	0.7	0.6	1.0	1.1	1.3	6.0		1.3	0.8	1.5	1.8	10.0	4.0	3.4	3.4	5.7	3.5		0.1	0.08	0.33	
Corrected apatite (U-Th)/He date			Datenew ±TAU ^X ±TAU ^V			42.9	48.2	42.8	48.0	52.7	50.8	54.4	53.9	53.2	52.1		71.1	85.9	123.2	145.2	159.6	90.7	104.0	116.2	115.7	95.4		3.3	3.2	5.3	
orrected	Massing Massing Object F1,col F1,col Bassing B					1%	1%	1%	1%	1%	1%	2%	2%	2%	2%		2%	1%	1%	1%	9%	4%	3%	3%	2%	4%		3%	3%	8%	
	iteorig		ETAU" ±			0.4	0.6	0.5	0.7	0.7	0.6	1.0	1.1	1.2	0.8		12	0.8	1.4	1.7	9.6	3.9	3.3	3.3	5.4	3.4		0.1	0.1	0.3	
	eU _{rev} F _{1,vel} Date _{ele} Date _{ele}		Dateore [®] ± TAU ⁴¹ ± TAU ^V			41.5	46.6	41.4	46.3	50.9	49.1	52.5	52.0	51.3	50.3		68.7	83.0	119.0	140.3	154.2	87.6	100.4	112.2	111.7	92.1		3.0	3.0	4.8	
j		TAU +	geom ⁵ 1	R		1%	1%	2%	2%	2%	1%	2%	2%	2%	3%		2%	1%	2%	1%	3%	2%	2%	2%	2%	2%		4%	4%	4%	
		± TAU ±	+ 1000	Gentli		0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.02		0.02	0.01	0.01	0.01	0.02	0.02	0.02	0.01	0.02	0.02		0.02	0.02	0.02	
ed F ₇	FT, new		± TAU ⁴			1%	1%	1%	1%	1%	1%	1%	1%	1%	1%		1%	%0	1%	1%	3%	2%	1%	1%	2%	1%		1%	1%	1%	
combined F _T			TALP			0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.01		10.0	00.0	0.00	00.0	0.02	0.01	0.01	0.01	0.01	0.01		0.00	0.01	0.01	
			, ,	T, new		0.79	0.78	0.79	0.73	0.70	0.69	0.68	0.67	0.65	0.62		0.72	0.81	0.72	0.83	0.68	0.77	0.77	0.78	0.74	0.79		0.60	0.60	0.60	
	FT, orig		c	FT, orig FT, new I IAU		0.82	18.0	0.82	0.76	0.73	0.71	0.70	0.69	0.67	0.64		0.74	0.84	0.75	0.86	0.71	0.80	0.80	0.81	0.77	0.82		0.66	0.65	0.65	
		: TAU +	geom	R		15%	15%	15%	14%	14%	14%	14%	14%	15%	15%		15%	14%	14%	15%	16%	15%	15%	15%	15%	15%		16%	16%	17%	
	Matching Matching Belleng Transment Date of the second sec	TAU + ±	geom' g	(mqq)		3.3	2.7	3.4	3.4	3.7	4.2	4.1	5.7	3.1	7.9		2.8	4.3	7.2	7.4	8.2	4.2	6.1	5.9	5.2	3.5		10.8	17.8	3.4	
۴Ū		*	±TAU ^k			2%	2%	2%	2%	2%	2%	3%	3%	3%	3%		2%	1%	2%	1%	8%	5%	4%	4%	5%	4%		2%	3%	3%	
Ĩ			eUnew ¹ ±TAU ¹ ±TAU ^k			0.4	0.4	0.4	0.5	0.6	0.5	0.8	1.0	0.7	1.3		0.4	0.4	0.9	0.7	3.4	1.2	1.6	1.4	1.7	6.0		1.0	3.2	0.7	
			eUnew ¹	(mdd)		23.0	18.8	23.3	23.7	25.4	29.0	28.2	39.5	21.5	53.5		18.9	29.5	50.0	50.7	52.6	27.3	42.0	40.4	34.4	23.5		66.1	108.1	20.7	
	eUorig		eU _{orig} ^h	(mqq)		19.1	15.6	19.3	19.7	21.1	24.1	23.4	32.8	17.9	44.4		15.7	24.5	41.5	42.1	43.7	22.7	34.8	33.5	28.5	19.5		48.9	80.0	15.3	
			4 5		- 1						20%	20%	20%	20%	20%		20%	20%	20%	20%	20%	20%	20%	20%	20%	20%		23%	23%	23%	
<u>،</u>	Mass _{new}		*+ (,+ ,* , *									0.4	0.4	0.4	0.3		0.9	2.7		3.4		1.2		1.4	0.9	1.8		0.3	0.2	0.2	
Mass			ŝ	18H		10.1	8.4	8.3	5.4	2.7	2.7	2.2	2.1	1.8	1.4		4.4	13.3	5.7	16.8	2.3	5.9	6.0	7.2	4.4	9.0		1.1	0.9	0.9	
	Massorig		Width ^c Massorig	191		12.1	10.1	6.6	6.5	3.2	3.2	2.6	2.5	2.2	1.7		5.3	16.0	6.9	20.3	2.8	7.2	7.2	8.6	5.3	10.8		1.5	1.2	1.2	
		Max.									189	110	124	106	96		96	168	145	225		154	153	154	133	159			79		
		ple	Lot Gan ^b	neo.				_	Hex.		Hex.	Hex.	Hex.	Hex.	Hex.	Hex.	Hex.	Hex.	Hex.	Hex.	16MFS05	Ellip.	Ellip.	Ellip.							
		Samp	aliquot		BF16-1	a01	a02	a03	a04	a05	a06	a07	a08	90e	a10	IMM	a01	a02	a03	a04	a05	a06	a07	a08	e0e	a10	16	a02	a03	a04	



	All uncertainties reported at the 1s level.		
575	All calculations done assuming F_T uncertainties are fully correlated (r = 1).		
	^a All BF16-1, MM1, and 16MFS05 data are published in Flowers and Kelley (2011), Weisberg et al. (2018), and Collett et al. (2019), respectively.		
	^b Geometry is defined as described in Figure 3 of Ketcham et al. (2011). All GEM A and B grains are hexagonal (hex.) and all GEM C grains are ellipsoid (ellip.).		
580	^c Maximum width is measured perpendicular to the <u>length/c-axis</u> .		Deleted: c-ais
	^d Mass _{orig} is the mass of the crystal determined by 2D microscopy measurements, the volume assuming the reported grain geometry, and the volume equations and mineral densities in Ketcham et al. (2011).		
I	^e Mass _{new} is computed the same as mass _{orig} , but the original V is corrected by applying the correction factor in Table 2 based on the grain geometry, and this new volume is used in the mass calculation.		
585	^f The 1s uncertainty on mass _{new} is calculated by propagating the uncertainty on V from Table 2 based on grain geometry through the mass equation.		
	^g The 1s percent uncertainty on mass _{new} .		
I	^h eU _{ariz} is effective Uranium concentration calculated using the mass _{orig} . Calculated as U + 0.238*Th + 0.0012*Sm after equation A <u>7 of Cooperdock et al. (2019)</u> ,		Deleted: 7
590	ⁱ eU _{new} is computed the same as eU _{orig} but uses the mass _{new} value.	(Deleted: of Cooperdock et al. (2019).
	^j The 1s total analytical uncertainty (TAU, which are the uncertainties on the parent isotopes) on eU. This calculation ignores the negligible contribution from Sm concentration uncertainty and uses 0% geometric uncertainty.		
	^k The 1s total analytical percent uncertainty on eU _{new} .		
595	¹ The 1s TAU + geometric uncertainty on eU _{new} . This <u>uncertainty includes the total analytical uncertainty and</u> , the uncertainty assigned based on grain geometry (Table 2), assumes that the geometric uncertainties on U and Th concentrations are perfectly correlated (<u>r = 1</u>), and ignores the negligible contribution from Sm concentration uncertainty. Although the correlation coefficient will vary with each data set, the dominant contribution to concentration uncertainty comes from the volumetric uncertainty, which is highly correlated. Additionally, assuming perfect correlation yields the maximum possible value, so we use this conservative approach.	(Deleted: calculation Deleted: uses
600	^m The 1s total analytical + geometric percent uncertainty on eU _{new} .		
	ⁿ $F_{T,orig}$ is the combined alpha-ejection correction for the crystal calculated from the original parent isotope-specific F_T corrections, the proportion of U and Th contributing to the ⁴ He production, and assuming homogeneous parent isotope distributions using equation A4 in Cooperdock et al. (2019). The parent isotope-specific alpha ejection-corrections were computed assuming the reported grain geometry in this table and the equations and alpha-stopping distances in Ketcham et al. (2011).		
605	^o F _{T,new} is computed the same as F _{T,orig} but uses isotope-specific F _{T,new} values corrected by applying the correction factors in Table 2 based on grain geometry and size.		
	$^{\text{p}}$ The 1s TAU on $F_{\mathrm{T,new}}.$ This calculation uses 0% geometric uncertainty.		
	^q The 1s total analytical percent uncertainty on F _{T,new} .		
610	^r The 1s TAU + geometric uncertainty. This <u>uncertainty includes the total analytical uncertainty and</u> uses the parent isotope- specific $F_{T,new}$ uncertainties assigned based on grain geometry and size (Table 2).	(Deleted: calculation

	^s The 1s total analytical + geometric percent uncertainty on $F_{T,new}$.	
	^t The corrected (U-Th)/He date _{orig} is calculated iteratively using the absolute values of He, U, Th, Sm, the isotope-specific $F_{T,orig}$ values, and equation 34 in Ketcham et al. (2011) assuming secular equilibrium.	
620	^u The 1s TAU uncertainty on date _{orig} includes the propagated total analytical uncertainties on the U, Th, Sm and He measurements. Uncertainty propagation done using HeCalc (Martin et al., 2023) ₂₇	 Deleted: (Martin et al., in review).
I	^v The 1s total analytical percent uncertainty on date _{orig} .	
	^w The corrected (U-Th)/He date _{new} is computed the same as date _{orig} , but uses the isotope-specific $F_{T,new}$ values corrected by applying the correction factors in Table 2 based on grain geometry and size.	
625	* The 1s TAU uncertainty on the corrected (U-Th)/He datence includes the propagated total analytical uncertainties on the U, Th,	
	Sm, He measurements, This calculation uses 0% geometric uncertainty. Uncertainty propagation done using HeCalc (Martin et	 Deleted: (Table 2)
	al., 2023)	 Deleted: (Martin et al., in review)
I	^y The 1s total analytical percent uncertainty on the corrected (U-Th)/He date _{new} .	
20	* The 1s total analytical + geometric uncertainty on the corrected (U-Th)/He date _{new} . This <u>uncertainty includes the propagated</u>	 (Deleted: calculation
630	total analytical uncertainties on the U, Th, Sm, He measurements and uses the parent isotope-specific $F_{T,new}$ uncertainties assigned based on grain geometry and size (Table 2).	
	^{aa} The 1s total analytical + geometric percent uncertainty on the corrected (U-Th)/He date _{new} .	
1	^{ab} R _{FT.orig} is the radius of a sphere with an equivalent alpha-ejection correction as the grain, calculated using the uncorrected parent isotope-specific F _g values in equation A6 in Cooperdock et al. (2019).	 Formatted: Subscript
635	^{ac} R _{FT,new} is computed from R _{FT,orig} by multiplying R _{FT,orig} by the correction factor in Table 2 based on grain geometry.	
	^{ad} The 1s uncertainty on R _{FT,new} is assigned based on grain geometry (Table 2).	
	^{ae} The 1s percent uncertainty on R _{FT,orig} .	

Code and Data Availability

Data and code used to produce the corrections and uncertainties and figures is stored through the Open Science Framework: <u>https://osf.io/fu98s/</u>. All analyses and plots were done in R (Wickham et al., 2019).

Author Contributions

RMF and JRM conceptualized the project; SDZ curated the data; SDZ and JRM performed the formal data analysis; RMF, JRM, and SDZ acquired funding; SDZ performed the investigation; JRM, SDZ, and RMF developed the methodology; RMF provided supervision; SDZ performed the validation; SDZ did the data visualizations; SDZ and RMF wrote the original draft; RMF, SDZ, and JRM reviewed and

645 did the data visualizations; SDZ and RMF wrote the original draft; RMF, SDZ, and JRM reviewed an edited the manuscript.

Competing Interests

The authors declare they have no conflict of interest, Deleted:

Acknowledgments

We thank Alison Duvall, Lon Abbott, Ray Donelick, and Jacky Baughman for samples. We are grateful to Jennifer Coulombe and Adrian Gestos for their assistance, support, and advice regarding nano-CT and Dragonfly. The use of Blob3D would not have been possible without support from Rich Ketcham and Romy D. Hanna. We appreciate numerous discussions with Morgan Baker. Thanks to Peter Martin for advice and for HeCalc. <u>Rich Ketcham and Christoph Glotzbach provided insightful reviews that</u> improved the clarity of this manuscript. This work was partially funded by National Science Foundation

660 GRFP DGE-1650115 to Zeigler. Funding for the Zeiss Xradia Versa X-Ray Microscope was provided by NSF CMMI-1726864.

References

Abbey, A. L., Niemi, N. A., Geissman, J. W., Winkelstern, I. Z., and Heizler, M.: Early Cenozoic exhumation and paleotopography in the Arkansas River valley, southern Rocky Mountains, Colorado,
 Lithosphere, 10, 239–266, https://doi.org/10.1130/L673.1, 2017.

Abbott, L. D., Flowers, R. M., Metcalf, J., Falkowski, S., and Niazy, F.: Post-Laramide, Eocene

epeirogeny in central Colorado—The result of a mantle drip?, Geosphere, 18, 1223–1246, https://doi.org/10.1130/GES02434.1, 2022.

670 Baker, M., Zeigler, S. D., Metcalf, J. R., and Flowers, R. M.: Estimating Uncertainties for Alpha-Ejection Corrections and eu Values for the Zircon U-Th Method, in: Fall Meeting 2020, American Geophysical Union, Online, https://doi.org/2020AGUFMV031.0004B, 2020.

Baughman, J. S., Flowers, R. M., Metcalf, J. R., and Dhansay, T.: Influence of radiation damage on titanite He diffusion kinetics, Geochimica et Cosmochimica Acta, 205, 50–64,
https://doi.org/10.1016/j.gca.2017.01.049, 2017.

Brown, R. W., Beucher, R., Roper, S., Persano, C., Stuart, F., and Fitzgerald, P.: Natural age dispersion arising from the analysis of broken crystals. Part I: Theoretical basis and implications for the apatite (U–Th)/He thermochronometer, Geochimica et Cosmochimica Acta, 122, 478–497, https://doi.org/10.1016/j.gca.2013.05.041, 2013.

680 Collett, C. M., Duvall, A. R., Flowers, R. M., Tucker, G. E., and Upton, P.: The Timing and Style of Oblique Deformation Within New Zealand's Kaikōura Ranges and Marlborough Fault System Based on Deleted: Finally, t

Low-Temperature Thermochronology, Tectonics, 38, 1250–1272, https://doi.org/10.1029/2018TC005268, 2019.

Cooperdock, E. H. G., Ketcham, R. A., and Daniel F. Stockli: Resolving the effects of 2-D versus 3-D grain measurements on apatite (U–Th)/He age data and reproducibility, Geochronology, 1, 17–41, https://doi.org/10.5194/gchron-1-17-2019, 2019.

690	Dragonfly. Object Research Systems. Montréal, Quebec, Canada. https://theobjects.com/dragonfly, 2020.		Deleted: Dragonfly: Dragonfly, 2020.
		•	Formatted: Normal
	Ehlers, T. A. and Farley, K. A.: Apatite (U-Th)/He thermochronometry: methods and applications to		
	problems in tectonic and surface processes, Earth and Planetary Science Letters, 206, 1–14,		
	https://doi.org/10.1016/S0012-821X(02)01069-5, 2003.		

Deleted:

695 Evans, N. J., McInnes, B. I. A., Squelch, A. P., Austin, P. J., McDonald, B. J., and Wu, Q.: Application of X-ray micro-computed tomography in (U–Th)/He thermochronology, Chemical Geology, 257, 101– 113, https://doi.org/10.1016/j.chemgeo.2008.08.021, 2008.

Farley, K. A., Wolf, R. A., and Silver, L. T.: The effects of long alpha-stopping distances on (U₄Th)/He ages, Geochimica et Cosmochimica Acta, 60, 4223–4229, https://doi.org/10.1016/S0016-700 7037(96)00193-7, 1996.

Flowers, R. M. and Kelley, S. A.: Interpreting data dispersion and "inverted" dates in apatite (U–Th)/He and fission-track datasets: An example from the US midcontinent, Geochimica et Cosmochimica Acta, 75, 5169–5186, https://doi.org/10.1016/j.gca.2011.06.016, 2011.

 Flowers, R. M., Shuster, D. L., Wernicke, B. P., and Farley, K. A.: Radiation damage control on apatite
 (U-Th)/He dates from the Grand Canyon region, Colorado Plateau, Geology, 35, 447–450, https://doi.org/10.1130/G23471A.1, 2007.

Flowers, R. M., Zeitler, P. K., Danišík, M., Reiners, P. W., Gautheron, C., Ketcham, R. A., Metcalf, J. R., Stockli, D. F., Enkelmann, E., and Brown, R. W.: (U-Th)/He chronology: Part 1. Data, uncertainty, and reporting, Geological Society of America Bulletin, 130, 33, 2022a.

710 Flowers, R. M., Ketcham, R. A., Enkelmann, E., Gautheron, C., Reiners, P. W., Metcalf, J. R., Danišík, M., Stockli, D. F., and Brown, R. W.: (U-Th)/He chronology: Part 2. Considerations for evaluating, integrating, and interpreting conventional individual aliquot data, Geological Society of America Bulletin, 130, 25, 2022b.

Gleadow, A., Harrison, M., Kohn, B., Lugo-Zazueta, R., and Phillips, D.: The Fish Canyon Tuff: A new
 look at an old low-temperature thermochronology standard, Earth and Planetary Science Letters, 424, 95–108, https://doi.org/10.1016/j.epsl.2015.05.003, 2015.

720	Glotzbach, C., Lang, K. A., Avdievitch, N. N., and Ehlers, T. A.: Increasing the accuracy of (U-Th(-Sm))/He dating with 3D grain modelling, Chemical Geology, 506, 113–125, https://doi.org/10.1016/j.chemgeo.2018.12.032, 2019.		
725	Harbert, S. A.: Landscape Response to Oblique Convergence: Insights from Numerical Modeling and from the Marlborough Fault System, New Zealand, Thesis, https://digital.lib.washington.edu:443/researchworks/handle/1773/44635, 2019. He, J. J. Y. and Reiners, P. W.: A revised alpha-ejection correction calculation for (U–Th) / He		Deleted: Harbert, S., Duvall, A. R., Upton, P., Flowers, R. M., and O'Sullivan, P. B.: Mid-Miocene Development of the Alpine-Wairau Fault Restraining Zone, South Island, New Zealand, in review.¶ Formatted: Normal
730	thermochronology dates of broken apatite crystals, 12, 2022. Herman, F., Braun, J., Senden, T. J., and Dunlap, W. J.: (U–Th)/He thermochronometry: Mapping 3D geometry using micro-X-ray tomography and solving the associated production–diffusion equation, Chemical Geology, 242, 126–136, https://doi.org/10.1016/j.chemgeo.2007.03.009, 2007.		Formatted: Bibliography
	Hoffman, P. F.: United Plates of America, The Birth of a Craton: Early Proterozoic Assembly and Growth of Laurentia, Annual Review of Earth and Planetary Sciences, 16, 543–603, https://doi.org/10.1146/annurev.ea.16.050188.002551, 1988.	.	Formatted: Normal
735	Hourigan, J. K., Reiners, P. W., and Brandon, M. T.: U-Th zonation-dependent alpha-ejection in (U-Th)/He chronometry, Geochimica et Cosmochimica Acta, 69, 3349–3365, https://doi.org/10.1016/j.gca.2005.01.024, 2005.		
	JCGM: Joint Committee for Guides in Metrology – The International Vocabulary of Metrology – Basic and General Concepts and Associated Terms, 2012.		
740	Ketcham, R. A.: Computational methods for quantitative analysis of three-dimensional features in geological specimens, Geosphere, 1, 32–41, https://doi.org/10.1130/GES00001.1, 2005.v	<	Deleted: Ketcham, R. A.: Forward and Inverse Modeling of Low- Temperature Thermochronometry Data, Reviews in Mineralogy and Geochemistry, 58, 275–314, https://doi.org/10.2138/rmg.2005.58.11, 2005.¶
745	Journal of Microscopy, 213, 158–171, https://doi.org/10.1111/j.1365-2818.2004.01277.x, 2004. Ketcham, R. A., Gautheron, C., and Tassan-Got, L.: Accounting for long alpha-particle stopping distances in (U–Th–Sm)/He geochronology: Refinement of the baseline case, Geochimica et Cosmochimica Acta, 75, 7779–7791, https://doi.org/10.1016/j.gca.2011.10.011, 2011.		Formatted: Normal
	Martin, P. E., Metcalf, J. R., and Flowers, R. M.: Calculation of uncertainty in the (U–Th)/He system, Geochronology, 5, 91–107, https://doi.org/10.5194/gchron-5-91-2023, 2023.		Deleted: & Deleted: thinsp:@thinsp;
750	Meesters, A. G. C. A. and Dunai, T. J.: Solving the production–diffusion equation for finite diffusion domains of various shapes: Part II. Application to cases with α -ejection and nonhomogeneous distribution of the source, Chemical Geology, 186, 57–73, https://doi.org/10.1016/S0009-		

2541(01)00423-5, 2002.

765	Rahl, J. M., Reiners, P. W., Campbell, I. H., Nicolescu, S., and Allen, C. M.: Combined single-grain (U- Th)/He and U/Pb dating of detrital zircons from the Navajo Sandstone, Utah, Geology, 31, 761–764, https://doi.org/10.1130/G19653.1, 2003.	Formatted: Bibliography
770	Geological & Nuclear Sciences (NZ), 2006. Rodgers, D. W.: Thermal and Structural Evolution of the Southern Deep Creek Range, West Central Utah and East Central Nevada., 1, 1988. Shuster, D. L., Flowers, R. M., and Farley, K. A.: The influence of natural radiation damage on helium	Formatted: Normal
775	diffusion kinetics in apatite, Earth and Planetary Science Letters, 249, 148–161, https://doi.org/10.1016/j.epsl.2006.07.028, 2006. Spell, T. L. and McDougall, I.: Characterization and calibration of 40Ar/39Ar dating standards, Chemical Geology, 198, 189–211, https://doi.org/10.1016/S0009-2541(03)00005-6, 2003.	
780	Van Schmus, W. R., Bickford, M. E., and Zietz, I.: Early and Middle Proterozoic provinces in the central United States, in: Geodynamics Series, vol. 17, edited by: Kröner, A., American Geophysical Union, Washington, D. C., 43–68, https://doi.org/10.1029/GD017p0043, 1987. Weisberg, W. R., Metcalf, J. R., and Flowers, R. M.: Distinguishing slow cooling versus multiphase	Formatted: Normal
	cooling and heating in zircon and apatite (U-Th)/He datasets: The case of the McClure Mountain syenite standard, Chemical Geology, 485, 90–99, https://doi.org/10.1016/j.chemgeo.2018.03.038, 2018.	
785	Wickham, H., Averick, M., Bryan, J., Chang, W., McGowan, L. D., François, R., Grolemund, G., Hayes, A., Henry, L., Hester, J., Kuhn, M., Pedersen, T. L., Miller, E., Bache, S. M., Müller, K., Ooms, J., Robinson, D., Seidel, D. P., Spinu, V., Takahashi, K., Vaughan, D., Wilke, C., Woo, K., and Yutani, H.: Welcome to the tidyverse, Journal of Open Source Software, 4, 1686, https://doi.org/10.21105/joss.01686, 2019.	

<i>I</i>			4
Page 35: [2] Deleted	Spencer Zeigler	2/8/23 8:37:00 AM	
,			
Page 37: [3] Deleted	Spencer Zeigler	1/24/23 10:09:00 AM	
Page 38: [4] Formatted	Becky Flowers	2/2/23 5:39:00 PM	
Subscript	becky Howers	2/2/25 5.55.00 FM	
-			
Page 38: [4] Formatted	Becky Flowers	2/2/23 5:39:00 PM	
Subscript			
Page 38: [4] Formatted	Becky Flowers	2/2/23 5:39:00 PM	
Subscript			
Page 38: [4] Formatted	Becky Flowers	2/2/23 5:39:00 PM	
Subscript			
Page 38: [4] Formatted	Becky Flowers	2/2/23 5:39:00 PM	
Subscript			
Page 38: [5] Deleted	Becky Flowers	2/2/23 3:25:00 PM	
			4
Page 38: [5] Deleted	Becky Flowers	2/2/23 3:25:00 PM	
			.
Page 38: [5] Deleted	Becky Flowers	2/2/23 3:25:00 PM	
raye so: [5] Deleteu	Becky riowers	2/2/23 3:23:00 FM	
Page 38: [5] Deleted	Becky Flowers	2/2/23 3:25:00 PM	

L

I

L

L

L