Cyclic magmatic-hydrothermal evolution in porphyry systems: High-precision U-Pb and Re-Os geochronology constraints from the Tibetan Qulong porphyry Cu-Mo deposit

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Abstract

We present high-precision chemical abrasion isotope dilution thermal ionization mass spectrometry (CA-ID-TIMS) U-Pb zircon and isotope dilution negative thermal ionization mass spectrometry (ID-N-TIMS) Re-Os molybdenite geochronology of the world-class Tibetan Qulong porphyry Cu-Mo deposit. The data is used to constrain the timing, duration and yield implications of the ore-forming processes. The U-Pb data suggest that the pre-ore Rongmucuola pluton crystalized at $17.142 \pm 0.014/0.014/0.023$ Ma (uncertainties presented as analytical / + tracer / + decay constant uncertainties), with emplacements of the syn-ore P porphyry and post-ore quartz diorite occurring at $16.009 \pm 0.016/0.017/0.024$ and $15.166 \pm 0.010/0.011/0.020$ Ma, respectively. The Re-Os analysis of multiple independent molybdenite separations from single molybdenite-bearing quartz veins yields sub-per-mil-level analytical precision (<1 ‰), which is comparable with that of modern CA-ID-TIMS U-Pb zircon geochronology. The new Re-Os data indicate that the majority of the metals at Qulong were deposited over a minimum duration of $266 \pm 13$ thousand years (kyr) between $16.126 \pm 0.008/0.060/0.077$ and $15.860 \pm 0.010/0.058/0.075$ Ma, with the main phase of mineralization being broadly synchronous with the emplacement of the P porphyry. However, our Re-Os data of molybdenite hosted within the Rongmucuola pluton imply that a portion of mineralization also predated the P porphyry, and suggest that the P porphyry is an intermineral porphyry stock, although mineralization cut by P porphyry has not been previously documented or observed in this study. Correlating the Re-Os ages with vein types (A-B-D veins) demonstrates that the mineralization process was cyclical with the presence of at least 3 short-lived ($38 \pm 11$ to $59 \pm 10$ kyr) mineralization pulses between $16.126 \pm 0.008$ and $16.050 \pm 0.005$ Ma, $16.040 \pm 0.007$ and $15.981 \pm 0.007$ Ma, and $\sim 15.981 \pm 0.007$ and $15.860 \pm 0.010$ Ma. Coupling the Re-Os molybdenite ages and quartz (co-precipitated with the dated molybdenite) fluid inclusion data suggests that the cooling history was also cyclic,
and implies a rapid cooling rate during the entire mineralization process (0.55 ± 0.11 °C/kyr),
with much faster cooling rates (1.19 ± 0.82 - 1.27 ± 0.53 °C/kyr) for the individual
mineralization pulses. The cyclic and rapid cooling process requires additional cooling
mechanism rather than the inefficient conduction, which we attribute to meteoric water
circulation.

The presence of mineralization predating the intermineral P porphyry stock and the
absence of evidence of an early porphyry stock at Qulong suggest that mineralization
potentially can take place without contemporaneous magmatism at mineralization levels. As a
result, dating magmatic events may not necessarily bracket the entire mineralization duration
of a porphyry system. This highlights the importance of dating ore minerals to fully reveal the
magma-hydrothermal process. In addition, the absence of contemporaneous magmatism
during mineralization have broad implications for the classification of porphyry copper
deposits and mineral exploration. The timescales of mineralization cycles constrained here
via direct dating of ore minerals (tens of kyr) are comparable with those recently proposed
through high-precision U-Pb zircon dating, diffusion modelling and numerical simulation.
We propose that the cyclic mineralization pulses are linked with the periodic release of
volatile from the lower crustal magma chamber, and such cyclic processes are common for
porphyry copper systems worldwide. As such episodic/cyclic metal-enrichment potentially is
one of the controlling factors of porphyry copper ore formation, and are therefore key to
differentiate the formation of economic and sub-economic porphyry deposits.

Finally, direct comparison of molybdenite Re-Os dates from different labs and with
the zircon U-Pb system needs to account for the much larger uncertainties from tracer
calibration and decay constants, respectively, which therefore loses the necessary resolution
to investigate the ore-forming process at the kyr level. As a result, calibration between the
two chronometers and using shared tracer solutions and a transparent data reduction platform within the community is required.
1 Introduction

Metals forming porphyry copper deposits are derived from the lower crust and transported to the shallow mineralization levels by magmatic fluids via porphyry stocks that act as conduits (Cooke et al., 2014; Richards, 2011; Sillitoe, 2010). Although this model is widely accepted, there is a paucity of detailed, precise and robust timeframes of igneous and hydrothermal systems to test this hypothesis. For example, at what level are mineralization and magmatism contemporaneous? Moreover, the duration of the porphyry mineralization process is poorly constrained, with studies suggesting orders of magnitude variation, from tens of thousands years to several millions of years (Myr) (Chiaradia et al., 2014).

The timescales of ore formation in porphyry copper deposits can be constrained through the dating of pre- and post-ore intrusions. Traditionally, in-situ zircon U-Pb dates are used to establish the timeframe of porphyry stocks and constrain the duration of ore formation (Deckart et al., 2012; Sillitoe and Mortensen, 2010). However, limited by the precision (~2%) of in-situ U-Pb zircon analysis (Klötzli et al., 2009; Li et al., 2015; Schaltegger et al., 2015; Schoene, 2014), the conclusions of these studies are controversial (von Quadt et al., 2011). High-precision (per mil level) dating techniques, e.g., CA-ID-TIMS zircon U-Pb geochronology, make it possible to refine the timeframe and timescales of porphyry copper systems (von Quadt et al., 2011). Recent high-precision zircon U-Pb geochronology case studies indicate that the lifetime of porphyry Cu deposits varies from tens to hundreds of kyr, and by inference suggest the presence of multiple magmatic-hydrothermal episodes (Buret et al., 2016; Chelle-Michou et al., 2015; Tapster et al., 2016; von Quadt et al., 2011). The latter is supported by numerical simulation and diffusion modelling, which suggest that the ore-forming event involves multiple short-lived (several tens of kyr) mineralization pulses (Cathles, 1977; Chelle-Michou et al., 2017; Mercer et al., 2015; Weis, 2015; Weis et al., 2012). However, these approaches (U-Pb dating, numerical
simulation and diffusion modelling) are either based on dating porphyry stocks, which do not
necessarily bracket the entire mineralization duration, and/or age constraints of silicate
minerals, which may not co-precipitate with ore-minerals, or numerical models that do not
fully reflect the complexity of a certain porphyry system.

The most straightforward approach to constrain the timing and duration of ore-
forming events is directly dating the ore minerals. The ubiquitous distribution of molybdenite
in porphyry Cu-Mo deposits and advances in molybdenite Re-Os geochronology permit
precise dating of the ore-forming event(s) directly (Spencer et al., 2015; Stein, 2014). High-
precision molybdenite Re-Os dating of the El Salvador and El Teniente porphyry Cu-Mo
deposits derived a mineralization duration of 0.6 Myr (Zimmerman et al., 2014), and
presence of multiple short-lived (<100 kyr) mineralization pulses over 1.7 Myr (Spencer et al.,
2015), respectively. As each complete mineralization pulse comprises petrographically
defined veinlets, which are termed as A (earliest), B, D (latest) veins (Gustafson and Hunt,
1975; Sillitoe, 2010), multi-pulsed mineralization events will result in a complex evolution
history (i.e., A veins that are younger than D veins). This has been supported by high-
precision Re-Os dating of the Los Pelambres Cu–(Mo) deposit, which shows that D veins are
~1 Myr older than B veins (Stein, 2014).

As discussed above, multiple magmatic-hydrothermal/mineralization pulses are
evident for deposits with prolonged formation intervals (e.g., several Myr). However, for
deposits with shorter formation intervals (e.g., several to tens to hundreds of kyr), which, in
some cases, are at the same magnitude as the precision of the dating method, there is an
enhanced need to apply high-precision dating to understand the chronology and duration of
the ore-forming process. Moreover, due to a lack of deposit-wide crosscutting relationships, it
is not clear yet whether these short-lived systems are formed via a single protracted pulse or
through multiple short-lived intermittent pulses. In addition, for deposits incrementally
formed by several successive mineralization pulses, the timescale of individual pulses remains unknown. The answers for these questions will aid our understanding of metal deposition processes in porphyry copper deposits and shape exploration strategy, e.g., are multiple mineralization pulses necessary to form giant deposits, or is a prolonged mineralization duration essential for economic deposits? Moreover, if multiple magmatic-mineralization events are common in porphyry copper deposits, what is the cooling history of the system? In addition, multiple magmatic-hydrothermal/mineralization events predict thermal resetting for chronometers with low closure temperatures, which need to be considered when interpreting thermochronology dates (e.g., zircon U-Th-He, apatite AFT and mica $^{40}$Ar-$^{40}$Ar).

In this paper, we examine the temporal models (e.g., the temporal relationship between magmatism and mineralization, and the ore-forming processes) of porphyry copper deposits via constraining the ore-forming process of the well-studied Qulong porphyry Cu-Mo deposit by employing CA-ID-TIMS U–Pb zircon and ID-N-TIMS Re-Os molybdenite geochronology with analytical precisions of ~1‰. Qulong has only one mineralization centre with one main porphyry stock, which makes the deposit a relatively simple and ideal candidate for the research proposed here. The data are used to examine the temporal relationship between magmatism and mineralization, and to propose that one of the main controls of economic porphyry copper systems is episodic/cyclic mineralization events.

2 Geological background

The Qulong porphyry Cu-Mo deposit is located in the eastern region of the Gangdese magmatic arc (Fig. 1A), which extends along the southern margin of the Lhasa terrane. The Lhasa terrane records the subduction of the Neo-Tethyan oceanic lithosphere and subsequent India-Asia collision (Zhang et al., 2014; Zhu et al., 2015). The eastern part of the Gangdese
magmatic arc is termed the Gangdese Porphyry Copper Belt which contains >20 porphyry
copper systems that formed in a collisional or post-collisional (India-Asian) tectonic setting
during the Cenozoic (Hou et al., 2009; Wang et al., 2015; Zheng et al., 2015), with the
magnas generated by partial melting of a thickened basaltic lower-crust (Hou et al., 2009;
Richards, 2015; Wang et al., 2014). Of these porphyry copper systems, eight of them have Cu
reserves in excess of 0.5 million tonnes (Mt), with Qulong being the largest both in this belt
and China, which hosts ~11 Mt Cu and >0.5 Mt Mo (Hu et al., 2015; Li et al., 2017b; Yang et
al., 2009; Zhao et al., 2016).

2.1 Miocene intrusive units at Qulong

The Mid-Miocene Qulong deposit is spatially associated with the Rongmucuola
pluton, aplite, P porphyry, X porphyry, two stages of breccia, and quartz diorite units (Fig. 1).
The P porphyry and orebody show a spatial association with a north-south striking normal
fault system (Fig. 1). The geology of the Qulong system has been described in detail by many
previous studies (Hu et al., 2015; Li et al., 2017b; Xiao et al., 2012; Yang et al., 2009; Zhao
et al., 2016; Zheng et al., 2004), which we summarise and discuss below. The relative
chronology of these units defined by crosscutting relationships is presented in Figure 2A (Hu
et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016).

The Mid-Miocene Rongmucuola pluton is the predominant host lithology to the
mineralization of Qulong, which intruded into the Jurassic volcanic (Yeba Formation) and
intrusive rocks (dacite-rhyolite porphyry). The pluton has a surface exposure of ~8 km² and a
deepth of >2 km as constrained by drilling (Fig. 1B-C). The Rongmucuola pluton’s
composition varies gradually from granodiorite to biotite monzogranite (Fig. 1B) from east to
west (Yang et al., 2009; Zhao et al., 2016). Despite this compositional variation, the entire
pluton possesses a similar mineralogy (plagioclase, K-feldspar, quartz, amphibole and biotite,
Fig. 3A), however mineralization is only present in the western Rongmucuola pluton (Fig. 1B). The western Rongmucuola pluton is cut by small aplite dikes, the P porphyry, the X porphyry, and a breccia pipe. The aplite hosted by the Rongmucuola pluton has limited distribution in the drill cores and occurs as several to tens of centimetres wide discontinuous fracture-controlled dikes (Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The aplite is characterized by intergrowths of fine-grained (≈1 mm) anhedral alkali feldspar and quartz with disseminated magnetite and pyrite (Li et al., 2017b; Yang et al., 2009).

The P porphyry, which has an exposure diameter of ≈200 m and a depth of >1.8 km as indicated by drilling, was emplaced into the centre of the western Rongmucuola pluton (Fig. 1B-C). No observed (this study) or documented crosscutting relationships between the aplite and the P porphyry are known (Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The P porphyry is monzogranitic in composition (plagioclase, quartz, K-feldspar and amphibole and biotite, Fig. 3B) and occurs in the centre (Fig. 1B-C) of the deposit (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The X porphyry crosscuts both the western Rongmucuola pluton (Fig. 1C) and the P porphyry (Zhao et al., 2016), and has a limited distribution only being present at shallow levels (<200 m) as discontinuous irregular dikes with variable thickness from a few centimetres to 0.5 m (Li et al., 2017b; Yang et al., 2009). The X porphyry possesses a similar composition and texture with that of the P porphyry, with the exception to its biotite content (3-5% in the X porphyry vs <3% in the P porphyry, Yang et al., 2009). A magmatic-hydrothermal breccia pipe with a diameter of ≈100 m (Fig. 1C) and unknown depth (>500 m) is observed to only crosscut the western Rongmucuola pluton (Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The breccia pipe contains two stages of breccia formation, with the first stage breccia comprising clasts (<2 cm) of the mineralized and hydrothermally altered Rongmucuola pluton and the X porphyry, which is crosscut by the second stage breccia, which is characterized by fragments (1 - 10 cm)
of mineralized and hydrothermally altered Rongmucuola pluton (Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). A cement of a monzogranitic composition, plus mineralized alteration assemblages (e.g., anhydrite, quartz, feldspar, biotite, pyrite can chalcopyrite) is characteristic to both stages of breccia (Yang et al., 2009).

The cessation of magmatism at Qulong is marked by a quartz diorite (plagioclase, quartz and hornblende, Fig. 3C), which intrudes the Rongmucuola pluton (Fig. 1C) as discontinuous dikes with a thickness of 2 - 6 m (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The quartz diorite exhibits a low degree of alteration as evidenced by the plagioclase phenocrysts. The alteration assemblage consists of calcite-chlorite-sericite and clay minerals (Li et al., 2017b; Yang et al., 2015).

2.2 Alteration and mineralization at Qulong

Mineralization at Qulong is predominately (>80 %) hosted by the western Rongmucuola pluton. Drill core logging indicates that the Cu-Mo metals are directly associated with potassic stage alteration assemblages (80 %), though propylitic and phyllic alteration assemblages also contain Cu-Mo mineralization (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The potassic stage alteration is ubiquitous in the western Rongmucuola pluton with the most intensive alteration assemblages being spatially associated with the P porphyry and the N-S trending fault system (Li et al., 2017b; Yang et al., 2009). Propylitic alteration assemblages predominately occur in the western Rongmucuola pluton, and overprint potassic alteration assemblages (Yang et al., 2009). Both potassic and propylitic stage alterations are overprinted by pervasive phyllic stage alteration assemblages in the western Rongmucuola pluton (Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). In the first order, the grades of Cu-Mo metals exhibit concentrically zoned patterns (Fig. 1D-E) around the P porphyry and the N-S trending fault (Yang et al., 2009), although the ore with
the highest grade does decouple from the P porphyry. The decoupling is most evident for the
grade of Mo in the orebody, with the highest grade of ore only occurs on the eastern side of
the P porphyry (Fig. 1E). Considering the small size, limited and discontinuous distribution
of the aplite and the X porphyry, it is currently accepted that the P porphyry is the main fluid
conduit at Qulong (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). The
ore minerals at Qulong are chalcopyrite and molybdenite with trace amounts of bornite. The
ore minerals are predominately hosted by quartz vein assemblages in the western
Rongmucuola pluton (Li et al., 2017b; Yang et al., 2009), and to a lesser extent by the
Jurassic Yeba Formation. Except for very minor disseminated molybdenite clusters (<1 mm)
or quartz veins with very minor molybdenite (equivalent to <0.5 mg), which also exhibit
intensive hydrothermal overprint by late stage propylitic and phyllic alteration, no
molybdenite-bearing veins with limited overprint and appreciable molybdenite (≥10 mg)
have been observed in the P porphyry during the course of study. Further, according to
previous drill core logging (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al.,
2016) and observation of this study, the P porphyry does not crosscut any mineralization. The
quartz veins are classified as A, B and D veins (Fig. 4) based on previous studies (Li et al.,
2017b; Yang et al., 2009) and by following the vein terminology/classification of Gustafson
and Hunt (1975) and Sillitoe (2010). Representative A, B and D veins observed at Qulong are
presented in Figure 4. The A veins are characterized by 0.3 - 0.8 cm wide discontinuous
granular quartz veinlets with discontinuous narrow (< 3 mm) K-feldspar selvages, which host
disseminated chalcopyrite and molybdenite (Fig. 4A). The B veins are represented by veinlets
(0.4 - 2 cm wide) comprising crystalline quartz with a crack-seal texture, and possess
irregularly distributed K-feldspar selvage, and predominantly host molybdenite and
chalcopyrite along the vein margin (Fig. 4B). The D quartz veins (0.5 - 6 cm) host euhedral
pyrite, with sericite and anhydrite (0.1 and 1.5 cm) selvages (Fig. 4D), and minor disseminated chalcopyrite and molybdenite.

The relative timing relationships between different vein types, e.g., barren A vein cut by later mineralized A vein and/or B vein, have been previously documented (Li et al., 2017b; Yang et al., 2009) and also observed in this study (Fig. 4E-F). However, importantly to avoid any potential mixing of different generations of mineralization and therefore to yield robust molybdenite Re-Os geochronology, this study focused on veins without cross-cutting relationships by later veins and that show the least evidence of overprinting (e.g., Fig. 4A, B, D).

2.3 Previous Geochronology

The Qulong deposit has been studied extensively with the principal goal to establish the timing of magmatism, mineralization and timeframe of cooling (Hou et al., 2004; Hu et al., 2015; Li et al., 2017b; Wang et al., 2006; Yang et al., 2009; Yang et al., 2015; Zhao et al., 2016; Zheng et al., 2004) and references therein). These studies established the timeframe of the Qulong porphyry system (Fig. 2A) via LA-ICPMS and SHRIMP/SIMS (U-Pb zircon, appendix Table A1) and ID-ICPMS (Re-Os molybdenite, appendix Table A2). Samples from the western Rongmucuola pluton dated by in-situ zircon U-Pb yield variable weighted mean $^{206}$Pb/$^{238}$U dates between 17.6 ± 0.4 Ma and 16.2 ± 0.3 Ma, with dates from the P porphyry varying from 17.58 ± 0.74 Ma to 16.2 ± 0.4 Ma. The X porphyry has a mean $^{206}$Pb/$^{238}$U date of 15.9 ± 0.4 Ma and post-ore quartz diorite yield mean $^{206}$Pb/$^{238}$U dates of 15.7 ± 0.2 and 15.3 ± 0.3 Ma. Molybdenite hosted by the Rongmucuola pluton yield variable dates ranging from 16.85 ± 0.19 to 15.36 ± 0.21 Ma. These prior studies provide a basic framework for the formation time of Qulong. However, the emplacement ages of the intrusions constrained by in-situ U-Pb zircon techniques and the mineralization ages determined by ID-ICP-MS...
molybdenite Re-Os methodology have significant uncertainties (1.3 - 4.2 % and 1.1 - 12.2 %, respectively), which equate to absolute uncertainties of 0.2 - 0.7 and 0.2 - 1.9 Myr, respectively, for the U-Pb and Re-Os dates of the ~16 Ma porphyry system. In order to evaluate the dates of the same intrusion from different studies, the mean and 2 standard deviation of zircon U-Pb $^{206}$Pb/$^{238}$U data from different studies with different dating techniques are presented in Figure 2B. The reason for using the mean and 2 standard deviation, rather than weighted mean, is that dates from different studies are not equivalent and cannot be regarded as one population. The significant variations demonstrated in Figure 2B either reflect the complexity of ages (e.g., complex magmatic evolution history) the zircon grains record (Chelle-Michou et al., 2014), and/or unaccounted analytical bias inherited in micro-beam analysis (Li et al., 2015). As demonstrated previously (von Quadt et al., 2011), the uncertainties of these data are mostly greater than the timescales proposed for the ore formation processes (Chiaradia et al., 2014). Therefore, these dates cannot be used to precisely define the durations of the magmatic and mineralization event(s) and further develop models for ore formation (Schaltegger et al., 2015).

3 Samples and analytical methods

To provide robust and precise time constraints for the Qulong porphyry Cu-Mo system, high-precision CA-ID-TIMS zircon U-Pb geochronology and ID-N-TIMS molybdenite Re-Os dating are employed. Representative samples with limited alteration of the western Rongmucuola pluton, P porphyry, and the quartz diorite (Fig. 3) were collected from drill core to conduct U-Pb zircon dating. The X porphyry has been exhausted by previous studies in the available drill core we studied, as such the X porphyry was not available for CA-ID-TIMS zircon U-Pb analysis for this study. A previous attempt to date the
aplite by SIMS zircon U-Pb analysis demonstrated that the aplite only contains inherited
grains (Li et al., 2017b), and in this case the aplite is not further investigated by CA-ID-TIMS.
The sample from the Rongmucuola pluton (1605-296, Fig. 3A) used for U-Pb dating
exhibits a low degree of hydrothermal alteration and mineralization, as evidenced by the rims
of the biotite and plagioclase, which are altered to chlorite and sericite, with the groundmass
possessing minor disseminated pyrite. The P porphyry sample (001-550, Fig. 3B) used for U-
Pb dating is extensively hydrothermally altered and overprinted by multiple stages of
alteration assemblages. The quartz phenocrysts exhibit an embayed morphology and K-
feldspar shows partial replacement by sericite. In addition, the groundmass displays different
degrees of hypogene alteration. For example, plagioclase and biotite either exhibit complete
or partial destruction with alteration to sericite/clay and chlorite, respectively. The quartz
diorite sample (1605-81, Fig. 3C) used for U-Pb dating only exhibits minor alteration, as
represented by the rims of the plagioclase phenocrysts which are altered to calcite-chlorite-
sericite and clay minerals.
To yield robust molybdenite Re-Os geochronology, e.g., avoid any potential
disturbance of Re-Os by overprinting and mixing different stages of mineralization, veins
without intersections and exhibiting the least evidence of overprint (e.g., Fig. 4A, B, D) were
selected for molybdenite Re-Os geochronology. Three A veins, 7 B veins and 3 D veins, were
selected to conduct high precision ID-N-TIMS Re-Os molybdenite geochronology. All of the
veins used for Re-Os molybdenite geochronology are hosted by the western Rongmucuola
pluton, with the exception of one vein (sample 313-145), which is hosted by an aplite dike.
The dated molybdenite-bearing veins are identical to those used previously for a fluid
inclusion study (Li et al., 2017b), and therefore permit coupling vein formation ages and
temperatures (see discussion). The sample number of studied igneous rocks and molybdenite-
bearing veins is made up by the drill core number and sample depth in meters. Locations of drill are marked in Figure 1B.

3.1 CA-ID-TIMS zircon U-Pb

The methods for U-Pb geochronology are the same as previously documented by (Tapster et al., 2016). Mineral separation was carried out at the NERC Isotope Geosciences laboratory (NIGL), British Geological Survey (BGS), Nottingham, UK. Zircons were isolated from ~1.5 kg drill core samples. After washing, samples were jaw crushed, disc milled and sieved (<355 µm mesh). Heavy minerals were concentrated using a Rogers table, then a Frantz electromagnetic separator and finally by standard heavy liquid (diiodomethane) concentration. Zircon grains from the concentrated materials were then handpicked under a binocular microscope and mounted in epoxy, and then polished with a 0.25 µm diamond paste and finished at near-equatorial sections to yield the best internal exposure of the zircon grains.

The mounts were coated with carbon for CL imaging (Li et al., 2017a), and zircon grains with least inclusions and no indication of inheritance cores (Fig. 3) were removed from the mounts and then individually thermally annealed in quartz crucibles at 900 °C for ~60 hours as part of the chemical abrasion process (Mattinson, 2005). The zircons were then individually ultrasonically cleaned in warm 4N HNO₃, and further rinsed with 4N HNO₃ and ultrapure acetone. The chemical abrasion was continued by loading the zircons individually into 300 µl FEP Teflon microcapsules with 200 µl 29 N HF with trace 4N HNO₃. The microcapsules were placed in a self-sealing Parr™ vessel at ~180 °C for 12 hours (Mattinson, 2005). The leachate was removed, with the zircon fractions being rinsed in 4N HNO₃, fluxed in 6N HCl at ~80 °C for several hours, and further rinsed in 4N HNO₃ and ultrapure water. The leached zircons were spiked with the mixed $^{202}$Pb-$^{205}$Pb-$^{233}$U-$^{235}$U EARTHTIME tracer
solutions (ET2535, Condon et al., 2015; McLean et al., 2015) before dissolution with ~120 µl of 29N HF and trace HNO₃ in Parr™ vessels at ~220 °C for 60 hours. Solutions of the equilibrated tracer and dissolved zircon were evaporated and re-dissolved in Parr™ vessels overnight in 200 µl 3N HCl at 180 °C. Uranium and Pb were separated by standard HCl-based anion-exchange chemistry (Krogh, 1973) and evaporated to dryness with 2 µl of 0.05N H₃PO₄.

Uranium and Pb were loaded together on a single zone-refined Re filament in a silica gel-phosphoric acid matrix (Gerstenberger and Haase, 1997). Isotope ratios were measured using a Thermo-Electron Triton Thermal Ionization Mass-Spectrometer. Uranium was measured as an oxide (UO₂) in static Faraday mode on Faraday detectors equipped with 10¹² Ω resistors for intensities greater than 4 mV or peak-hopping mode on a SEM detector for low-U samples. The isobaric interferences are corrected using a ¹⁸O/¹⁶O value of 0.00205 (Condon et al., 2015). Lead isotopic compositions were measured by peak-hopping mode on a secondary electron multiplier detector (SEM). Linearity characterization and dead time corrections (Pb = 24.5 ns; U = 22 ns) for SEM measurements were made from long-term monitoring of the NBS 982, NBS 981 and U 500 standards.

Measured U and Pb isotopic data were processed using the TRIPOLI software package (Bowring et al., 2011). Data reduction, error propagation, data calculation and presentation were conducted using ET_Redux and related algorithms (McLean et al., 2011). Mass bias for Pb isotope measurement was corrected with real-time ²⁰²Pb/²⁰⁵Pb ratios (Condon et al., 2015; McLean et al., 2015). The U mass fractionation was calculated in real-time based on the isotopic composition of the EARTHTIME tracer solutions. Decay constants used here are from Cheng et al. (2000) for ²³⁰Th, Jaffey et al. (1971) for ²³⁸U and ²³⁵U, with the value of ²³⁸U/²³⁵U from Hiess et al. (2012).
All common Pb (Pbc) was attributed to an isotopic composition of the NIGL laboratory blank of $^{206}\text{Pb}^{204}\text{Pb} = 18.099 \pm 3.02 \%$, $^{207}\text{Pb}^{204}\text{Pb} = 15.545 \pm 1.758 \%$ and $^{208}\text{Pb}^{204}\text{Pb} = 37.824 \pm 2.814 \%$ (2σ uncertainties) for the analytical period. For most zircon grains, the radiogenic Pb are high compared with common Pb (with a ratio >5 in most cases), showing that the results are insensitive to the common Pb corrections (Table 1). The U blanks were assigned a value of 0.10 ± 0.02 pg (2σ) based upon total procedural blanks.

The preferential inclusion of $^{238}\text{U}$ and exclusion of $^{230}\text{Th}$ during zircon crystallization (Schärer, 1984), and thus eventual deficit in $^{206}\text{Pb}$ (younger apparent dates), was accounted via Th disequilibrium correction. For corrections, the Th/U values of the bulk samples were used to represent the Th/U value of the melt at the time of zircon crystallization (Schärer, 1984).

The Th/U values of the Rongmucuola pluton, the P porphyry and the quartz diorite are 4.0, 3.6 and 5.0, respectively (Yang et al., 2009). These values were used with an uncertainty of ± 2 (2σ) to correct the initial $^{230}\text{Th}$ disequilibrium. However, the Th/U values of individual zircons may vary due to fractional crystallization, partition coefficients, temperature and oxidation state (Burnham and Berry, 2012; Luo and Ayers, 2009; Rubatto and Hermann, 2007). To account for all these complexities, we further investigated the sensitivity of the $^{230}\text{Th}$ correction by varying the Th/U values within geologically feasible values between 1 and 7 (Fig. 5). As demonstrated by using the youngest group comprising 3 zircon dates from the P porphyry (see below for detailed discussion), the weighted average of the youngest group varied by less than 40 kyr over the range of Th/U values between 2 and 7 (Fig. 5A), and the statistically acceptable MSWD values (e.g., <1.5, Fig. 5B) were maintained throughout. In addition, within uncertainties, the weighted average from our preferred solution (using the bulk sample's Th/U value for correction) overlapped with those corrected from Th/U values of 2 - 7 (Fig. 5A). However, when magmatic Th/U value
approaches unity (Th/U = 1), a significantly younger date was determined. Given that the Th/U values of the Rongmucuola pluton, the P porphyry and the quartz diorite are 4.0, 3.6 and 5.0, respectively (Yang et al., 2009), we consider that correcting for initial $^{230}$Th disequilibrium using a Th/U $\leq 1$ is a geologically unreasonable assumption. Therefore, using Th/U values ($\pm 2$, 2 $\sigma$) of the bulk samples is a reasonable and valid approach.

3.2 ID-NTIMS molybdenite Re-Os

The molybdenite separation and Re-Os dating methods are adopted from previous studies (Lawley and Selby, 2012; Selby and Creaser, 2001). Molybdenite-bearing (0.3 - 3 mm grain size) quartz veins were selected based on previous petrography and fluid inclusion study (Li et al., 2017b), and then cut into several parts depending on molybdenite abundance (Fig. 6). The fractions were then individually crushed by an agate pestle to 10 mesh (~2 mm) and then handpicked under a binocular microscope to remove non-molybdenite-bearing phases and to examine the genetic association between quartz and molybdenite. The pre-purified materials (quartz-bearing molybdenite) of individual separations were loaded into Teflon beakers with 10 ml 32 N HF to dissolve quartz at room temperature overnight (Lawley and Selby, 2012). The HF purification step was repeated until most of the quartz fractions were dissolved, e.g., most of the molybdenites are liberated. The molybdenites were then rinsed with MQ three times and further rinsed with ethanol, and then dried at $\sim 35$ °C. The concentrated molybdenite aliquots were further purified (removal of any pyrite/chalcopyrite and undissolved silicate phases) by hand under a binocular microscope. The purity of the mineral separate was estimated to be better than 95 %.

This approach, dating multiple independent molybdenite separations from a single vein (Fig. 6A-B), offers the opportunity to check the consistency and closed system behavior of the Re-Os system within the vein and ultimately verify the accuracy of the dates as further
discussed below. In addition, similar to that of zircon U-Pb geochronology, this approach permits calculating a weighted average of dates from the same vein and statistically reduces the effect of the analytical uncertainties.

For molybdenite digestion, a known amount of molybdenite (typically >20 mg) and tracer solution ($^{185}$Re plus isotopically normal Os, Selby and Creaser, 2001) was loaded into a Carius tube (Shirey and Walker, 1995) with 3 ml 15.5 N HCl and 6 ml 16 N HNO$_3$, sealed and placed in steel jackets. The Carius tube bearing jackets were placed in an oven to digest at 220 °C for ~24 hours, which permitted the achievement of both rhenium and osmium isotopic equilibrium between the tracer and sample (Selby and Creaser, 2001). Once cooled to room temperature, the outsides of the tubes were carefully rinsed by MQ to remove any particles and then dried. The bottom parts of the Carius tubes were frozen in a mixture of dry ice and ethanol and then opened from the top by a H$_2$O$_2$ flame in a fume hood. Osmium was isolated from the acid medium using solvent extraction method (CHCl$_3$; 3 x 3 mL) at room temperature (Shen et al., 1996), and then back extracted into HBr (Cohen and Waters, 1996) on a rocker overnight. The Os-bearing HBr solution was dried with the Os further purified by micro-distillation (Birck et al., 1997; Roy-Barman, 1993). After Os extraction, the Re-bearing acid medium was dried and then separated by NaOH-acetone solvent extraction (Cumming et al., 2013; Matthews and Riley, 1970). The Re was further purified via modified standard anion column chromatography (Morgan et al., 1991).

Rhenium and Os were loaded onto outgassed Ni and Pt filaments, respectively. The isotopic compositions of Re and Os were measured by negative thermal ionization mass spectrometry (N-TIMS) as ReO$_4^-$ and OsO$_3^-$, respectively (Creaser et al., 1991; Völkening et al., 1991). The mass spectrometer analysis was conducted on a Thermo Scientific TRITON mass spectrometer at Durham University, with Re and Os isotopic compositions measured using static Faraday cups collection mode. The measured Re and Os isotopic compositions
were corrected using a $^{18}\text{O}/^{16}\text{O}$ and $^{17}\text{O}/^{16}\text{O}$ value of 0.002045 and 0.001113 (Nier, 1950).

The isotopic composition of Re was corrected for instrumental fractionation by fractionation factors defined by the differences between standard Re analyses and the value of Gramlich et al. (1973; $^{185}\text{Re}/^{187}\text{Re} = 0.59738$) before and after the sample Re measurements. The Os mass fractionation was monitored in real-time by monitoring the Os isotopic composition of the tracer and corrected using a $^{192}\text{Os}/^{188}\text{Os}$ of 3.08761. The uncertainties of sample Re and Os isotopic composition measurements, tracer calibration, sample and tracer weighting, reproducibility of Re and Os isotope standards, as well as blank abundances and isotopic compositions during the course of study were all propagated. During this study, the Re and Os blanks were 2 pg and 0.5 pg, respectively, with a $^{187}\text{Os}/^{188}\text{Os}$ value of 0.24 ± 0.01 (n = 6, 2σ). The high Re and Os abundances (Table 2) in these samples in comparison to the blanks result in a negligible blank correction to the final date (<0.035 %).

The molybdenite Re-Os model date was calculated using the equation $t = \ln \left( \frac{^{187}\text{Os}/^{187}\text{Re} + 1}{\lambda} \right)$, in which $\lambda$ is the decay constant (Smoliar et al., 1996). Using the $^{187}\text{Re}$ decay constant of (Selby et al., 2007) yield model dates that are ~0.008 Ma younger than those calculated with $^{187}\text{Re}$ decay constant of (Smoliar et al., 1996). This small systematic discrepancy is a level similar to the analytical uncertainty and the model dates calculated with two $^{187}\text{Re}$ decay constants overlap within uncertainties. Therefore, we consider that using either $^{187}\text{Re}$ decay constants does not impact the interpretation of the final date.

To evaluate the accuracy and reproducibility of the molybdenite Re-Os analytical approach, the Henderson molybdenite reference material (RM8599) was run during the course of this study. Nine analyses of RM8599 yield a weighted mean of 27.695 ± 0.038 (2σ, MSWD = 0.53, Fig. 6C), which is in good agreement with the recommended value 27.66 ± 0.10 Ma (Markey et al., 2007; Zimmerman et al., 2014), and previous analysis at Durham (Lawley and Selby, 2012).
A further consideration, although negligible, is the presence of trace amounts of common\(^{187}\)Os in the molybdenite (Selby and Creaser, 2001; Stein, 2014; Stein et al., 2001), which is not accounted for by the analytical method using a tracer solution with a normal Os isotopic composition (Markey et al., 2007; Selby and Creaser, 2001). However, the levels of common\(^{187}\)Os in most published molybdenite is \(<23\) ppt, and typically \(<6\) ppt (Spencer et al., 2015; Stein, 2014; Stein et al., 2001; Zimmerman et al., 2014). Therefore, this level of common\(^{187}\)Os will only alter the model dates of the studied samples by 1-10 kyr (33 of the 42 dates \(<3\) kyr). Further, our approach of obtaining multiple independent dates from a single vein (Fig. 6A-B) is an independent test for the consistency of the Re-Os systematics along the vein. The reproducible dates (Table 2) from the same vein verifies the accuracy of the model dates, and support the negligible presence of common\(^{187}\)Os in molybdenite since a single vein can be regarded as a specific geological event (Stein, 2006). Finally, as other tracers (e.g., \(^{190}\)Os-\(^{188}\)Os tracer) are calibrated against the common Os solution/tracer (Markey et al., 2007), their precision will never be better than that of the common Os tracer, and as a result dates obtained using a common Os tracer, will, in principle, have smaller uncertainties inherited from tracer calibration.

3.3 Uncertainties of duration and cooling rates

As this study is utilizing high-precision dates to constrain the ore-forming process, e.g., the duration and cooling rates, it is particularly important to evaluate the uncertainties associated with the dates. For durations calculated from the same radio-isotopic system, e.g., Re-Os or U-Pb, either analytical uncertainty or analytical + tracer calibration uncertainties are considered for dates using the same tracer or different tracer, respectively. For example, when calculating the duration of mineralization (Re-Os system), or the temporal gap between the emplacement of the Rongmucuola pluton and the P porphyry (U-Pb system), only analytical
uncertainties needs to accounted. Considering that molybdenite Re-Os and zircon U-Pb dates are ‘independent’ chronometers, the uncertainty related to calibration of both systems must be considered (i.e., tracer calibration and decay constants) when comparing the two chronometers to derive a geological inference.

The uncertainties of the duration or rates are propagated using a Monto Carlo simulation. When calculating the uncertainties of the cooling rates, in addition to considering the analytical uncertainties of the Re-Os dates, the uncertainty in the trapping temperatures of quartz hosted fluid inclusions are used (± 20 °C) (Li et al., 2017b).

4 Results

The uncertainties of the U-Pb and Re-Os dates are presented as ± x/y/z (analytical uncertainty /+tracer calibration /+decay constant uncertainties) in Figures 7 and 8 (Tables 1-2), respectively.

4.1 Zircon U-Pb geochronology

All the analysed zircon grains from the Rongmucuola pluton, with the exception of two grains, yielded concordant 206Pb/238U and 207Pb/235U dates (Fig. 7A).

The 8 zircons from the western Rongmucuola pluton yielded 206Pb/238U dates of 17.975 - 17.057 Ma (Fig. 7B) with Th/Uzircon values of 0.47 - 0.59 (Table 1). The youngest date obtained from a single analysis is 17.057 Ma, but this date could not be reproduced by other zircon grains. A weighted mean, 17.142 ± 0.014/0.014/0.023 Ma, was calculated from the youngest cluster of dates that yielded a statistically acceptable population (MSWD = 1.12; n = 3).

Seven analyses from the P porphyry yielded 206Pb/238U dates between 16.115 and 15.998 Ma (Fig. 7B), and the youngest cluster of dates gives a weighted mean of 16.009 ±
0.016/0.017/0.024 (MSWD = 0.90; n = 3) with Th/U\textsubscript{zircon} values of 0.92 - 1.11 (Table 1). This

group included two fragments of a single elongate prismatic zircon grain (z4A and z4B, Fig. 3D) which yielded indistinguishable dates.

Seven zircons from the quartz diorite yield 206\textsuperscript{Pb}/238\textsuperscript{U} dates of 15.189 - 15.116 Ma (Fig. 7B) with 6 yielding a weighted mean of 15.166 ± 0.010/0.011/0.020 (MSWD = 1.67).

4.2 Molybdenite Re-Os geochronology

For the 13 molybdenite-bearing veins, multiple (2 - 7) individual molybdenite separates (e.g., Fig. 6) were obtained for each vein, except for 3 veins which only yielded one separate each as a result of a low concentration of molybdenite. In total 42 analyses were conducted, of which six of them were reprocessed after Li et al. (2017b) either through re-analysis of the purified Re fraction or analysis a new aliquot of the identical mineral separate. Overall, the Re and 187\textsuperscript{Os} abundances of these samples range from 60 to 504 ppm, and from 10 to 85 ppb, respectively (Table 2).

The weighted mean dates were calculated for each vein, with the exception of the three veins with only one analysis. The weighted mean dates or model dates (three samples with one analysis) for these veins span from 16.126 ± 0.008/0.060/0.077 (n = 3, MSWD = 1.9) to 15.860 ± 0.010/0.058/0.075 Ma (n = 2, MSWD = 0.021) and do not form a single statistic population (Fig. 8). Dates of A, B and D veins vary from 16.126 ± 0.008/0.060/0.077 (n = 3, MSWD = 1.9) to 16.040 ± 0.007/0.058/0.075 Ma (n = 4, MSWD = 0.69), from 16.107 ± 0.015/0.065/0.082 (n = 1) to 15.939 ± 0.006/0.058/0.075 Ma (n = 4, MSWD = 1.6), and from 16.088 ± 0.007/0.059/0.076 (n = 3, MSWD = 1.6) to 15.860 ± 0.010/0.058/0.075 Ma (n = 2, MSWD = 0.021), respectively. No relationship is observed between dates and vein types, although all A veins are older than 16.040 ± 0.007/0.058/0.075 Ma (Fig. 8).
5 Discussion

5.1 A refined chronology of the Qulong porphyry system

Regarding the emplacement ages of the dated igneous units, our preferred interpretation is the weighted averages calculated from the youngest groups of $^{206}\text{Pb}/^{238}\text{U}$ dates. For the Rongmucuola pluton, the youngest date cannot be reproduced by other grains. To avoid biasing the emplacement age to a younger date, which could potential be related to lead loss, we consider the weighted average of the youngest cluster, $17.142 \pm 0.014/0.014/0.023$ (MSWD = 1.12; n = 3), as our preferred estimate of the emplacement age of the western Rongmucuola pluton (Fig. 7).

The youngest cluster of zircon grains from the P porphyry comprises two fragments of a single elongate prismatic zircon grain (z4A and z4B, Fig. 3D), therefore supporting the hypothesis that the lead loss domains, if occurred, was efficiently removed by the chemical abrasion process. As such, we consider the weighted mean, $16.009 \pm 0.016/0.017/0.024$ (MSWD = 0.90; n = 3), as the best estimate of the emplacement age of the P porphyry.

The best estimated emplacement age of the quartz diorite is $15.166 \pm 0.010/0.011/0.020$ (MSWD = 1.67, n = 6), represented by the weighted average calculated from 6 of the 7 zircon Miocene dates (Fig. 7B).

The oldest discordant zircon from the Rongmucuola pluton is interpreted as a result of the incorporation of xenocrystic cores from an older magmatic source. A similar scenario is preferred to explain the $\sim36.267$ Ma grain from the quartz diorite (Table 1). The older dates from each pluton are interpreted as a mixture of two age domains (Chelle-Michou et al., 2014), e.g., in addition to the zircon rims formed during the emplacement of the intrusions, the zircons also contain xenocrystic cores, or represent a protracted zircon crystallisation history in an upper crustal magma chamber, and are not discussed further.
As discussed above, our new CA-ID-TIMS U-Pb zircon ages provided here agree with the relative chronology based on field observations (Fig. 2A) and timeframe determined via *in-situ* U-Pb zircon techniques (Fig. 2B), but at a much higher temporal resolution (~0.1 %).

Field relationships show that the Rongmucuola pluton and the quartz diorite are pre-ore and post-ore without further evidence of Miocene magmatism outside this interval (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Yang et al., 2015; Zhao et al., 2016). In this regard, the U-Pb ages of the Rongmucuola pluton and the quartz diorite constrain the duration of mineralization-related magmatism at Qulong to 1.976 ± 0.017 Myr. The aplite is observed only in the Rongmucuola pluton without direct crosscutting relationship with both the P and X porphyries (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016), and previous SIMS U-Pb zircon geochronology showed only evidence of inheritance (Li et al., 2017b). The minimum emplacement age of the aplite is estimated as 16.126 ± 0.077 Ma (including all sources of uncertainty, e.g., analytical uncertainty + tracer calibration + decay constant uncertainties) based on a Re-Os molybdenite date from an A-type quartz-vein hosted by the aplite (sample 313-145, discussed below). Given the emplacement age of the P porphyry is 16.009 ± 0.024 Ma (with all uncertainties), the aplite predates the P porphyry and is at least 0.117 ± 0.081 Myr older.

For molybdenite Re-Os dates, given the similar morphology of molybdenite grains and alteration assemblages within a vein (e.g., Fig. 4), we assume that molybdenite grains in each vein formed at the same time. This assumption, in part, has been supported by the fact that model ages of multiple analyses of each vein (except 1605-33) overlap within uncertainty. For veins with multiple analyses, when the MSWD is acceptable for the number of analyses (Wendt and Carl, 1991), the weighted means are interpreted as molybdenite crystallization ages. By inference, e.g., close spatial association between molybdenite and the
gangue minerals, quartz-bearing fluid inclusions with opaque minerals (Li et al., 2017b), we suggest that molybdenite crystalized at the same time as the gangue minerals, and therefore can be used to represent the vein-formation ages. For 3 veins with one analysis, given the confidence of data reproducibility from multiple analyses of the single vein, the model ages were interpreted as vein formation ages. However, an exception is sample 1605-33, which gives a MSWD of 5 (Fig. 8) and is interpreted to reflect the presence of multiple generations of molybdenite, which nonetheless are not recognized petrographically. As such, the mean calculated from the youngest two analyses is interpreted as the best estimated formation age of that vein.

The Re-Os ages of this study overlap with the majority of the dates from previous studies within uncertainties (Fig. 2B). However, the Re-Os dates of this study are considerably more precise (~0.1 %, Fig. 7B) and suggest that the majority of the mineralization occurred between 16.126 ± 0.008/0.060/0.077 and 15.860 ± 0.010/0.058/0.075 Ma with a duration of ~ 266 ± 13 kyr. This duration is a minimum estimate as there is no geological evidence to exclude the presence of mineralization beyond this interval, and observations at Qulong suggest that Cu mineralization as whole is earlier than the Mo ore. However, we consider that this duration is reasonable as it has been independently shown that, with an average magma flux (pluton filling rate) of 0.01 km³/yr, a few tens of million tons of Cu, comparable to the amount of Cu at Qulong, can be accumulated within 200 - 300 kyrs (Chelle-Michou et al., 2017).

The Re-Os dates from previous studies (Fig. 2B and appendix Table A2) have appreciable uncertainties of 1.3 - 12.1 %, which we attribute to analytical challenges associated with initial Re-Os studies, e.g., imprecise determinations of the stoichiometry of Os salts used for tracer calibration (Yin et al., 2001), poor external reproducibility of ICP-MS analysis (Stein, 2014) potentially caused by memory effects of Os and unstabilized ion signal.
in the long term, and potential bias between labs (e.g., tracer calibration, data reduction). Given dates of the RM8599 reference materials measured during the course of study agree with the recommended value (Fig. 6C), and the agreement between molybdenite Re-Os and zircon U-Pb chronometers from a preliminary cross-calibration (Selby et al., 2007) with the same Re-Os tracer used in this study, we suggest that, to the best of our knowledge and effort, the molybdenite Re-Os ages from this study are the best estimates of molybdenite crystallization ages at Qulong. For future molybdenite Re-Os geochronology study, we reiterate that measuring and reporting the age values of RM8599 reference materials is essential, coupled with the analysis of a suitable sized aliquant of a mineral separate (e.g., Selby and Creaser, 2004; Stein et al., 2003), and employ the multiple-analysis from a single vein approach proposed here.

5.2 Integrating magmatic and hydrothermal timescales

At Qulong, field observations demonstrate a relatively close spatial association (Fig. 1D-E) between the P porphyry and high-grade Cu ore (Yang et al., 2009; Zheng et al., 2004). In addition, the aplite and X porphyry are very small in size and only distribute discontinuous. Further, the is no other porphyry stocks within 2 km as demonstrated by extensive drilling. Therefore, the P porphyry is regarded as the only observed porphyry stock that can act as a conduit (Hu et al., 2015; Li et al., 2017b; Yang et al., 2009; Zhao et al., 2016). However, this scenario has not been tested due to a lack of cross-cutting relationships at the deposit level. Testing this scenario is possible by dating the P porphyry and mineralization at Qulong with high accuracy and precision, and the null hypothesis is that the porphyry stock is expected to be older than the mineralization.

Considering the emplacement age of the P porphyry is 16.009 ± 0.024 Ma (with all uncertainties at the 2σ level, Fig. 7), 13 of the 16 published molybdenite Re-Os ages overlap
with this emplacement age, while the remaining 3 are older (including all sources of
uncertainty, e.g., analytical uncertainty /+tracer calibration /+decay constant uncertainties).

The three Re-Os dates that are older than the P porphyry agree with the field observations
that all mineralization at Qulong postdate the Rongmucuola pluton and predate the quartz
diorite (Fig. 2A). For our molybdenite Re-Os dates, only sample 313-145, which is hosted by
the aplite, is 0.117 ± 0.081 older than the P porphyry, with other 11 samples that overlap
within uncertainties, and the remaining 1 sample that is nominally younger (Fig. 8).

Excluding analytical issues, the presence of Re-Os dates from this study that predate
the P porphyry implies that mineralization did occur before the emplacement of the P
porphyry. As mentioned above, the aplite is unlikely to be a conduit given its exceptional
small volume (with a width of 0.05 - 2 m) and discontinuous distribution. Together with the
absence of other documented/observed porphyry stocks that predate the P porphyry, the most
likely scenario is that the P porphyry is an intermineral porphyry stock and potentially a part
of the mineralization (e.g., the one yielding ages that predate the P porphyry) was deposited
before the emplacement of the P porphyry. This scenario is arguably supported by the
observation that the highest grade of Mo ore does not occur within the P porphyry (Fig. 1D-E)
(Yang et al., 2009), as intermineral porphyry stocks commonly result in low-grade cores in
porphyry deposits (Sillitoe, 2000). The decoupling between shallow magmatism and volatile
transportation thus makes it unlikely that all metals were transported by the P porphyry stock
at Qulong. However, we reiterate that such a scenario is the most plausible one based on the
available data. Further detailed mapping (crosscutting relationships, alteration) could confirm
this hypothesis.
5.3 Cyclical mineralization pulses and their timescales

The veins in each mineralization pulse of porphyry copper deposits are classified as A, B and D veins (Fig. 3). The temporal succession of these veins records a progressive change in the fluid nature, e.g., temperature, pH and/or oxygen fugacity, and is widely used to trace fluid evolution processes (Gustafson and Hunt, 1975; Sillitoe, 2010). Given the timescale of hydrothermal activity that can be sustained by a single magmatic episode is limited to several tens of kyr (Buret et al., 2016; Cathles et al., 1997; Chelle-Michou et al., 2015; Mercer et al., 2015; Tapster et al., 2016; Weis et al., 2012), porphyry deposits which show prolonged duration (e.g., several Myr, Chiaradia et al., 2014) clearly imply multiple mineralization pulses, as supported by the presence of repetitive A-B-D veins from field observations and high precision dating (Sillitoe, 2010; Spencer et al., 2015; Stein, 2014). However, it is still not clear whether multiple mineralization pulses are only present in some deposits with prolonged duration, or if they are common in porphyry copper deposits in general. Also questionable is the role of multiple pulses for the formation of economic deposits.

Our geochronological data suggest that the mineralization process at Qulong was likely discontinuous during the 266 ± 13 kyr interval (Fig. 8), and characterized by the presence of at least two A-B-D cycles and one incomplete (B-D) cycle. Our Re-Os data constrain the lifetime of the two short-lived A-B-D cycles to 38 ± 11 and 59 ± 10 kyr with a gap of 48 ± 10 kyr between them. The timescales of mineralization cycles constrained here (tens of kyr) by direct dating of ore minerals are comparable with those recently proposed through high-precision dating (Buret et al., 2016; Chelle-Michou et al., 2015; Tapster et al., 2016), titanium diffusion in quartz (Mercer et al., 2015) and numerical modeling (Chelle-Michou et al., 2017; Weis et al., 2012). However, we are aware that although the sample set used here covers the entire hydrothermal fluid evolution path at Qulong (Li et al., 2017b), it is impossible to argue that all stages of mineralization phases have been sampled. In addition,
even though this study arguably presents the most detailed Re-Os geochronology study for a single porphyry Cu-Mo deposit so far, it is very likely that it is still under-sampled. Summarizing, we propose that according to the current study it is likely that the metals at Qulong system were deposited via a cyclic process, but uncertainties remain.

5.4 Rapid cooling during mineralization

Although the A, B and D veins in porphyry copper deposits record the evolution paths of ore-forming fluids (Cooke et al., 2014; Sillitoe, 2010), in most cases the relative chronology of vein types cannot be confidently correlated at the deposit-scale due to a lack of deposit-wide crosscutting relationships. Here, we explore the possibility of coupling high-precision molybdenite Re-Os and quartz fluid inclusion data to reconstruct the cooling paths of the Qulong porphyry system.

Correlating the Re-Os dates in this study and fluid inclusion data (Li et al., 2017b) suggests that the entire hydrothermal system cooled from 425 to 280 °C within 266 ± 13 kyr ($r^2 = 0.797$; Fig. 9). This suggests that the Qulong porphyry system as a whole cooled over the period of ore formation with a cooling rate of 0.55 ± 0.11 °C/kyr. In addition to the long-term cooling trend, the cooling rates of the two complete A-B-D cycles are 1.19 ± 0.82 and 1.27 ± 0.53 °C/kyr, respectively (Fig. 9). A potential concern is an underestimation of the duration caused by the undersampling of Re-Os dates as discussed above, by which the cooling rates could be overestimated. However, we argue that if undersampling is indeed an issue, it is unlikely that the duration will significantly increase given the number and nature (vein type, characteristics of ore related fluid inclusions) of sample set we studied, and the agreement of durations constrained in this study and other independent estimates (see above for detailed discussion). As such we suggest that the estimated cooling rates here, within uncertainties, are reasonable for Qulong. The cooling overarate estimated here propose a
much faster rate than that (0.2 °C/kyr) established from thermochronology (Zhao et al., 2016).

Despite the cooling trend of the first mineralization cycle overlaps with that of the long-term cooling rate within uncertainties, the cooling rates of both cycles are much faster than the long-term cooling rate of the Qulong porphyry system. To the best of our knowledge, the cooling rates presented in this study (between 0.55 and 1.27 °C/kyr) are faster than the majority (with the exception of Grasberg) of the reported cooling rates (McInnes et al., 2005; McInnes and Evans, 2005) with highest temporal resolution (<1 ‰ level).

The cooling process of a hydrothermal system is balanced between heat gain (e.g., magmatism injection and volatile fluxing associated with heat from depth) and heat loss (e.g., conduction to wall rocks, cooling by the circulation of meteoric water) (Cathles et al., 1997; Weis et al., 2012). At Qulong, the known thermal contributors are the Rongmucuola pluton, aplite, P porphyry, X porphyry, quartz diorite and exsolved volatile, with heat loss controlled by cooling via conduction. The Rongmucuola pluton, which hosts the majority of the ore at Qulong, is ~1 Myr older than the P porphyry (Fig. 5). Heat loss modelling predicts that a 40 km wide, 2 km thick sill intruded between 16 to 18 km depth can only sustain a maximum hydrothermal discharge >200 °C for 0.8 Myr, and this duration decreases to 1 kyr for systems at shallower depths (<3 - 5 km) (Cathles et al., 1997). Given the much smaller size (<200 m in diameter vs 40 km wide and 2 km thick) and much shallower (potentially ~3 km as inferred from a fluid inclusion study by Li et al., 2017b vs 16 to 18 km) emplacement depth of the Rongmucuola pluton, we conclude that the Rongmucuola pluton was cooled to <200 °C before the emplacement of the P porphyry. As the volumes of the aplite and X porphyry are small, they would have had limited thermal contribution during ore formation as an intrusion with similar size can only sustain a thermal anomaly for less than a few kyr as indicated by quartz titanium diffusion and heat loss models (Cathles et al., 1997; Mercer et al., 2015). As a result, the only possible main thermal contributors are the P porphyry and
exsolved volatile, although we cannot confidently exclude possible contributions from unexposed and unidentified intrusions. With the assumption that the crystallization/cooling process of the P porphyry lasted for less than tens of kyr according to thermal simulation and diffusion modeling (Cathles et al., 1997; Mercer et al., 2015), the ~266 ± 13 kyr long thermal anomaly accompanying the mineralization process needs heat sources external to the contribution from the P porphyry itself. Therefore, we suggest that a periodic magmatic-hydrothermal fertilization process, e.g., an episodic hydrothermal fluid release during the incremental building of the source pluton where the metals and magmatic fluids were derived, provided the additional thermal contribution that maintained the mineralization process.

The thermal anomalies and faster cooling process discussed above require a mechanism to explain the rapid heat removal. However, an increase of heat loss through conduction is unlikely, and we suggest that the most reasonable scenario is rapid thermal removal via the meteoric water circulation model (Fekete et al., 2016; Weis et al., 2012). Significant meteoric water circulation is only expected in highly permeable rocks, granitoid units which dominate porphyry deposits are less favorable for such a scenario unless they are intensely fractured. A simple calculation suggests that 10 - 25 volume % meteoric water (~100 °C after heating) mixing with 90 - 75 volume % magmatic water is enough to cool the hot (> 500 °C) magmatic fluid to the temperatures recorded by ore-related fluid inclusions (425 - 280 °C). If this is the case, depleted oxygen isotopic composition is expected for the ore-forming fluid at Qulong, which has been demonstrated by our unpublished data and it is the object of a future publication.

Given each mineralization cycle will bring additional heat to the cooling porphyry system, the rapid and periodic cooling process discussed above predicts resetting of thermal chronometers of earlier mineralization cycles by later mineralization cycles. This prediction is supported by the identical Ar-Ar ages of the earlier potassic stage biotite and later phyllic
stage sericite, and the significant scatter of zircon U-Th-He systematics (Zhao et al., 2016), although the uncertainties of these dates do not permit a robust evaluation for the timeframe of these events (e.g., timeframe of formation and cooling history).

5.5 Formation of the world class Qulong Cu-Mo deposit

The detailed high-precision zircon U-Pb and molybdenite Re-Os geochronology presented here allow us to propose a model for the formation of the world-class Qulong porphyry Cu-Mo deposit (Fig. 10).

At ~17 Ma, magma generated in the lower crust was emplaced into the upper crust (the emplacement of the Rongmucuola pluton and the aplite), and was potentially associated with minor mineralization (Fig. 10A). Due to a lack of porphyry stock at this time, the potential conduit transporting magma and volatile to shallow mineralization level is the north-south trending fault. The meteoric water could circulate downward into the system through the fault system.

After the emplacement of the Rongmucuola pluton and aplite, a temporary cessation of magmatism is observed at shallow levels. At ~16 Ma, potentially triggered by a magma injection event, the magma and metal-bearing fluid rose to the shallow crustal levels periodically, as indicated by the formation of mineralization, and the P and X porphyries (Fig. 10B). The formation of these intermineral porphyry stocks may have remobilized/destroyed the earlier mineralization, and resulted in a relatively low Mo grade core within the P porphyry. Episodic volatile releasing events are potentially the trigger of the mechanical failure of the cupola and the upper crust, as evidenced by the formation of two stages of breccia (Fig. 10C). Hydrothermal fluids associated with the breccia pipe caused further alteration and mineralization assemblages, and may have also remobilized/destroyed a portion of earlier stage mineralization. The system is thermally balanced by heat input from
the magmatism and volatiles release events, and heat remove by conduction to wall rock and
meteoric water circulation and mixing.

After ~15.860 Ma, the system experienced a further quiet period until ~15.2 Ma. Most
likely initiated by injection of new magma that caused thermal rejuvenation, the quartz diorite
was formed, which marks the cessation of Miocene magmatism and mineralization at Qulong
(Fig. 10D). Minor amount of magmatic fluid associated with the emplacement of the quartz
diorite, plus the circulation of meteoric water caused further alteration as represented by the
low temperature alteration in the quartz diorite.

6 Implications for porphyry copper system genesis

Radio-isotopic dating is widely applied to constrain the timescales of porphyry copper
deposits formation, and give variable timescales from tens of kyr to several Myr (Chiaradia et
al., 2014). Given the timescale is a primary feature of the systems studied, which should be
an independent parameter regardless of the dating methods used, the strong positive
relationship between analytical precision and the quoted duration is not expected (Fig. 11).
Therefore, this relationship (Fig. 11) suggests that at least some of the prolonged durations
proposed for porphyry systems may be an artefact of over interpreting low-precision data
without evaluation of their accuracy. Recent simulation studies suggest that the duration of
ore formation is potentially one the of controlling factors for the formation of giant deposits
(Chelle-Michou et al., 2017; Chiaradia and Caricchi, 2017). However, it is obvious that such
a “long” duration (e.g., tens to hundreds of kyr) is still beyond the precision of micro-beam
analysis (Chiaradia et al., 2014). Consequently, we reiterate that high-precision
gEOchronology is essential for resolving the magmatic-mineralization timescales of porphyry
copper systems at the tens of kyr level.
The genetic model of porphyry copper deposits suggests that the porphyry stocks are the conduits conveying metal-bearing volatile from the lower crustal magma chamber to shallow levels (Cooke et al., 2014; Richards, 2011; Sillitoe, 2010). However, in most cases this scenario is very difficult to test by crosscutting relationship for two reasons. First, the porphyry stock is exceptionally small compared to the size of the deposit (Cooke et al., 2014; Richards, 2011; Sillitoe, 2010), such that the crosscutting relationship between the porphyry stock and mineralization phases are only available in a very limited space, e.g., surrounding the porphyry stock. Secondly, if there are mineralization phases before the emplacement of the porphyry stock, they will likely be overprinted/remobilized/destroyed by the emplacement of the porphyry stock where the potential crosscutting relationships are available. Therefore, the best approach to test this hypothesis is via high precision dating, and our high precision dating at Qulong arguably suggests that it is possible to transport metal-bearing volatile without the presence of a contemporary porphyry stock. This scenario has been supported by observations at modern volcanoes whereby degassing can occur before/without magma eruption at shallow levels (Shinohara, 2008). If this finding is correct, it further suggests that the absence of a contemporary porphyry stock does not indicate that the mineralization is not a porphyry style deposit, which has broad implication for deposit type clarification and mineral exploration. In addition, the presence of mineralization before the emplacement of a porphyry stock or even the absence of porphyry stocks implies that dating porphyry stocks does not necessarily brackets the entire porphyry ore formation process. In this case dating the mineralization directly is critical to yield a comprehensive understanding of the timescale of ore formation, as well as of any associated magmatism (Stein, 2014).

The cyclic mineralization process inferred by our Re-Os dates suggests that the periodic mineralization process also operates for deposits with a short duration (e.g.,
hundreds of kyr overall duration), which is the same as those observed in porphyry systems with prolonged (Myr level) duration (Spencer et al., 2015; Stein, 2014; Zimmerman et al., 2014). Cyclic magmatic process in porphyry Cu deposits is ubiquitously recorded in many magmatic minerals, e.g., zoning/resorption textures of plagioclase, quartz and zircon (Buret et al., 2016; Mercer et al., 2015; Tapster et al., 2016; Williamson et al., 2016). By inference, we suggest that the periodic magmatic-hydrothermal processes during ore formation proposed here are common in porphyry copper deposits, and are linked with the episodic magmatic process occurring at depth within the source pluton. The significance of this cyclic process is that each mineralization pulse can enhance the economic resource in a geologically focused area. As such, the cyclic process presented here potentially is one of the contributing factors to form giant deposits, and is the key to differentiate economic and sub-economic deposits. However, it is not clear yet what fundamentally controls this cyclic process, although the periodic re-fertilization of volatile linked with source pluton evolution might be critical (Williamson et al., 2016).

The cyclic cooling process proposed here, together with the potential thermal resetting of the U-Th-He and Ar-Ar chronometers, should be considered in future cooling history studies of porphyry copper systems. In addition, the rapid cooling (0.55 - 1.27 °C/kyr) and periodic volatile recharge process proposed here suggest that external fluid potentially is a critical cooling agent, as recently proposed by numerical modeling and isotopic study of porphyry deposits (Fekete et al., 2016; Weis et al., 2012).

A final note concerns the uncertainties when comparing Re-Os dates from different labs and with U-Pb and other chronometers. Reporting the measured values of the RM8599 is a must to evaluate the (relative) accuracy of the Re-Os dates (Stein, 2014), and also the analysis of a suitable sized aliquant from a mineral separate is of primary importance (Selby and Creaser, 2004; Stein et al., 2003). Currently the decay constant uncertainties (238U and
$^{187}\text{Re}$) are the main source of uncertainties when comparing Re-Os and U-Pb systems. In addition, the inter-laboratory agreement of Re-Os systems has only been demonstrated between a few labs (Markey et al., 2007), so direct comparison of dates beyond these laboratories potentially involves an unaccounted bias. Given that significant progress has been achieved for the precision and accuracy of the zircon U-Pb chronometers in the past ten years via the community-driven EARTHTIME initiative (Condon et al., 2015; McLean et al., 2015), and the analytical precision of molybdenite Re-Os system is down to 1‰, now it is an appropriate time to start a further evaluation between the molybdenite Re-Os and zircon U-Pb systems (Selby et al., 2007) as suggested by Chiaradia et al. (2014). Following the example of the U-Pb community, shared tracer/reference solutions and transparent data reduction platform should be adopted in the Re-Os community with the aim to standardize the Re-Os chronometer, reduce the potential bias between laboratories and ultimately improve the accuracy. Once achieved, data from all laboratories can be integrated without loss of resolution (due to inter-laboratory bias) and will enable better utilization of data from the literature.

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List of figures and Tables
Fig. 1. Geological setting of the Lhasa terrane and deposit geology map of Qulong porphyry Cu-Mo deposit. A) Simplified geological setting of the Lhasa terrane, Gangdese Magmatic Belt and Gangdese Porphyry Copper Deposits Belt. B) Geological map of the Qulong porphyry Cu-Mo deposit. C) Cross section showing the crosscutting relationship of the Miocene units at the Qulong porphyry Cu-Mo deposit. D-E) Cross section showing the Cu and Mo grades. Revised after Yang et al. (2009) and Zhao et al. (2016).

Fig. 2. Geological timeframe of the intrusion and mineralization at Qulong constrained by crosscutting relationships and geochronology. A) Relative chronology of the Miocene magmatism and mineralization at Qulong porphyry Cu-Mo deposit. B) Published geochronology data of the Miocene magmatism and mineralization at Qulong porphyry Cu-Mo deposit. See text for discussion. Detailed dates are provided in Table A1 - A2. For source of dates, 1, Hu et al., 2015. 2, Wang et al., 2006. 3, Li et al., 2004. 4, Zhao et al., 2015. 5, Hou et al., 2004. 6, Li et al., 2017. Revised after Li et al. (2017b).

Fig. 3. Petrography of the Rongmucuola pluton (A), the P porphyry (B), and post-ore quartz diorite (C). CL images of the analyzed zircon grains of the Rongmucuola pluton, the P porphyry and the quartz diorite (D).

Fig. 4. Representative A (A), B (B) and D (D) veins from the Qulong porphyry Cu-Mo deposit. A) The A veins are characterized by 0.3 - 0.8 cm wide discontinuous granular quartz veinlets with narrow (<3 mm) K-feldspar selvages, which host disseminated chalcopyrite and molybdenite; B) The B veins are represented by veinlets (0.4 - 2 cm wide) comprising crystalline quartz with suture in the centre, and possess irregularly distributed K-feldspar selvage, and predominantly host molybdenite and chalcopyrite along the margin of the vein; C) The definition of vein types are illustrated following those of Gustafson and Hunt (1975) and Sillitoe (2010). D) The D quartz veins (0.5 - 6 cm) host euhedral pyrite, with sericite and anhydrite (0.1 and 1.5 cm) selvages, and minor disseminated chalcopyrite and molybdenite. E-F) Representative sections showing the crosscutting relationships between different stages of veins, e.g., A type quartz vein cut by later A and B type veins. See text for details.

Fig. 5. The impact of Th disequilibrium correction as illustrated by the youngest group of zircon dates of the P porphyry. A) the weighted mean of the youngest three dates as a function of variable Th/U values used for Th disequilibrium correction. B) the MSWD of the weighted mean of the youngest three dates as a function of the variable Th/U values used for
Th disequilibrium correction. Also showing in the figures are the applied Th/U value with uncertainty for the P porphyry (see text for detailed discussion).

Fig. 6. A) Cartoon illustrating the approach of obtaining several independent molybdenite separates from a single vein, which permits testing the consistency of Re-Os systematics along the vein, verify the accuracy of the Re-Os ages and reducing analytical uncertainties. B) The multiple independent molybdenite separates approach illustrated by a B vein analysed in this study. C) The model ages of the Henderson molybdenite reference materials (RM8599) analysed during the course of study, which yield a weighted mean age of 27.695 ± 0.038 (n = 9, MSWD = 0.53) and overlap with the recommended value of 27.66 ± 0.10 (Markey et al., 2007; Zimmerman et al., 2014).

Fig. 7. The CA-ID-TIMS U-Pb zircon dates of the Rongmucuola pluton, the P porphyry and the quartz diorite at Qulong porphyry Cu-Mo deposit. A) The Wetherill Concordia diagram of the analysed zircon grains. B) $^{206}\text{Pb}/^{238}\text{U}$ ages of the analysed zircon grains and the weighted mean ages of the Rongmucuola pluton, the P porphyry and post-ore quartz diorite.

Fig. 8. The ID-NTIMS molybdenite Re-Os geochronology data of molybdenite veins at Qulong, with probability density plot of different type of veins shown.

Fig. 9. Cooling history of Qulong porphyry Cu-Mo system during ore formation, with vein-formation ages and temperatures constrained by molybdenite Re-Os dating in this study and fluid inclusion data of Li et al. (2017b). The uncertainties of fluid inclusions are shown as ±10 °C (1 sigma), while the uncertainties of Re-Os dates are smaller than the width of the symbol.

Fig. 10. Cartoon showing the genetic model of the Qulong porphyry Cu-Mo deposit. See figure and text for detailed information.

Fig. 11. Analytical precision used in porphyry copper deposits studies and inferred timescales exhibiting a positive relationship. Data are from Chiaradia et al. (2014). See text for discussion.

Table 1. CA-ID-TIMS U-Pb data of Miocene intrusive rocks at Qulong.

Table 2. Re-Os data of molybdenite samples from Qulong.
Supplementary tables

Table A1, Published zircon U-Pb geochronology dates at Qulong

Table A2, Published molybdenite Re-Os and zircon U-Th-He geochronology dates at Qulong
Fig. 1

A) ASIA  INDIA

B) Cu-Mo orebody

C) Dacite-rhyolite porphyry Rongmucuola pluton (east)
P porphyry X porphyry Quartz diorite Breccias

D) Elevation (m)

Cu grade Mo grade

Cu grade, wt % >0.6 >0.3 >0.1 <0.1 Mo grade, wt % >0.06 >0.03 >0.01 <0.01

B) C) 200 m 500 m

N 200 km

ZK313 ZK001 ZK1405 ZK1605

D) E) 200 m

100 km

E) Porphyry Cu deposits
The document contains a table and a graph showing the dates of various geological events. The table compares different intrusive and mineralization events, with columns for Rongmucuola pluton, P porphyry, X porphyry, Post-ore Q-diorite, and Mineralization. The graph illustrates the relative ages of these events with dates in blue and purple, indicating LA and SHRIMP/SIMS dates from different studies, which are considered as emplacement ages. The table provides mean and 2 standard deviations of all data from the same intrusion dated by the same method, with specific references to LA-ICP-MS, SIMS, and Zircon U-Th-He methods. The graph also includes data from Ar-Ar and Re-Os methods, with a note indicating the mean and 2 standard deviation of all data from the same intrusion dated by the same method.
A) quartz-sericite halo

B) K-feldspar halo

C) A quartz vein with trace molybdenite and chalcopyrite

D) quartz-molybdenite ± chalcopyrite (± suture)

E) Later barren A quartz vein cut by later A vein

F) Note the ubiquitous chlorite alteration

G) Discontinued granular quartz vein

H) Crystalline quartz vein

I) Note the disseminated Cu-Mo mineralization

J) Note the ubiquitous chlorite alteration

K) scale bar is 1 cm
Fig. 5

(A) Weighted mean date (Ma) vs. Th/U

(B) MSWD of mean vs. Th/U

applied value for P porphyry
Th/U = 3.6 ± 2
A) multiple independent molybdenite separations from a single vein

molybdenite bearing quartz vein

separation 1 separation 2 separation 3 separation 4

B) multiple independent molybdenite separations from a single vein

1 cm

C) 27.40 27.50 27.60 27.70 27.80 27.90

Date (Ma)

box heights are 2σ

Mean = 27.695 ± 0.038 Ma, n = 9, MSWD = 0.53

Recommended value = 27.66 ± 0.10
Rongmucuola pluton
17.142 ± 0.014 /0.014 /0.023 Ma
n = 3, MSWD = 1.12

P porphyry
16.009 ± 0.016 /0.017 /0.024 Ma
n = 3, MSWD = 0.90

Quartz diorite
15.166 ± 0.010 /0.011 /0.020 Ma
n = 6, MSWD = 1.67
Fig. 8

16.126±0.008
/0.060/0.077
MSWD=1.9
16.098±0.013
/0.066/0.083
MSWD=0.6
16.107±0.015
/0.065/0.082
16.088±0.007
/0.058/0.075
MSWD=0.43
16.088±0.007
/0.059/0.076
MSWD=1.6
16.050±0.005
/0.056/0.073
MSWD=0.33
16.040±0.007
/0.058/0.075
MSWD=0.69
15.981±0.007
/0.059/0.076
MSWD=1.3
15.943±0.007
/0.058/0.075
MSWD=0.09
15.939±0.006
/0.058/0.075
MSWD=1.6
15.877±0.006
/0.056/0.073
MSWD=5.0
15.860±0.010
/0.058/0.075
MSWD=0.021
15.886±0.007
/0.058/0.075
MSWD=0.87
16.107±0.015
/0.065/0.082
16.088±0.007
/0.058/0.075
MSWD=0.43
16.088±0.007
/0.059/0.076
MSWD=1.6
16.036±0.012
/0.065/0.082
16.011±0.015
/0.064/0.081
15.981±0.007
/0.059/0.076
MSWD=1.3
15.943±0.007
/0.058/0.075
MSWD=0.09
15.939±0.006
/0.058/0.075
MSWD=1.6
15.877±0.006
/0.056/0.073
MSWD=5.0
15.860±0.010
/0.058/0.075
MSWD=0.021
15.886±0.007
/0.058/0.075
MSWD=0.87

**Weighted Mean ± Uncertainties**

- **Analytical Uncertainty**
- **Tracer Uncertainty**
- **Decay Constant Uncertainty**

**Sample Descriptions**
- **Sample Number (Drill Core No.-Depth)**
- **Vein Type**
- **Box Heights are 2σ**

**Legend**
- **A vein**
- **B vein**
- **D vein**

**Emplacement Age of P porphyry**
16.009±0.024

**Density Plot**
- **Date (Ma)**
- **Plot Density**
- **A vein**
- **B vein**
- **D vein**
$0.55 \pm 0.11 \, ^\circ \text{C/kyr (} r^2 = 0.797 \)$

$1.27 \pm 0.53 \, ^\circ \text{C/kyr}$

$1.19 \pm 0.82 \, ^\circ \text{C/kyr}$

A vein
B vein
D vein

Fig. 9

Temperature ($^\circ \text{C}$)

Age (Ma)

samples with depths >211 m
A) ~17 Ma, Magma generated at the lower crust emplaced into the upper crust as represented by the emplacement of the Rongmucuola pluton and the aplite. Potential mineralization could be formed at this stage, in which case the N-S fault is a conduit for the transportation of magmatism and hydrothermal fluid. The meteoric water also can circulate downward through the fault system.

B) ~16 Ma, Potentially accompanied with new magma injection events, metals bearing fluid periodically rose up via the fault and caused intensive alteration and mineralization. The P porphyry was emplaced at this time, and was followed by the formation of the X porphyry. The P porphyry and the fault were the main conduits. As an inter-mineral porphyry stock, P porphyry potentially remobilized/destroyed the existed mineralization and formed the low Mo grade core. The system is thermally balanced by heat input from magmatism and volatile release, and heat remove through conduction and meteoric water circulation.

C) Following this was the mechanical failure of the cupola and the crust in the upper crust, as represented by the formation of the breccia pipe.

D) After ~15.860 Ma, the system experienced a relatively quite peroid until 15.2 Ma when a thermal rejuvenation was initiated by a magma injection event, as represented by the quartz diorite. The minor fluid caused low temperature alteration to the quartz diorite without further mineralization.
Fig. 11

\[ r^2 = 0.6031 \]

Duration in Myr

Precision in Myr

Qulong

\[ r^2 = 0.6031 \]
<table>
<thead>
<tr>
<th>Composition</th>
<th>Isotopic Ratios</th>
<th>Dates (Ma)</th>
</tr>
</thead>
<tbody>
<tr>
<td>z4</td>
<td>0.59</td>
<td>1.95</td>
</tr>
<tr>
<td>z7</td>
<td>0.55</td>
<td>3.38</td>
</tr>
<tr>
<td>z1</td>
<td>0.57</td>
<td>4.82</td>
</tr>
<tr>
<td>z5</td>
<td>0.52</td>
<td>2.71</td>
</tr>
<tr>
<td>z3</td>
<td>0.47</td>
<td>0.59</td>
</tr>
<tr>
<td>z9</td>
<td>0.52</td>
<td>9.02</td>
</tr>
<tr>
<td>z8</td>
<td>0.51</td>
<td>9.31</td>
</tr>
<tr>
<td>z6</td>
<td>0.59</td>
<td>1.59</td>
</tr>
<tr>
<td>z2</td>
<td>1.11</td>
<td>4.58</td>
</tr>
<tr>
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<tr>
<td>z3</td>
<td>0.84</td>
<td>5.10</td>
</tr>
<tr>
<td>z6</td>
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<tr>
<td>z5</td>
<td>1.11</td>
<td>4.78</td>
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<tr>
<td>z4B</td>
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<td>6.07</td>
</tr>
<tr>
<td>z4A</td>
<td>0.92</td>
<td>3.40</td>
</tr>
</tbody>
</table>

1. Corrected for initial Th/U disequilibrium using radiogenic 206Pb and Th/U([magma]).
2. Measured ratio corrected for fractionation, tracer and blank contribution only.
3. Measured ratios corrected for fractionation, tracer and blank.
4. Assuming concordance between the U-Pb and Th-Pb systems.
5. Total mass of radiogenic Pb.
6. 4% discordance = 100 - (100 * (206Pb/238U date) / (207Pb/235Pb date))
7. Ratios of radiogenic Pb (including 208Pb) to common Pb.
8. Ratio of radiogenic Pb (including 208Pb) to common Pb.
9. Measured ratio corrected for fractionation and spike contribution only.
10. Measured ratios corrected for fractionation, tracer and blank.
### Table 2 Re-Os data of molybdenite samples from Qulong

<table>
<thead>
<tr>
<th>Sample</th>
<th>wt (g)</th>
<th>Re (ppm)</th>
<th>2 sigma (abs)</th>
<th>167Re (ppm)</th>
<th>2 sigma (abs)</th>
<th>166Os (ppm)</th>
<th>Date (Ma)</th>
<th>2 sigma (abs)</th>
<th>average (Ma)</th>
<th>2 sigma abs uncertainties</th>
<th>Xσ</th>
<th>Yσ</th>
<th>Zσ</th>
<th>vein type</th>
<th>depth (meter)</th>
<th>Temperature (°C)</th>
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<td>1605-211_2-4</td>
<td>0.042</td>
<td>21.2</td>
<td>0.7</td>
<td>132.1</td>
<td>0.4</td>
<td>35.4</td>
<td>0.1</td>
<td>16.084</td>
<td>0.007</td>
<td>16.088</td>
<td>0.007</td>
<td>0.58</td>
<td>0.075</td>
<td>A</td>
<td>211</td>
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<td>59.9</td>
<td>0.3</td>
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<td>16.132</td>
<td>0.008</td>
<td>16.126</td>
<td>0.008</td>
<td>0.06</td>
<td>0.077</td>
<td>A</td>
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<tr>
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<td>16.126</td>
<td>0.008</td>
<td>0.06</td>
<td>0.077</td>
<td>A</td>
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<td>425</td>
</tr>
</tbody>
</table>

*data reprocessed (re-run the Re separations or digest a new aliquot of the same mineral separations) after LI et al. 2017, Mineralium Deposita

*analytical and tracer uncertainty

*analytical, tracer and decay constant uncertainty of Smoliar et al., 1996

*vein formation temperatures constrained by fluid inclusion study from Li et al., 2017

*analytical uncertainty

*tracer uncertainty