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# Multi-aliquot method for determining (U + Th)/He ages of hydrothermal hematite: Returning to Elba



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#### ABSTRACT

We have used a multi-aliquot method to obtain precise (U + Th)/He ages of hydrothermal hematite and to assess the extent to which He loss from fine-grained hematite caused by diffusion and recoil. Hematite (n = 6) from the Rio Marina mine, Elba (Italy) yields (U + Th)/He ages that range from 5.36  $\pm$  0.33 to 5.64  $\pm$  0.11 Ma, giving a weighted mean age of 5.53  $\pm$  0.14 Ma and an isochron age of 5.25  $\pm$  0.20 Ma.  $^{40}\text{Ar}/^{39}\text{Ar}$  data from cogenetic adularia yield flat age spectra with analytically indistinguishable plateau ages (5.575  $\pm$  0.008 and 5.583  $\pm$  0.013 Ma). An additional adularia has a more complex spectrum and yields an interpreted age of 5.64  $\pm$  0.03 Ma. The hematite (U + Th)/He ages overlap the  $^{40}\text{Ar}/^{39}\text{Ar}$  ages, albeit they are less precise (2–6% vs. 0.2–0.5%). This indicates that the loss of in situ radiogenic  $^4\text{He}$  from complex fine-grained hematite, either by diffusion and recoil, is insignificant. The study shows that multi-aliquot method has the potential to reliably deliver precise and accurate ages for iron oxide mineralisation that has not suffered significant post-crystal-lisation thermal perturbation.

#### 1. Introduction

The incorporation of modest amounts of U and Th (few ppm) in iron oxide minerals such as goethite, hematite and magnetite makes them candidates for (U + Th)/He dating. Early studies yielded geologically reasonable (U + Th)/He ages from Pliocene to Mesozoic hematite (Wernicke and Lippolt, 1993, 1994a, 1994b, 1997; Lippolt et al., 1995). Systematic laboratory studies have shown that the  $^4\text{He}$  closure temperature (Tc) of hematite depends on the grain size; massive hematite with grain diameters in excess of 100  $\mu\text{m}$  has Tc  $> 200\,^{\circ}\text{C}$ , while it approaches 50  $^{\circ}\text{C}$  in fine-grained crystallites typical of botryoidal hematite (< 1  $\mu\text{m}$ ) (e.g. Lippolt et al., 1993; Bähr et al., 1994; Farley and Flowers, 2012). This has been confirmed by computational methods (Balout et al., 2017).

Most (U—Th)/He dating studies of iron oxides/oxyhydroxides measure the He, U and Th in the same sub-mg-sized fragments of material (e.g. Farley and Flowers, 2012; Vasconcelos et al., 2013). This technique allows dating of intergrown and fine-grained minerals and in recent years it has proved to be a powerful method for determining the timing of processes that have hitherto proved difficult, such as the

timing of fault movement (e.g. Evenson et al., 2014; Ault et al., 2015; McDermott et al., 2017) and regional weathering events (Shuster et al., 2005; Cooper et al., 2016; Deng et al., 2017). However, the method is analytically challenging and has complexities that are often not evident from age data. For instance, over-heating of the iron oxides/oxy-hydroxides can result in the volatilization loss of U (and Th) leaving the resultant ages significantly younger than the crystallisation age (e.g. Vasconcelos et al., 2013; Cooper et al., 2016). Under-heating of samples can lead to the incomplete degassing of He yielding ages that are significantly younger than the crystallisation age (e.g. Danišík et al., 2013). Multiple analyses of same age material typically yield strongly (  $\pm$  20%) over-dispersed ages, even after rejection of obviously overheated grains (e.g. Cooper et al., 2016).

Here we present new (U + Th)/He ages of hydrothermal hematite where  $^4He$  is measured on separate aliquots of the same sample from U and Th. We revisit the seminal study of Lippolt et al. (1995) who determined (U + Th)/He ages of hydrothermal hematite and compared them to K-Ar ages of cogenetic adularia from the Miocene-aged hydrothermal mineralisation from the island of Elba, Italy. We report

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Fig. 1. Images of hematite ore samples from San Marino mine, Elba (Italy) used in this study. (A) Italy1: micaceous specular hematite and adularia. (B) Italy2: micaceous specular hematite and euhedral adularia (1–2 mm). (C) Italy3: micaceous specular hematite and prismatic adularia. (D) M1392: massive euhedral, hematite rhombohedrons. (E) Italy4: micaceous specular hematite with coarse grained vein

multi-aliquot hematite He ages and high precision adularia  $^{40}\text{Ar}/^{39}\text{Ar}$  ages with the aim of assessing the age precision attainable using the multi-aliquot technique, and determining the extent to which diffusion and  $\alpha$  recoil result in  $^4\text{He}$  loss from hematite.

#### 2. Sample location and description

Several iron ore deposits are present on the eastern coast of Elba between Monte Calendozio and Capo Calamita. Three major iron ore types can be distinguished on Elba Island (Tanelli et al., 2001; Dünkel, 2002): Rio Marina type (deposits of Rio Marina and Rio Albano, northeastern Elba, characterized by lenticular bodies of hematite-pyrite (+limonite)), Ortano type (including the deposits of Ortano and Terra Nera (central east), characterized by hematite-magnetite-pyrite), and Calamita type (deposits of Sassineri, Ginevro and Calamita, comprising lenses and massive bodies of magnetite ± hematite associated to skarn bodies in the Calamita Peninsula (southeast)). The deposits were likely formed contemporaneously with the shallow intrusion of the nearby Porto Azzurro quartz monzonitic intrusion at 5.9 ± 0.2 Ma (Maineri et al., 2003). There is little consensus on the source of the iron in the Elba deposits; it may be sedimentary in origin, remobilised by hydrothermal fluids (Tanelli and Lattanzi, 1986; Dünkel, 2002) or of magmatic origin (Lotti, 1929; Gillieron, 1959).

The ore bodies of the first type at the Rio Marina mine are typically stratiform and vein-type, hosted by quartz-phyllite units of the Carboniferous-Triassic Tuscan Nappe, often close to the contact with overlying carbonates. The hematite in the deposits is typically lamellar/micaceous, associated with pyrite, quartz, and chlorite. Iron ores from the Valle Giove stope also contain abundant adularia and have been targeted in this study.

We have analysed vein hematite from the Valle Giove stope in the Rio Marina mine district. Three samples (Italy-1, -2 and -3) are dominated by specular (micaceous) hematite and contain abundant cogenetic adularia (Fig. 1). Hand specimen Italy-4 contains both coarse-(Italy-4c) and fine-grained (4f) micaceous hematite. Sample M1392 is a massive (oligistic) hematite mostly formed of flattened rhombohedrons that are up to 2 cm long. Hematites from Italy-1 to -4 are generally comprised of aggregates of plate-shaped crystals that are 20-5 µm thick (Fig. 2A and B). The oligistic hematite of M1392 is poly-crystalline, comprised of less well-ordered hematite plates that are significantly thinner than the micaceous hematites (Fig. 2C). All hematites are compositionally homogeneous and there is no evidence of mineral inclusions or recrystallisation/zonation (Fig. 2D and E). Back-scattered electron (BSE) and electron microprobe element mapping of adularias show that they are homogeneous and contain no significant Ca (Fig. 2 F-J). This is consistent with the Ar/Ar data (Table 2). There is no

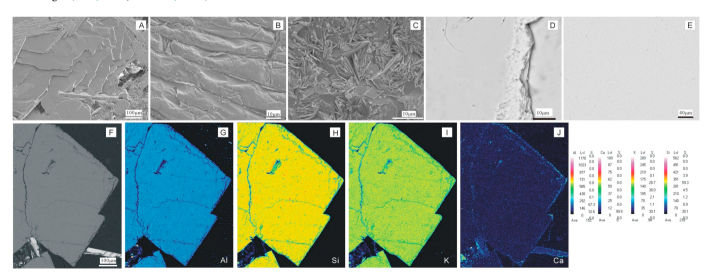


Fig. 2. (A and B). Scanning electron microscope (SEM) back-scattered images of polished fragments of Italy-1 hematite showing the polycrystalline aggregates of plate-shaped hematite. These features are typical of all Italy-x samples. (C) SEM back-scattered image of rhombohedron hematite of M1392. The thickness of each layer of rhombohedron is probably  $< 1 \, \mu m$ . (D) back-scattered image of plate-shaped hematite show the homogeneous texture. These features are typical of all Italy-x samples. (E) back-scattered image of rhombohedron hematite of M1392 show the homogeneous texture. (F–J) show back-scattered electron micoprobe and element maps of Italy-1 adularia showing homogeneous texture and lack of zonation and alteration. These features are typical of all adularia samples.

L.-Y. Wu et al. Chemical Geology 504 (2019) 151–157

indication of multiple generations of adularia growth nor significant sericite alteration.

#### 3. Analytical procedures

#### 3.1. Hematite $(U + Th)/^4He$ dating

The analytical procedure reported here differs from the majority of previous studies that measure He, U and Th concentrations on the same aliquot of hematite. In this study we have measured He and U—Th in separate aliquots of hematite from the same sample. This avoids the problem of incomplete He extraction and U volatilization due to underand over-heating respectively, though it requires significantly more laboratory effort and the availability of larger sample mass (Farley and McKeon, 2015).

Several grams of hematite were carefully removed from each sample to minimize the incorporation of gangue minerals. The hematite fragments were crushed and any visible gangue minerals removed from the  $100-200\,\mu m$  fraction by handpicking under a binocular microscope. The handpicked separate was then gently crushed and fully mixed.

Aliquots of 10-20 mg of  $\sim 30 \,\mu\text{g}$  grains were weighed into  $10 \times 2.5 \, \text{mm}$  Pt-foil tubes. The tubes were crimped at both ends then placed into recesses in a fully degassed Cu pan then pumped to  $< 10^{-8}$  Torr prior to degassing at 80 °C for 3 h. Helium was extracted by heating each packet to ~1300 °C for 10 min by rastering a diode laser beam along the length of the tube. The temperature of the packet was assessed by visual observation of the color emitted from heated Pt tubes. A sapphire cover glass was used to avoid volatilized metal from absorbing onto the sapphire viewport. The liberated gases were purified by exposure to two liquid nitrogen-cooled charcoal traps during heating and for a further 15 min. <sup>4</sup>He abundances along with H (mass 2) and CH<sub>4</sub> (mass 16) were determined by an electron multiplier in a Hiden HAL3F quadrupole mass spectrometer operated in static mode following procedures in Foeken et al. (2006). Absolute He concentrations were calculated by peak height comparison against a calibrated He standard. The precision of He measurements was determined by repeated measurements of the gas standard. Within-day reproducibility of measurement of the  $^4$ He standard was typically  $\pm 0.4\%$  (2 $\sigma$ , n = 10-12). Laser heating of empty Pt tubes yielded He levels  $(1.6 \times 10^{-11} \, \text{ml}, \, n = 9)$  that were close to system background levels. To verify complete extraction of He each sample was re-heated to 1300-1400 °C for a further 10 min. Between three and five aliquots of each sample were measured.

U and Th concentrations were measured in three to five aliquots of hematite (3-11 mg) that had not been used for He extraction using an Agilent 7500ce Q-ICP-MS. An initial analysis of natural U and Th in each sample was done in order to optimise the spike amount for sample/spike ratios that were between 0.1 and 10. Each sample was loaded into a Teflon beaker with a dilute-spike solution equivalent to 2-4 ng of  $^{230}$ Th and 1-2 ng of  $^{235}$ U in 2 ml 12 M HCl. The beakers were then heated to 100 °C for 2 days to completely dissolve the iron oxide. The HCl was then evaporated off and the dried residues were refluxed with 2 ml of 1.5 M HNO<sub>3</sub> at 80 °C overnight. Pre-cleaned columns were filled with pre-cleaned TR-B50-S resin, and washed with 9 ml of 0.2 M HCl and 9 ml of 0.1 M HCl + 0.3 M HF. The resin was pre-conditioned using 9 ml of 1.5 M HNO3 before the sample was introduced in 2 ml 1.5 M HNO<sub>3</sub>. Matrix elements were removed by rinsing the columns with 12 ml 1.5 M HNO<sub>3</sub> and 2.5 ml 3 M HCl. U and Th were eluted using 12 ml mixture of 0.1 M HCl 0.3 M HF. The final elutes were evaporated to dryness on a hotplate at 80 °C then set to reflux with 2 ml of 0.8 M  $HNO_3$  for 24 h. The long-term blanks averaged 0.52 ng (n = 11) for  $^{238}$ U and 0.16 ng (n = 11) for  $^{232}$ Th. These levels were negligible in comparison to sample U and Th concentrations. The Sm content of Elba hematites have been shown by Lippolt et al. (1995) to be insignificant to the production of <sup>4</sup>He so were not determined.

#### 3.2. Adularia <sup>40</sup>Ar/<sup>39</sup>Ar dating

The  $^{40}$ Ar/ $^{39}$ Ar age determinations were performed at the New Mexico Geochronology Research Laboratory. Adularia was hand-picked from crushed samples then irradiated at the TRIGA reactor in Denver, Colorado for 12 h in the NM-290 package along with Fish Canyon Sanidine interlaboratory standard FC-2 with an assigned age of 28.201 Ma (Kuiper et al., 2008). Ages are calculated with a total  $^{40}$ K decay constant of 5.463  $\times$  10 $^{-10}$  a $^{-1}$  (Min et al., 2000).

After irradiation, six crystals of FC-2 from each of 8 monitor holes from a 24-hole irradiation tray, along with ~1 mg sample aliquots were loaded into a copper tray, evacuated and baked at 140 °C for 2 h. FC-2 crystals were fused and adularia was step-heated both with a CO<sub>2</sub> laser and the extracted gas was cleaned with a SAES NP10 getter operated at 1.6 A and a D50 getter at 20 °C for 30 s. The gas was analysed for argon isotopes using a ThermoScientific ARGUS VI multi-collector mass spectrometer equipped with five Faraday cups, and one ion counting multiplier (CDD). The configuration had <sup>40</sup>Ar, <sup>39</sup>Ar, <sup>38</sup>Ar, <sup>37</sup>Ar and <sup>36</sup>Ar on the H1, AX, L1, L2, and CDD detectors, respectively. H1, AX and L2 utilized Faraday detectors equipped with  $10^{13}\Omega$  resistors whereas L1 had a Faraday with a  $10^{14} \Omega$  resistor. The CDD is an ion counting detector with a dead time of 14 ns. All data acquisition was accomplished with NM Tech Pychron software and data reduction used Mass Spec (v.7.875) written by Al Deino at the Berkeley Geochronology Laboratory. Extraction line blank plus mass spectrometer background values are averages of numerous measurements interspersed with the unknown measurements. These values are 12  $\pm$  4%, 0.2  $\pm$  9%,  $0.08 \pm 10\%$ ,  $0.4 \pm 3\%$ ,  $0.06 \pm 6\%$ ,  $\times 10^{-17}$  mol for masses 40, 39, 38, 37, and 36, respectively.

Plateau ages are calculated for the indicated steps and represents the inverse variance weighted mean age and the error is the square root of the sum of  $1/\sigma^2$  values. The error is also multiplied by the square root of the MSWD for MSWD > 1. J-error (0.03%) is included for the plateau age error and all errors are reported at  $2\sigma$ .

#### 4. Results

Hematite <sup>4</sup>He concentrations range from 1.39 to  $17.1 \times 10^{-10}\,\text{ml/mg}$  and the multiple aliquots usually differ by < 3% in each sample (Table 1). <sup>238</sup>U concentrations range from 4.6 to  $66.1 \times 10^{11}\,\text{atoms/mg}$  and <sup>232</sup>Th concentrations range from 0.1 to  $13.7 \times 10^{11}\,\text{atoms/mg}$ . Within-sample U concentrations vary by 3–5%. Variations for <sup>232</sup>Th are larger, though the low concentrations and low production ratio of <sup>4</sup>He relative to U mean that this has a rather minor effect on ages. U/Th ratios of the individual aliquots range from 0.8 to 19.7 (Table 1). The within-sample consistency of U/Th ratios, and the lack of variation in He, U and Th concentration in multiple aliquots suggest that the crushed hematite has been adequately homogenized.

(U + Th)/He ages calculated using the average He, U and Th concentrations for each sample range from 5.36 to 5.64 Ma (Fig. 3; Table 1). The uncertainty of the He ages calculated using the standard deviation of the average He and U (+Th) concentrations, ranges from 2 to 6%. Uncertainties in the average He and U (+Th) concentrations make approximately equal contributions to the age uncertainty.

Two of the three adularia samples (Italy-1 and 3) yield essentially flat  $^{40}$ Ar/ $^{39}$ Ar age spectra giving plateau ages of 5.575  $\pm$  0.008 and 5.583  $\pm$  0.013 Ma (Fig. 4a, c; Table 2). Significant parts of these spectra that yield > 50% of the total  $^{39}$ Ar released have steps concordant at 2 $\sigma$ . For instance, steps C–K of Italy-1 containing 60% of the spectrum yields an age of 5.568  $\pm$  0.009 Ma and for Italy-3 steps *E*-K (68% of the spectrum) yield an age of 5.574  $\pm$  0.009 Ma. Based on the criteria for Fleck et al. (1977) these ages could be argued to be the preferred ages, but we choose to include the entire data set despite the very minor discordance and the fact that the overall interpretation of the data are not altered by choosing subsets of the data. Several features of the Ar/Ar data indicate that they provide a robust age for adularia

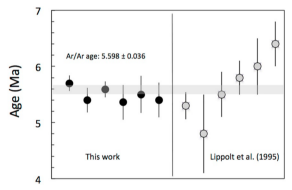
L.-Y. Wu et al. Chemical Geology 504 (2019) 151–157

**Table 1** (U-Th)/He ages and trace element contents for several aliquots of hematite samples from Elba, Italy.

Sample	Aliquot	Mass (mg)	<sup>4</sup> He *	Mean <sup>4</sup> He*	Aliquot	Mass (mg)	$^{238}U^{+}$	<sup>232</sup> Th <sup>+</sup>	eU+	Mean eU <sup>+</sup>	He age (Ma)
Italy-1	I1He1	19.4	1.37	1.36 ± 0.01	I1U1	9.1	0.19	39	0.21	19.9 ± 0.5	5.64 ± 0.11
	I1He2	15.4	1.36		I1U2	11.9	0.19	29	0.20		
	I1He3	10.2	1.35		I1U3	7.0	0.19	23	0.20		
Italy-2	I2He1	14.3	3.93	$3.81 \pm 0.12$	I2U1	7.1	0.56	4	0.57	$58.5 \pm 1.9$	$5.39 \pm 0.23$
	I2He2	13.0	3.81		I2U2	8.4	0.57	28	0.58		
	I2He3	12.0	3.68		I2U3	6.1	0.59	29	0.61		
Italy-3	I3He1	19.0	1.63	$1.61 \pm 0.01$	I3U1	7.2	0.18	188	0.23	$23.8 \pm 1.0$	$5.59 \pm 0.15$
	I3He2	13.0	1.61		I3U2	6.5	0.18	238	0.24		
	I3He3	11.8	1.60		I3U3	6.3	0.19	246	0.25		
Italy-4C	I4CHe1	15.8	2.74	$2.76 \pm 0.02$	I4CU1	8.2	0.37	450	0.47	$56.5 \pm 17$	$5.36 \pm 0.33$
	I4CHe2	12.8	2.75		I4CU2	7.4	0.63	555	0.76		
	I4CHe3	13.7	2.79		I4CU3	8.5	0.35	444	0.46		
Italy-4F	I4FHe1	20.3	3.44	$3.47 \pm 0.09$	I4FU1	8.1	0.41	527	0.53	$52.0 \pm 3.3$	$5.49 \pm 0.35$
	I4FHe2	18.6	3.62		I4FU2	7.7	0.42	522	0.54		
	I4FHe3	15.3	3.44		I4FU3	9.0	0.37	451	0.48		
	I4FHe4	14.4	3.36								
	I4FHe5	14.4	3.50								
M1392	MHe1	11.1	16.76	$16.71 \pm 0.31$	MU1	8.9	2.37	142	2.42	$250 \pm 16$	$5.39 \pm 0.33$
	MHe2	13.2	16.22		MU2	10.2	2.61	205	2.68		
	MHe3	9.7	16.54		MU3	5.6	2.42	102	2.46		
	MHe4	9.9	17.06		MU4	5.4	2.22	78	2.25		
	MHe5	9.6	16.78		MU5	4.3	2.60	0	2.56		
					MU6	4.7	2.60	106	2.64		

<sup>\*</sup> He concentrations are reported in  $10^{-10}$  ml/mg.

<sup>&</sup>lt;sup>+</sup> U concentrations and mean eU+ are reported in ppm, and Th concentrations are ppb.



**Fig. 3.** Summary diagram of (U + Th)/He ages of hematite from Rio Marina mine, Elba; this work (dark circles) and Lippolt et al. (1995) (light circles). The average of the three new  $^{40} \rm{Ar}/^{39} \rm{Ar}$  ages of cogenetic adularia (5.598  $\pm~0.036$  Ma) is shown by the grey bar. The multi-aliquot hematite ages are more precise than those of Lippolt et al. (1995) and they overlap within uncertainty the adularia crystallisation age.

deposition. The nearly flat age spectra indicate simple argon systematics and suggest that they record no significant Ar-loss during protracted cooling. Additionally, the nearly undetectable <sup>37</sup>Ar signal indicates that the samples are essentially devoid of Ca and the 39Ar intensity coupled with sample weight and J-factor indicate K2O contents of > 16 wt%. Both are indicative of pure adularia. Microprobe analyses also confirm that the samples are pure adularia (Fig. 2H). In contrast to Italy-1 and Italy-3, Italy-2 has a more complex 40Ar/39Ar spectrum with initially old ages stepping down to a minimum segment that has a weighted mean age of 5.64  $\pm$  0.03 (Fig. 4b). We suggest that this sample maybe contaminated with excess <sup>40</sup>Ar and the age can be treated as a maximum. There is no simple isochron array, however initially old steps in the age spectrum correlate to lower radiogenic yields that could support and excess argon component. The lack of a well-defined isochron likely suggests that both excess argon component and an atmospheric component are not thermally distinct and thus record mixing between multiple sources with different initial <sup>40</sup>Ar/<sup>36</sup>Ar ratios (cf. Heizler and Harrison, 1988).

Maineri et al. (2003) dated a magmatic muscovite, a hydrothermal sericite and a partly chloritized biotite from the region and obtained overall complex results. They concluded that complexity of the muscovite and sericite was related to mixing of variable amounts of the two

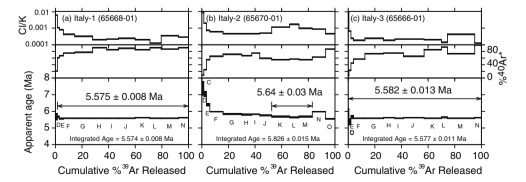


Fig. 4. <sup>40</sup>Ar/<sup>39</sup>Ar age spectra for three adularia samples from hematite mineralisation at Rio Marina, Elba. The spectra show the apparent ages, %<sup>40</sup>Ar\* and Cl/K ratios measured during each heating step.

L.-Y. Wu et al. Chemical Geology 504 (2019) 151-157

Table 2 <sup>40</sup>Ar/<sup>39</sup>Ar data of adularia cogenetic with hematite from Elba, Italy.

ID	Power	$^{40}\mathrm{Ar}/^{39}\mathrm{Ar}$	$^{38}$ Ar/ $^{39}$ Ar	$^{37}\mathrm{Ar}/^{39}\mathrm{Ar}$	$^{36}$ Ar/ $^{39}$ Ar	$^{39}Ar_{K}$	K/Ca		Cl/K	<sup>40</sup> Ar*	<sup>39</sup> Ar	Age	$\pm~1\sigma$
	(Watts)				(×10 <sup>-3</sup> )	(×10 <sup>-15</sup> mol)				(%)	(%)	(Ma)	(Ma
Italy-1	, Adularia, 1.1	4 mg, J = 0.00	)28246 ± 0.0	1%, IC = 1.00	859 ± 0.0017725	, NM-290F, Lab#	= 65,668-	-01, Arg	gus VI				
C	0.3	5.591	0.0416	0.0004	15.19	2.08		1339	0.0059	19.6	1.5	5.654	0.09
D	0.4	1.830	0.0180	-0.0027	2.475	2.68	-		0.00109	59.8	3.5	5.627	0.03
E	0.4	1.613	0.0167	0.0012	1.799	3.81	4	411.6	0.00082	66.9	6.2	5.538	0.02
F	0.5	1.436	0.0148	0.0006	1.188	11.6	7	785.7	0.00042	75.4	14.7	5.556	0.01
G	0.6	1.399	0.0143	0.0011	1.053	18.3	4	452.4	0.00033	77.6	28.0	5.573	0.00
Н	0.7	1.209	0.0136	0.0008	0.4058	14.8	$\epsilon$	524.3	0.00019	90.0	38.8	5.579	0.00
ī	0.7	1.268	0.0138	0.0006	0.6189	11.9		307.2	0.00023	85.5	47.5	5.559	0.00
J	0.8	1.220	0.0138	0.0001	0.4552	17.5		6951	0.00023	88.9	60.3	5.559	0.00
K	0.9	1.291	0.0137	0.0002	0.6772	14.9		2479	0.0002	84.4	71.1	5.587	0.00
L	1.0	1.214	0.0137	-0.0003	0.4176	11.7	_	21/	0.00012	89.8	79.6	5.585	0.00
M	1.2	1.256	0.0133	0.0005	0.5671	17.7		1031	0.00012	86.6	92.5	5.572	0.00
N	2.0	1.202	0.0143	0.0003	0.3720	10.28		2290	0.00033	90.8	100.0	5.593	0.00
		1.202	0.0141		0.3720			2290	0.0003	90.8			
	ated age ± 2σ			n = 12		137.2					$K_2O = 16.37\%$	5.574	0.00
	u ± 2σ on ± 2σ	steps C-N steps C-N		n = 12 n = 12	MSWD = 2.61 $MSWD = 3.06$	137.2	<sup>40</sup> Ar/ <sup>36</sup>	5Ar=		295.5	100.0	5.575 5.575	0.00
		•	020247 ± 0.0		859 ± 0.0017725	NM 200E Lob# .	,		nio VI	2,010	_ 2.0	0.070	0.00
C	0.3	1 mg, 5 = 0.00 15.87	0.0591	0.0016	48.85	, NWI-290F, Lab# - 1.76		-01, A18 322.8	0.00844	9.0	1.8	7.36	0.2
D	0.3	5.397	0.0391	0.0010	13.64	1.83		415.7	0.00209	25.2	3.7	7.006	0.08
E	0.4	2.286	0.0243	0.0012	3.531	2.22		132.0	0.00209	54.2	6.0	6.369	0.04
F	0.5	1.818	0.0157	0.0013	2.219	9.71		383.6	0.00058	63.8	16.1	5.957	0.01
G	0.6	1.601	0.0150	0.0008	1.574	13.4		557.2	0.00046	70.8	30.0	5.820	0.01
H	0.7	1.663	0.0151	0.0009	1.816	7.39		579.5	0.00047	67.6	37.7	5.772	0.01
I	0.7	1.536	0.0150	0.0008	1.367	5.79		544.1	0.00047	73.6	43.7	5.800	0.01
J	0.8	1.875	0.0154	0.0001	2.559	8.71		3771	0.0005	59.5	52.7	5.733	0.01
K	0.9	1.984	0.0181	0.0000	2.979	12.9		2,183	0.0011	55.5	66.1	5.656	0.01
L	1.0	1.956	0.0205	-0.0001	2.916	7.09	-		0.00164	55.8	73.4	5.606	0.02
M	1.2	2.027	0.0176	0.0006	3.123	9.27	8	876.5	0.0010	54.3	83.1	5.656	0.01
N	2.0	1.840	0.0169	0.0006	2.288	9.86	ç	913.7	0.00085	63.1	93.3	5.962	0.01
O	4.0	1.257	0.0149	-0.0009	0.5975	6.47	-		0.00046	85.9	100.0	5.532	0.01
Integrated age ± 2σ			n = 13		96.4					$K_2O = 16.18\%$	5.826	0.01	
Platea	u ± 2σ	steps K-M		n = 3	MSWD = 2.21	29.229					30.3	5.644	0.03
		steps K-M		n = 3	MSWD = 2.13		$^{40}$ Ar/ $^{36}$	$^{6}Ar =$		333 ±	56	5.06	0.8
Italy-3	, Adularia, 1.0	5  mg,  J = 0.00	)28255 ± 0.0	2%, IC = 1.00	859 ± 0.0017725	, NM-290F, Lab# :	= 65,666-	-01, Arg	gus VI				
C	0.3	17.92	0.0631	0.0025	56.92	1.090	2	200.9	0.0090	6.1	0.9	5.65	0.2
D	0.4	4.833	0.0181	0.0002	12.83	1.62		2761	0.00068	21.4	2.2	5.343	0.09
E	0.4	2.438	0.0147	-0.0007	4.518	2.64	_		0.00025	45.0	4.3	5.652	0.04
F	0.5	1.883	0.0140	0.0006	2.703	10.58		326.9	0.00018	57.4	12.8	5.554	0.01
G	0.6	1.507	0.0141	0.0002	1.429	17.2		3028	0.00010	71.8	26.7	5.554	0.00
Н	0.7	1.448	0.0141	0.0002	1.210	10.84		3020	0.00024	75.2	35.4	5.586	0.00
I	0.7	1.458	0.0137	-0.0001	1.261	7.68	_		0.00013	74.3	41.6	5.560	0.01
J	0.7	1.694	0.0139	0.0001	2.039	19.7	_		0.00021	64.2	57.4	5.589	0.01
K	0.8	1.284				16.0		1097	0.00023	84.7	70.3	5.579	0.01
	1.0		0.0137	0.0005	0.6584	5.40							
L		1.184	0.0134	0.0013	0.2936			394.6	0.00014	92.6	74.6	5.621	0.01
M	1.2	1.504	0.0148	0.0006	1.405	25.3		791.6	0.00041	72.2	95.0	5.579	0.00
N	2.0	1.120	0.0133	0.0015	0.0864	6.20	3	348.9	0.00013	97.7	100.0	5.609	0.01
_	Integrated age $\pm 2\sigma$			n = 12		124.2					$K_2O = 16.07\%$	5.577	0.01
Plateau ± 2σ		steps C-N		n = 12	MSWD = 3.92	124.2	40 00				100.0	5.582	0.01
Isochr	on ± 2σ	steps C-N		n = 12	MSWD = 3.72		$^{40}$ Ar/ $^{36}$	Ar=		294.0	± 1.1	5.592	0.00

#### Notes:

Isotopic ratios corrected for blank, radioactive decay, and mass discrimination, not corrected for interfering reactions.

Errors quoted for individual analyses include analytical error only, without interfering reaction or J uncertainties.

Integrated age calculated by summing isotopic measurements of all steps.

Integrated age error calculated by quadratically combining errors of isotopic measurements of all steps.

Plateau age is inverse-variance-weighted mean of selected steps.

Plateau age error is inverse-variance-weighted mean error (Taylor, 1982) times root MSWD where MSWD > 1.

Plateau error is weighted error of Taylor (1982).

Isochron and error determined using York (1969).

Isotopic abundances after Steiger and Jäger (1977).

X preceding sample ID denotes analyses excluded from plateau age calculations.

i symbol preceding sample ID denotes analyses excluded from isochron age calculations.

- K/Ca not determined as no measureable <sup>37</sup>Ar above blank level.

 $IC = measured {}^{40}Ar/{}^{36}Ar$  of air standard divided by 295.5.

Ages calculated relative to FC-2 Fish Canyon Tuff sanidine interlaboratory standard at 28.201 Ma.

Decay Constant (LambdaK (total)) = 5.463e-10/a.

Correction factors:

 $(^{39} Ar)^{37} Ar)_{Ca} = 0.00073 \ \pm \ 0.000020.$   $(^{36} Ar)^{37} Ar)_{Ca} = 0.0002725 \ \pm \ 0.0000009.$ 

 $(^{38}\text{Ar}/^{39}\text{Ar})_{\text{K}} = 0.012718 \pm 0.00008.$ 

 $(^{40}\text{Ar}/^{39}\text{Ar})_{\text{K}} = 0.0088 \pm 0.0004.$ 

L.-Y. Wu et al. Chemical Geology 504 (2019) 151–157

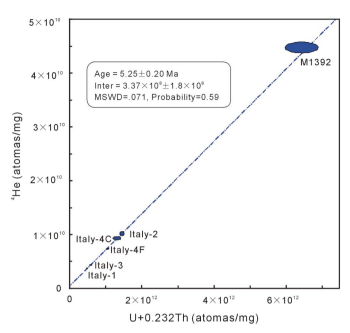


Fig. 5. Isochron plot of (U + 0.232Th) vs.  $^4$ He using the multi-aliquot (U-Th)/He ages of hematite from Rio Marina, Elba.

phases that have discordant ages and variable Cl concentrations. They provided no explanation of the biotite age spectrum discordance and suggested that variable K/Ca values were related to Ca-rich impurities. They utilized the Cl/K (i.e.  $^{38} \mbox{Ar}_{\rm Cl}/^{39} \mbox{Ar}_{\rm K}$ ) data to help constrain the mixing of younger sericite with the older muscovite and concluded that sericite alteration occurred at  $\sim\!6.7$  Ma. Cl/K spectra (Fig. 4) suggest that values are highest (> 0.005) in the initial heating steps and fall to values that vary between 0.0001 and 0.0005. There is no simple correlation between Cl/K and age which further supports variable trapped initial  $^{40} \mbox{Ar}/^{36} \mbox{Ar}$  components to explain the complexity of Italy-2. Therefore, unlike the complex results of Maineri et al. (2003), the adularia data are straightforward (especially Italy-1 and 3) to interpret and indicate hydrothermal precipitation of pure adularia well after an earlier sericite alteration event.

## 5. Discussion: Accuracy and precision of multi-aliquot (U + Th)/He ages

The new (U + Th)/He ages of the Rio Marina hematite samples overlap within uncertainty. They display significantly less variation than the age range determined in the previous study (4.8 to 7.3 Ma; Lippolt et al., 1995) (Fig. 3). This is unlikely to reflect variation in the timing of hematite mineralisation of the Rio Marina deposit that was not sampled in this study as the quartz monzonitic intrusion which drove the hydrothermal system provides an upper limit to the age of mineralisation (5.9  $\pm$  0.2 Ma; Maineri et al., 2003). The uncertainty on these hematite ages (3–5%) is an improvement on the 10–30% age range that can result from multiple single aliquot measurements even after rejection of data affected by over-heating or inclusions (e.g. Cooper et al., 2016). This conclusion has, however, to be tempered by the recognition that the multi-aliquot technique requires significantly more sample material and analytical effort (e.g. column chemistry).

The mean (U + Th)/He age of the three hematite samples with coexisting adularia (5.56  $\pm$  0.15 Ma) overlaps the plateau ages of the adularia with the nearly flat age spectra. This hematite age is slightly younger than the maximum  $^{40}{\rm Ar}/^{39}{\rm Ar}$  age determined for the Italy-2 adularia, however Italy-2 likely contains excess argon based on its overall saddle-shaped age spectrum.

The mean (U + Th)/He age of all five Rio Marina hematite samples  $(5.53 \pm 0.14 \, \text{Ma})$  is indistinguishable from the age of the three that

contain coexisting adularia, consistent with all samples originating from the same hydrothermal event. In this case the data can be used to plot an isochron (Fig. 5). The isochron slope is equivalent to an age of  $5.25\pm0.20\,\mathrm{Ma}$ . The isochron intersects the origin within uncertainty, indicating that the hematite does not contain significant parentless <sup>4</sup>He, such as would be present in hydrothermal fluid inclusions (e.g. Stuart et al., 1995).

The similarity of the multi-aliquot hematite (U + Th)/He ages and Ar/Ar ages is strong evidence that there has been no  $^4\text{He}$  loss since mineralisation that can be distinguished within analytical uncertainty. This implies that diffusive loss during post-mineralisation cooling or reheating during subsequent hydrothermal events, and  $\alpha\text{-ejection}$  to grain boundaries can be ignored in these samples. This is supported by the observation that the coarse and fine fractions of Italy-4 hematite yield ages that are indistinguishable within uncertainty (Table 1).

Dünkel (2002) estimate that the Rio Marina hematite precipitated at 300–320 °C based on fluid inclusion homogenization temperatures in cogenetic quartz. At this temperature hematite is expected to retain no  $^4\text{He}$ . The closure temperature of the Elba hematites likely range from 80 to 200 °C (Lippolt et al., 1993; Bähr et al., 1994; Balout et al., 2017; Farley, 2018). The similarity of the (U + Th)/He and  $^{40}\text{Ar}/^{39}\text{Ar}$  ages of the samples is consistent with rapid cooling to < 50 °C after precipitation and the absence of significant re-heating.

It also implies that recoil of  $\alpha$  particles to hematite grain boundaries does not result in the loss of significant proportion of the  $^4\text{He}$  from the  $10\text{--}30\,\mu\text{m}$  fragments analysed in this study. This may be a consequence of the grain boundaries being volumetrically insignificant, or that they do not form a connected network that promotes the loss of He from intergrown sub- $\mu\text{m}$  hematite crystallites.

#### 6. Conclusion

(U + Th)/He ages have been determined from several fine-grained vein hematite samples from Rio Marina mine, Elba, using a multiple aliquot technique. The low variability in He and U concentrations in 3-5 aliquots from each sample suggests that crushing to  $< 30 \,\mu m$  is adequate to homogenise 5-10 mg of hematite in this case. The data define a coherent isochron consistent with an absence of parentless <sup>4</sup>He (e.g. in hydrothermal fluid inclusions) in the hematite. The new He ages overlap high precision 40Ar/39Ar age of coexisting adularia which implies that diffusion and recoil loss of <sup>4</sup>He from hematite microcrysts are not significant, and that the He ages record the time of mineralisation. The multi-aliquot technique avoids the analytical challenges associated with techniques where He, U and Th are measured on the same aliquot, and demonstrates that the method has the potential to reliably deliver precise and accurate ages for iron oxide precipitation that has not suffered significant post-crystallisation thermal perturbation. The coherence of the hematite-adularia ages, and the apparent homogeneity of He, U and Th (at the sample mass analysed here) suggests that Elba hematite may prove a useful secondary mineral standard analogous to the Durango apatite for (U + Th)/He dating.

The  $^{40}$ Ar/ $^{39}$ Ar ages of the Rio Marina mine adularia are the first high precision ages of the Elba iron oxide mineralisation. The ages overlap the K–Ar age for the Porto Azzurro monzonite (5.9  $\pm$  0.2 Ma; Maineri et al., 2003) confirming the genetic link. Further they provide a baseline for future chronology studies aimed at determining the history of iron mineralisation on Elba.

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