**In-situ Lu – Hf geochronology of calcite**

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**Abstract.** The ability to constrain the age of calcite formation is of great utility to the Earth Science community, due to the ubiquity of calcite across a wide spectrum of geological systems. Here, we present the first in-situ laser ablation inductively coupled tandem quadrupole mass spectrometry (LA-ICP-MS/MS) Lu–Hf ages for calcite, demonstrating geologically meaningful ages for IOCG and skarn mineralisation, carbonatite intrusion and low grade metamorphism. The analysed samples range in age between ca. 0.9 Ga and ca. 2 Ga with uncertainties between 1.7% and 0.6% obtained from calcite with Lu concentrations as low as ca. 0.5 ppm. The Lu–Hf system in calcite appears to be able to preserve primary precipitation ages over a significant amount of geological time, although further research is required to constrain the closure temperature. The in-situ approach allows calcite to be rapidly dated while maintaining its petrogenetic context with mineralization and other associated mineral processes. Therefore, LA-ICP-MS/MS Lu–Hf dating of calcite can be used to resolve the timing of complex mineral paragenetic sequences that are a feature of many ancient rock systems.

**1 Introduction**

Calcite (\text{CaCO}_3) is the main mineral phase of most carbonate sedimentary rocks and their metamorphic equivalents, is a common diagenetic phase and is a major component of carbonatites. Calcite is also a common product of hydrothermal alteration and constituent of mineralising systems where it may precipitate from fluids during pre-ore, ore-stage, and post-ore forming processes (Debruyne et al., 2016). The ability to directly date calcite unlocks the possibility to constrain the timing of a vast array of geological processes that can be difficult to date using conventional methods.

Accurate in-situ U–Pb geochronology of calcite has been applied to a variety of geological systems (e.g; Li et al., 2014; Ring and Gerdes, 2016; Roberts and Walker, 2016). However, calcite often incorporates significant quantities of Pb during crystallisation (i.e. ‘initial’ or ‘common’ Pb), which can limit the utility of U–Pb geochronology (Rasbury and Cole, 2009). Moreover, Pb is highly fluid mobile (Brugger et al., 2016), so it can be difficult to obtain primary age information with the U–Pb method in hydrothermal or strongly-altered systems (Roberts et al., 2020; Simpson et al., 2021b). Further, given the propensity for calcite to undergo recrystallisation, calcite U–Pb geochronology is rarely applicable to Precambrian systems as the calcite U–Pb system invariably does not remain closed over long timescales (Whitehouse and Russell, 1997).
Alternative dating systems involving the radioisotopic decay of rare earth elements (REE) such as Sm–Nd and Lu–Hf, have previously been applied to calcite (e.g. Barker et al., 2009; Maas et al., 2020; Nie et al., 1999; Peng et al., 2003), based on the moderate to strong compatibility of REEs in carbonates in many systems (Debruyne et al., 2016; Elzinga et al., 2002; Terakado and Masuda, 1988; Zhong and Mucci, 1995). Although it should be noted that REE compatibility will be dependent on the conditions of calcite formation, and can vary. Importantly for geochronology, experimental evidence indicates that Lu and Hf are highly immobile in many hydrothermal fluids (Brugger et al., 2016; Migdisov et al., 2016), meaning that the Lu–Hf system is potentially during post-formation processes relative to the U–Pb system. However, concentrations of Lu and Hf are generally low (ppm to ppt range) in calcite, necessitating the dissolution of large quantities of material (up to 2g) per sample for conventional Lu–Hf geochronology (Maas et al., 2020). These large quantities significantly reduce the spatial resolution of the technique and have the additional problem of potential contamination from inclusions. Furthermore, age variation is difficult to detect, and bulk samples may produce meaningless average age derived from mixing of age domains. The dissolution process also removes calcite from its petrological context. The recent development of in-situ Lu–Hf geochronology of individual minerals by LA-ICP-MS/MS allows for rapid acquisition of spatially resolved data, and has been demonstrated for garnet (Ribeiro et al., 2021; Tamblyn et al., 2021) and apatite (Glorie et al., 2021).

In this study, we present the first in-situ Lu–Hf dating of calcite from a variety of geological environments. We demonstrate that in-situ calcite Lu–Hf geochronology can produce meaningful ages for complexly deformed and hydrothermally-altered systems, such as mineral deposits, as well as carbonatite intrusions and low grade metamorphism.

2 Geological Background of Samples

The analysed samples were selected; (1) to demonstrate that calcite Lu–Hf can date primary calcite formation in carbonatites; (2) to reveal the potential of the method to unravel complex ore systems or later events, and; (3) to characterize large calcite samples that would make suitable reference materials for in-situ analysis.

2.1 Phalaborwa Carbonatite, South Africa

The Phalaborwa Igneous Complex is located ~450 km northeast of Johannesburg, in the Limpopo Province, South Africa. The igneous complex is the result of several distinct pulses of alkaline intrusions that were emplaced into Archean granitic gneiss (Staff, 1976). The Loolekop pipe is located in the centre of the Phalaborwa Igneous Complex and was intruded by two episodes of carbonatite emplaced at the intersection of five major faults and shear zones (Basson et al., 2017; Staff, 1976). The oldest carbonatite is termed the “transgressive banded” carbonatite and has an emplacement age of 2060.0 ± 2.2 Ma (baddeleyite SIMS U-Pb; Wu et al., 2011). This is intruded by a slightly younger carbonatite termed the “banded” carbonatite and has an emplacement age of 2059.8 ± 1.3 Ma (baddeleyite SIMS U-Pb; Wu et al., 2011). The Phalaborwa carbonatite is unique as it is the only known example of a carbonatite containing economic Cu mineralisation (Groves and Vielreicher, 2001). In the banded carbonatite-phoscorite, Cu mineralisation is primarily in the form of bornite inter-grown with valleriite with minor chalcopyrite.
(Staff, 1976). In the transgressive carbonatite, Cu mineralisation is present as chalcopyrite inter-grown with cubanite and valleriite (Staff, 1976). Cu mineralisation is interpreted to be magmatic-hydrothermal in origin, with Cu leached by high-temperature hydrothermal fluids at depth, precipitating along fractures within the hosting carbonatite (Le Bras et al., 2021). The sample used in this study (P01) is representative of carbonatite hosted Cu-mineralisation from within the Loolekop pipe (Fig. 1). The sample is mineralogically composed of chalcopyrite inter-grown with cubanite and pyrrhotite alongside an assemblage of magnetite, dolomite, calcite, biotite, pyroxene and valleriite. As the Phalaborwa carbonatite has a well constrained crystallisation age, it provides an ideal case study to demonstrate the utility of the in-situ Lu–Hf method for dating igneous calcite directly associated with Cu mineralisation.

2.2 The Eastern Fold Belt, Mt Isa Block, Queensland, Australia

The Eastern Fold Belt of the Mount Isa Domain has experienced multiple episodes of deformation, magmatism, metamorphism, mineralisation and pervasive hydrothermal alteration across the Paleo- to Mesoproterozoic, and hence represents one of the most metasomatized crustal blocks on Earth (Oliver et al., 2008). Hydrothermal calcite is common across the Mount Isa region, in the Mary Kathleen Domain (Oliver et al., 1993) and in many of the IOCG deposit of the Cloncurry District. For this study we have selected calcite samples from the Lime Creek calcite quarry and the Mt Elliott IOCG deposit for Lu–Hf analysis. The Mt Isa Domain has both regional and deposit level age constraints, making it a good area to demonstrate the technique.

The Lime Creek quarry is one of a number of large calcite pods or veins that are exposed in Mary Kathleen Domain. The Lime Creek quarry is hosted within the ca. 1760 Ma Argylla Formation and lies along the steeply dipping NNW-trending Tribulation-Lime Creek Fault, which offsets regional-scale ‘D2’ folds (Marshall, 2003). Breccias along this fault contain clasts of calc-silicate rocks and metadiorite with a matrix consisting of albite-actinolite-diopside-biotite-titanite-apatite that are subsequently overprinted by the undeformed Lime Creek calcite-dominated veins (Marshall, 2003). These veins are extremely coarse-grained with calcite crystals larger than 1 m$^3$, actinolite crystals over 1 m in length, andapatite, biotite, diopside and titanite grains over 20 cm in diameter (Marshall, 2003; Oliver et al., 1993). Based on cross-cutting relationships, it is interpreted that the Lime Creek vein system, and other calcite pods/veins of this style precipitated post-faulting during late-‘D3’ deformation (ca. 1550–1500 Ma) of the Isan Orogeny (Giles and Nutman, 2002; Marshall, 2003). This style of veinning is common throughout the Mary Kathleen Domain and provides evidence of km-scale fluid transport during late-stage metamorphism (Oliver et al., 1993). Based on C and O isotope analysis of calcite from these veins, they are interpreted to have formed from hydrothermal fluids likely associated with the intrusion of the ca. 1530 to 1500 Ma Williams-Naraku Batholiths (Oliver et al., 1993). Although no direct dating has been completed on the Lime Creek Quarry, titanite from the nearby and cognate Knobby Quarry have produced three titanite U–Pb ages of 1521 ± 5 Ma, 1527 ± 7 Ma, and 1555 ± 5 Ma (Oliver et al., 2004). The sample analysed in this study (LC1) consists of very coarse-grained calcite with coarse-grained diopside collected from a large calcite pod in the Lime Creek Quarry (Fig. 1).
Mt Elliott is an IOCG deposit located in the Eastern Fold Belt of the Mount Isa Inlier (Duncan et al., 2011). The deposit is situated within northwest striking splay of the Mount Dore Fault (Duncan et al., 2011; Wang and Williams, 2001), and is hosted within skarn-altered and deformed phyllites and schists (Garrett, 1992; Wang and Williams, 2001). The host rocks were metamorphosed to lower amphibolite facies during the ca. 1600-1580 Ma ‘D2’ deformation of the Isan Orogeny (Garrett, 1992; Wang and Williams, 2001). The formation of early albite-hematite (red rock) alteration enhanced brittle fracturing and brecciation of the shale (Garrett, 1992). This was infilled by two stages of open-space skarn development: (1) diopside-magnetite-hematite-calcite-titanite-allanite-phlogopite, and: (2) actinolite-scapolite-magnetite-andradite-calcite-epidote-allanite-chlorite and biotite (Garrett, 1992; Wang and Williams, 2001). Sulphides in the second stage include chalcopyrite, pyrrhotite and pyrite (Garrett, 1992). Although the two skarn assemblages are difficult to distinguish mineralogically, the second episode is the most widespread and represents the major Cu–Au event (Garrett, 1992; Wang and Williams, 2001). A variety of geochronological techniques have been applied to constrain the age of Cu–Au mineralisation. The earliest phase of skarn development has been dated at 1530 ± 11 Ma (U–Pb titanite; Duncan et al., 2011). The second stage of skarn development associated with Cu–Au mineralisation has been dated at 1513 ± 5 Ma (molybdenite Re–Os; Duncan et al., 2011) and 1510 ± 3 Ma (actinolite Ar–Ar; Wang and Williams, 2001). Two outcrop samples from the Mount Elliot Cu–Au deposit were selected for Lu–Hf geochronology. Mt Elliott 1 (ME 1) consists of coarse-grained pink-coloured calcite that is coeval with the formation of diopside, scapolite and magnetite (Fig. 1). Although the paragenesis of this sample is relatively unconstrained, the lack of sulphides may indicate that this sample belongs to the early pre-mineralisation skarn assemblage. Calcite from sample Mt Elliott 2 (ME 2) is coeval with the formation of andradite, pyrite, chalcopyrite, pyrrhotite and magnetite (Fig. 1). The close relationship between calcite and chalcopyrite in this sample indicates that it is associated with the main Cu–Au bearing skarn assemblage.

2.3 Flin Flon Volcanic Massive Sulphide (VMS) Deposit, Canada

The Flin Flon Greenstone Belt stretches across central Manitoba through to east central Saskatchewan and hosts several world-class Zn–Cu VMS deposits including the Flin Flon, Callinan and 777 deposits (Koo and Mossman, 1975). Zn–Cu mineralisation is interpreted to have formed contemporaneously with deposition of the 1888.9 ± 1.6 Ma Millrock member during the Trans-Hudson Orogeny (Gibson et al., 2012; Koo and Mossman, 1975; Rayner, 2010). The Flin Flon Zn-Cu orebody is recognised to have undergone six distinct deformation events that have affected the shape of the deposit (Lafrance et al., 2016; Schetselaar et al., 2017). ‘D1’ and ‘D2’ were associated with the intra-oceanic accretion of the Flin Flon Arc to other volcanic terrains before ca. 1872 Ma (Lafrance et al., 2016). ‘D3’ occurred from 1847–1842 Ma as a response to the final accretion of the Flin Flon Terrane to the Glennie Terrane, producing west-verging folds within stacked, east dipping thrust sheets of basement and cover rocks bounded by NNW-striking thrust faults (Lafrance et al., 2016). ‘D4’ resulted from the collision between the Flin Flon and Glennie complex with the Sask Craton and is broadly coeval with the ca. 1840 Ma Phantom Lakes dyke (Gibson et al., 2012; Lafrance et al., 2016). ‘D5’ deformation produced a penetrative regional cleavage (S5) that is
defined by a continuous chloritic foliation ubiquitous in the volcanic basement rocks (Gibson et al., 2012; Lafrance et al., 2016). EWE–WNW directed compression during ‘D₅’ deformation produced a second regional penetrative cleavage and reactivated a variety of regional-scale faults (Gibson et al., 2012; Lafrance et al., 2016). Regional greenschist to granulite facies metamorphism is associated with D₅–₆ deformation at ca. 1820–1790 Ma (Schneider et al., 2007). The Flin Flon mine horizon was imbricated during ‘D₃’ thrusting with the shape of the ore lenses moulded during ‘D₄’ and ‘D₅’ deformation (Schetselaar et al., 2018). Regional greenschist to amphibolite grade metamorphism occurred between 1820–1790 Ma (U–Pb monazite; Schneider et al., 2007), with rocks in the Flin Flon deposit reactivating greenschist facies (Koo and Mossman, 1975). The sample selected for this study is from the hydrothermally altered and sheared footwall of the Flin Flon VMS deposit. This sample is composed of highly foliated chlorite and calcite with disseminated pyrite and residual titanomagnetite. A band of highly foliated calcite was selected for Lu–Hf analysis (sample FF014; Fig. 1) to constrain the age of syn-metamorphic shearing of the deposit.

2.3 Yates U-Th prospect, Otter Lake Area, Grenville Province, Canada

The Otter Lake area is located in SE Ontario within the Grenville Province. The Grenville Province can be distinguished from surrounding provinces based on various structural, metamorphic and isotopic signatures attributed to the overprinting ca. 1080–980 Ma Grenvillian Orogeny (Rivers, 2015). This orogenic event produced widespread metamorphism from granulite to amphibolite facies (Breemen and Corriveau, 2005), accompanied by widespread hydrothermal alteration in the Otter Lake area (Kretz et al., 1999). The Yates U-Th prospect is located approximately 100 km northwest of Ottawa, and is renowned for the occurrence of pegmatites that contain large euhedral crystals of apatite set within a matrix of predominantly orange-pink calcite, with diopside, allanite, titanite, fluorite, thorite and phlogopite (Schumann et al., 2019). A wide range of dates have been produced from the Yates mine; including titanite Pb–Pb and U–Pb ages between ca. 1020 and 998 Ma (Frei et al., 1997; Kennedy et al., 2011); apatite Pb–Pb, and U–Pb ages of 913 ± 7 Ma (Barfod et al., 2005), 933 ± 12 Ma and 920 – 850 Ma (Chew et al., 2011; Xiang et al., 2021) respectively, and an apatite Lu–Hf age of 1031 ± 6 Ma (Barfod et al., 2005). In addition, Simpson et al. (2021a) obtained an in-situ Lu-Hf apatite age of 1000 ± 11 Ma (when corrected for laser induced elemental fractionation). Importantly, the apatite Lu–Hf and Pb–Pb ages have been obtained from the same large apatite crystal, indicating that the Lu–Hf and U–Pb systems have been decoupled (as opposed to multiple generations of apatite growth). Barfod et al. (2005) argued that late-stage fluid interactions may have affected Pb retentivity in the apatite, as the apatite was unlikely to be above the apatite Pb closure temperature at ca. 913 Ma. Calcite from a specimen containing coarse-grained euhedral apatite with pink calcite, quartz and diopside was selected for calcite Lu–Hf analysis (OL-MB, Fig. 1). The apatite is enclosed in the sampled calcite, and is interpreted to have crystallised just prior to the calcite, but during the same hydrothermal event.
3 Method

The samples were mounted in 2.5 cm diameter epoxy mounts and screened for Lu concentration by LA-ICP-MS to determine suitability for Lu–Hf analysis. Mineral liberation analysis (MLA) maps were obtained using a Hitachi SU3800 scanning electron microscope (SEM) to reveal the petrogenetic context of the analysed calcite.

3.1 In-Situ Lu-Hf Dating Method

Analysis was conducted at Adelaide Microscopy, The University of Adelaide. Calcite samples were analysed using a RESOlution 193 nm laser ablation system (Applied Spectra) with a S155 sample chamber (Laurin Technic). The laser ablation system was coupled to an Agilent 8900 tandem mass spectrometer (ICP-MS/MS). The methodology largely follows that of Simpson et al. (2021a) including an initial instrument tune conducted with no NH₃ in the reaction cell to achieve robust plasma conditions (U/Th = 1.00-1.05) and minimal oxide interferences (ThO/Th <0.2%). A carrier gas of 3.5 mL/min N₂ was added after the sample cell in order to increase sensitivity (Hu et al., 2008). Analytical conditions are included in appendix Table 1.

Methods for separation of $^{176}$Hf from $^{176}$Lu and $^{176}$Yb follow that of Simpson et al. (2021a). In more detail, the Agilent 8900x utilises a reaction cell between two quadrupole mass analysers, which can be used to separate isobaric interferences. The first quadrupole is used as a mass filter (e.g., when set to mass 176, only $^{176}$Lu, $^{176}$Yb, and $^{176}$Hf can pass), thereby minimizing potential background interferences and other, unwanted reactions. Following this, a mixture of 10% NH₃ and 90% He is added to the reaction cell (at a rate of 3 mL/min). This mixture is optimized to promote formation of the Hf reaction product $\text{Hf}((\text{NH})_2(\text{NH}_3)_3)^+$ and the second quadrupole is set to 82 amu higher than the first (e.g., Q1 = 176 amu and Q2 = 258 amu). This method minimizes the equivalent Lu and Yb reaction products (~0.03% for Lu, and below detection for Yb), such that the isobaric interferences on $^{176}$Hf are negligible (Simpson et al., 2021a). Lens voltages were tuned to increase sensitivity on the Hf reaction product (Simpson et al., 2021a). In order to calculate Lu/Hf ratios, $^{176}$Hf (+82) was measured directly, $^{175}$Lu was measured as a proxy for $^{176}$Lu, and $^{178}$Hf (+82) measured as a proxy for $^{177}$Hf (Simpson et al., 2021a). $^{176}$Hff/$^{176}$Lu, $^{176}$Lu/$^{177}$Hf, and $^{176}$Hf/$^{177}$Hf ratios were calculated as part of the normalization to NIST610, as opposed to separately converting measured $^{175}$Lu and $^{178}$Hf into $^{176}$Lu and $^{178}$Hf. In more detail, if we assume that the $^{176}$Lu/$^{175}$Lu ratio (or $^{177}$Hf/$^{178}$Hf ratio) is identical between NIST SRM 610 and all analysed samples, a correction factor calculated from the % difference between the $^{178}$Lu/$^{176}$Hf ratio measured in NIST SRM 610 and the published $^{176}$Lu/$^{177}$Hf will correct the unknowns for matrix independent fractionation and differences in isotopic abundance. $^{43}$Ca was measured for internal normalization of trace element abundances, and the following isotopes were measured to monitor for inclusions: $^{27}$Al, $^{47}$Ti, $^{89}$Y, $^{90}$Zr, $^{140}$Ce, and $^{172}$Yb.

Lutetium abundances in most calcite samples are low (< 6 ppm), so we employed a large laser diameter of 257 µm, and a repetition rate of 10 Hz to maximize sensitivity. High sensitivity is important in order to either measure common Hf (in this case $^{178}$Hf), or demonstrate that $^{176}$Hf is sufficiently above detection limits that the effects of common Hf are negligible. Smaller
spot sizes could be employed for higher Lu and/or higher Hf samples. An extra 20 seconds delay was added after each interval of sample ablation in order to ensure the washout had reached background levels. NIST SRM 610 glass ($^{176}\text{Lu}/^{177}\text{Hf}$: 0.1379 ± 0.005, $^{176}\text{Hf}/^{177}\text{Hf}$: 0.282122 ± 0.000009; Nebel et al., 2009) was used as the primary reference material and was analyzed using a spot size of 43 µm. The smaller spot size was required to ensure that $^{175}\text{Lu}$ was measured in pulse counting mode (<4 Mcps). Consistent with observations in Simpson et al. (2021a), Lu and Hf showed no measurable down-hole fractionation in the analysed carbonates (fig. 2), as such, no down-hole correction was applied to the data.

A side effect of the use of large ablation spots is ‘plasma loading’ for which the introduction of a large amount of material reduces the ionizing efficiency of the plasma (Kroslakova and Günther, 2007). Plasma loading was observed in the time resolved signals, with a reduction in signal intensity for all isotopes after ~10 to 15 seconds of ablation. Following this, the signal stabilized after ~18 seconds of ablation (Fig. 2). Importantly, this variation in signal intensity was not observed in the calculated time resolved isotope ratios (Fig. 2), which means that identical ratios were calculated whether this decrease in signal intensity was included or not in the ratio calculation. Importantly, plasma loading can be affected by sample matrix (Kroslakova and Günther, 2007), especially for minerals containing easily ionized elements such as Ca. This necessitates matrix matched calibration, despite the observed lack of down-hole changes in Lu–Hf ratios (Simpson et al., 2021a).

The large ablation volume caused accumulation of ablated material in the tubing and on the interface cones during the first analytical session, which coincided with a decrease in signal intensity over time. Consequently, session 1 records slightly more signal drift compared to session 2. However, there was no measurable corresponding drift observed in the calculated isotopic ratios, apart from a slight decrease in precision due to the lower sensitivity toward the end of the run. Therefore, we recommend that cones are cleaned prior to analysis, and suggest a maximum session duration of approximately 7 hours when using spot diameters of >200 µm. In addition to this, the accumulated material was sometimes mobilized in later analyses, potentially contaminating data. This was observed by increases in Al during the start of ablation that decayed down to background levels. Importantly, similar Al spikes were not observed during background measurement, indicating contamination due to material remobilized during ablation is likely, hence why the additional 20 seconds of washout did not fix this. This contamination did not generally produce a measurable effect on calculated Lu/Hf ratios. However, we stress that this contamination is important to monitor as Hf concentrations are sometimes in the ppt level. As such we recommend close monitoring of signals, particularly Al concentrations, and the removal of 1-3 seconds of each analysis after signal stabilisation if necessary.

3.2 Data Processing

For both LA-ICP-MS and LA-ICP-MS/MS analysis, a stoichiometric Ca concentration of 40.04 wt.% for calcite was used for internal normalization of trace element concentrations. Although the high Ca cps for all analysed samples indicate that they are close to stoichiometric calcite, there may be slight inaccuracies in calculated element concentrations due to major element
substitutions from Mg, Fe, and Mn that are common in carbonates. However, element concentrations were largely used as relative proxies to monitor for inclusions.

Background subtractions, element concentrations and ratio calculations were performed using LADR software (Norris and Danyushevsky, 2018). Where $^{178}\text{Hf}$ was measured above detection limits (~2 ppt for $^{178}\text{Hf}$), common Hf corrections were applied to the data after background subtractions, but prior to normalization to the standard. In more detail, the $^{178}\text{Hf}$ cps measurement for each sampling cycle of the analysis period of each laser spot was used to calculate the common Hf component of the corresponding $^{176}\text{Hf}$ measurement, using the following equation:

$$i^{176}\text{Hf}_r = i^{176}\text{Hf}_m - \left(\frac{i^{176}\text{Hf}}{i^{178}\text{Hf}}\right) c \times i^{178}\text{Hf}_m$$

Where $^{176}\text{Hf}_r = \text{radiogenic }^{176}\text{Hf}$, $^{176}\text{Hf}_m = \text{measured }^{176}\text{Hf}$, $^{178}\text{Hf}_m = \text{the measured }^{177}\text{Hf}$ and $\left(\frac{i^{176}\text{Hf}}{i^{178}\text{Hf}}\right) c = \text{the initial or ‘common’ }^{176}\text{Hf}/^{178}\text{Hf} \text{ ratio. These corrections were applied using an assumed initial }^{176}\text{Hf}/^{178}\text{Hf} \text{ ratio of 0.192 ± 0.004, which is equivalent to a }^{176}\text{Hf}/^{177}\text{Hf} \text{ ratio of 0.2816 ± 0.006. This value is based on the Hf evolution of the crust, with uncertainty that comfortably covers likely natural variation. The uncertainty on the initial }^{176}\text{Hf}/^{178}\text{Hf} \text{ ratio used for the common Hf corrections has been propagated to the final ages, in order to account for any inaccuracies introduced by value used. However, as most analyses have <1% common Hf (Table 1), any inaccuracy related to the initial }^{176}\text{Hf}/^{178}\text{Hf} \text{ ratio is negligible compared to the total uncertainty estimates given Hf isotopes do not vary significantly with time (Fisher and Vervoort, 2018; Vervoort, 2014). Such corrections, however, should be used with caution for samples with higher common Hf, although the dataset presented in this study is not sufficient to determine what an appropriate cutoff should be.}

Subsequent to this correction, isotopic ratios were corrected using an external reference material bracketing approach (commonly used in LA-ICP-MS geochronology), with primary and secondary reference materials interspaced with unknowns through each analytical session. The data was normalized to NIST SRM 610 glass to correct for drift and matrix independent fractionation. The Lu–Hf isotopic ratios published in Nebel et al. (2009) where used for the NIST610 SRM normalisation. Following this, $^{176}\text{Hf}/^{176}\text{Lu}$, $^{176}\text{Lu}/^{177}\text{Hf}$, and $^{176}\text{Lu}/^{176}\text{Hf}$ ratios were corrected to MKED calcite. Although the age of MKED calcite is currently not independently constrained, calcite is interpreted from textural evidence to have formed with the MKED titanite reference material, and therefore the titanite TIMS U–Pb age was used (1517.32 ± 0.32 Ma; Spandler et al., 2016). Further details are outlined in appendix A. This correction method is similar to that used by Roberts et al. (2017) for calcite U–Pb, where the observed analytical offset between the measured and expected Lu–Hf ratio in the standard is applied (as a % correction factor) to the ratios of the unknowns. This offset is inferred to be due to a combination of laser induced (matrix-dependent) elemental fractionation and plasma loading. The uncorrected ages for MKED calcite as well as for ME 1 across four analytical sessions are constant within uncertainty, indicating the age offset is a systematic analytical bias that is applicable.
to the calcite samples of unknown age (Fig. A2). Weighted average ages were calculated using ISOPLOTR (Vermeesch, 2018), using the $^{176}\text{Lu}$ decay constant determined by Söderlund et al. (2004); $0.00001867 \pm 0.00000008 \text{ Myr}^{-1}$

Correct handling of uncertainties in geochronology is important in order to draw accurate conclusions about the resulting ages. As per the recommendations for LA-ICP-MS U-Pb uncertainty propagation in Horstwood et al. (2016), uncertainties are categorised as random, in which case they are propagated to individual analyses, or systematic, in which case they are propagated to the final calculated age. As such, the uncertainties associated with the measurement of the primary standard (NIST SRM 610) have been propagated to the uncertainties of individual analyses. The following systematic uncertainties have been propagated to the final ages: Measurement uncertainty on the secondary standard (MKED C), uncertainty on the titanite U-Pb age used as the reference age for MKED C, uncertainties associated with the $^{176}\text{Lu}$ decay constant, the reference $^{176}\text{Hf}/^{177}\text{Hf}$ ratios for NIST SRM 610. Although for completeness it would be good to propagate uncertainty relating to potential differences in $^{175}\text{Lu}/^{176}\text{Lu}$ and $^{177}\text{Hf}/^{178}\text{Hf}$ between NIST SRM 610 and samples (i.e. natural variation in these ratios), currently there appears to be no data on this. These uncertainties are likely to be negligibly small relative to the overall uncertainty estimates for the analyses. The uncertainty associated with the reference $^{176}\text{Lu}/^{177}\text{Hf}$, $^{176}\text{Lu}/^{176}\text{Hf}$, and $^{176}\text{Hf}/^{176}\text{Lu}$ ratios of NIST SRM 610 are not propagated, as the correction factor associated with NIST610 SRM is cancelled during the correction to MKED calcite (as the NIST610 SRM correction factor is applied equally to MKED calcite and the unknowns samples, and thus becomes redundant). Uncertainty relating to long term reproducibility of the standards has not been propagated, as the standard data for all sessions does not show scatter outside of what would be expected from a single population. More data, however, is required to fully constrain this.

4 Lu-Hf Results

The analysed calcite generally contain $<1\%$ common Hf, apart from sample P01, which contains up to $13\%$ common Hf in individual analyses (Table 1). Consequently, the common Hf corrections are small (or effectively non-existent), and the resultant ages are not significantly affected by the assumed initial $^{176}\text{Hf}/^{177}\text{Hf}$ ratio. Corrected and uncorrected data are included in Supplementary File 1. The inverse isochron and weighted mean single-spot Lu–Hf ages, reported below, are corrected against MKED calcite for matrix-dependant fractionation and common-Hf corrected (where relevant) (Fig. 3). For analyses with inclusions, the signals have been cropped to remove inclusions or, in the case of more significant signal disturbances, excluded from age calculations. Inclusions were detected in the following samples: MKED calcite (6), LC1 (1), P01 (19) and FF014 (6). Excluded data points are included in supplementary file 1. Due to the large number of inclusions, P01 was analysed over two sessions. Data is presented as inverse isochrons (Li and Vermeesch, 2021), and as common Hf corrected weighted average ages (Fig. 3).
5 Discussion

The Phalaborwa carbonatite sample produced a $^{176}$Hf corrected weighted average Lu–Hf age of $2050 \pm 30$ Ma (Fig. 3), consistent with previous baddeleyite U–Pb SIMS ages ($\sim2060$ Ma; Wu et al., 2011). Importantly, the consistency between the calcite Lu–Hf age and existing constraints on carbonatite formation demonstrates that calcite Lu–Hf dating can produce primary age information for early Paleoproterozoic calcite. This result also demonstrates that calcite Lu–Hf geochronology is a viable technique to directly date carbonatite magmatism and associated mineralisation, even in the case of old calcite samples with only $\sim0.5$ ppm Lu.

The weighted average Lu–Hf ages for samples ME 1 and ME 2 are $1538 \pm 9$ Ma and $1504 \pm 13$ Ma, respectively (Fig. 3). The ages of these samples are consistent with the paragenetic timing of alteration at Mt Elliott, providing evidence for calcite precipitation during at least two temporally distinct alteration events. Sample ME 1 is from a coarse calcite-diopside-scapolite-magnetite vein that does not contain sulfides (Fig. 1), the age is, therefore, consistent with formation prior to the major $\sim1510$ Ma Cu–Au mineralisation event (Duncan et al., 2011; Wang and Williams, 2001). In addition, this age overlaps with a titanite U–Pb age from the Mt Elliott deposit ($1530 \pm 11$ Ma; Duncan et al., 2011), and is potentially related to regional Na-Ca alteration between ca. 1555 and ca. 1521 Ma (Oliver et al., 2004). The $1504 \pm 13$ Ma age obtained from sample ME 2 that has an ore stage paragenesis conforms with the $^{207}$Pb/$^{206}$Pb age of cogenetic andradite ($1507 \pm 35$ Ma; appendix B), and overlaps with the ca. 1510 Ma main mineralisation event (Duncan et al., 2011; Wang and Williams, 2001). Additionally, data for ME1 was pooled from all four analytical sessions in order to test reproducibility. Similar to the standard (MKED1; appendix A), ME1 does not show excess scatter between sessions (fig. 3).

Sample LC1, from the Lime Creek quarry, Eastern Fold Belt, Mt Isa Inlier produced an age of $1513 \pm 26$ Ma, consistent with published titanite U–Pb ages ($1521 \pm 5$ Ma $1527 \pm 7$ Ma) from the nearby Knobby Quarry (Oliver et al., 2004). Additionally, this age is consistent with the intrusion of the ca. 1530-1500 Ma Williams-Naraku batholiths, which is interpreted to be the source of the fluids from which the calcite precipitated (Oliver et al., 1993; Page and Sun, 1998). Our results for this sample further demonstrate that calcite Lu–Hf geochronology is an effective technique for constraining the age of calcite mineralisation.

Sample OL-MB from Otter Lake produced a Lu–Hf age of $892 \pm 12$ Ma (Fig. 3). This age is significantly younger than the apatite solution Lu–Hf age of $1030 \pm 6$ Ma (Barfod et al. (2005) and the in-situ apatite Lu–Hf age of $1000 \pm 11$ Ma (Simpson et al. 2021), but is similar to the apatite Pb-Pb age of $913 \pm 7$ Ma (Barfod et al. 2005) and the latest stage of extensional activity on the nearby Bancroft Shear Zone (1045 - 893 Ma, Ar-Ar phlogopite; Cosca et al., 1995). Given the similarity between the ca. 0.9 Ga ages, obtained by different methods, it seems likely that the calcite either grew or records Lu–Hf isotopic resetting during the same event that induced resetting of the apatite Pb–Pb system. The slight difference between the calcite Lu–Hf age
(894 ± 12 Ma) and apatite Pb-Pb age (913 ± 7 Ma) may be due to analytical (i.e. mixing of age domains in the solution Pb-Pb age) rather than geological reasons, particularly given an individual crystal of apatite from the Yates mine produced a U-Pb age range of 920-850 Ma (Xiang et al., 2021). The age difference may also be due to underestimation of uncertainties. Large (~3 cm) apatite crystals such as the one analysed by Barfod et al. (2005) are expected to have Pb closure temperatures of up to 600 °C (Barfod et al., 2005; Krogstad and Walker, 1994), giving a possible upper limit to Lu–Hf closure in calcite. We note that this is significantly higher than the closure temperature of Ar–Ar in phlogopite (ca. 400 °C), indicating that the Otter Lake area potentially had a different thermal history and/or that isotopic resetting in the apatite and calcite was aided by late fluid interactions, as hypothesised by Barfod et al., (2005). As such, further work is required to constrain the Lu–Hf closure temperature in calcite.

The in-situ Lu–Hf age of 1810 ± 18 Ma for the cleavage-hosted calcite vein from the Flin Flon VMS deposit (FF14; Fig. 3), as expected, is younger than the timing of initial mineralisation at the deposit (Koo and Mossman, 1975; Rayner, 2010; Stern et al., 1995). Instead, the age is in excellent agreement with ca. 1820-1790 Ma regional peak greenschist to amphibolite grade metamorphism (Schneider et al., 2007), suggesting the calcite precipitated during metamorphism related to deformation stage ‘D₅’ or ‘D₆’, associated with the final collision between the Flin Flon-Glennie Complex and the Sask Craton (Lafrance et al., 2016). This regional event locally reached maximum greenschist-facies metamorphism (Koo and Mossman, 1975), suggesting the calcite grew under low-grade metamorphic conditions. Sample FF014, therefore, demonstrates that calcite Lu–Hf geochronology has the potential to date low grade metamorphism, which has been difficult using traditional dating methods (e.g. Henrichs et al., 2018).

In summary, we demonstrate that in-situ Lu–Hf geochronology can produce geologically meaningful ages for calcite from a variety of mineralisation styles (e.g. IOCG, carbonatite, and skarn alteration) as well as greenschist-facies metamorphism. The technique also has great potential to date a range of other geological settings and processes (e.g., chemical sedimentation, carbonation reactions) provided calcite contains sufficient Lu for analysis.

5.1 Limitations

The success rate of the in-situ Lu–Hf dating approach in calcite is intrinsically related to; (1) the concentration of Lu, and; (2) the ingrowth time for radiogenic Hf (Fig. 4). Generally, the method is more suitable for REE-rich calcite typically observed in mineral deposits and carbonatites, and/or for Precambrian samples. In addition, the currently available mass-spectrometers require large laser beam diameters (257 µm) for successful calcite Lu–Hf dating, limiting spatial resolution compared to most laser-ablation dating techniques. We note that for high Lu samples, such as ME 1 (or samples that incorporate common Hf), smaller spot sizes are feasible. Additionally, particularly in hydrothermal settings, calcite often forms large, mm to cm scale crystals, reducing the need for small ablation volumes. While individual calcite crystals in other settings can sometimes be < 260 µm, the total amount of calcite is often large enough that aggregates of pure (or close to pure) calcite can be ablated.
Caution should be used with such analysis however, as this may affect laser induced fractionation, individual crystals may be of different ages, and there may be micro inclusions of other minerals.

### 5.2 Advantages of in-situ Lu-Hf dating of calcite compared to other geochronological methods

The Previous dissolution-based Lu–Hf geochronology has produced scattered isochrons, indicative of isotopic disturbances (Maas et al., 2020). While individual data points are significantly less precise than dissolution based methods, the ability to gain spatially resolved data on a much smaller scale (>260 microns), as well as obtain a large number of analyses in a single session can make data interpretation easier (Simpson et al., 2021a). Importantly, trace element data can be obtained simultaneously to interrogate each data point for inclusions or age zonation. Furthermore, calcite Lu–Hf dating can overcome two issues often encountered during U–Pb dating; (1) in contrast to Pb, calcite does not incorporate significant concentrations of common Hf, and; (2) Lu is comparatively resistant to thermal diffusion in calcite (Cherniak, 1998), increasing the likelihood of primary precipitation ages to be preserved. Although it should be acknowledged that fluid mobility and re-crystallisation of the calcite may affect Lu-Hf ages, and are difficult to predict. This opens the possibility that time constraints can be obtained for carbonates from the first three-quarters of Earth history that are generally difficult to date by other methods. Importantly, calcite is commonly associated with ore formation, meaning in-situ Lu–Hf dating affords the possibility to directly constrain the age of mineralising events and the temporal evolution of mineral deposit systems.

From our work, we suggest samples ME1 and OL-MB calcite could be developed as primary reference materials due to being (1) common-Hf free, (2) homogenous in age across crystals up ~1 cm in size, and (3) available in large quantities. We aim to characterize such reference materials and make them available to the wider geochronology community.

### 6 Conclusions and Future Directions

Calcite is among the most common of rock-forming minerals, meaning that in-situ Lu–Hf geochronology of calcite has enormous potential to constrain the age of formation and/or alteration of a range of igneous, sedimentary, metamorphic, and hydrothermal rock systems, including rock-types that are considered very difficult to date (e.g., marbles). This technique has particular application to mineral deposits as it allows for the ability to constrain the age of pre-ore, ore-stage and post-ore events (e.g., Fig. 3). Furthermore, given the successful dating of old (~2 Ga) calcite with <1 ppm Lu (e.g., sample P01; Table 1), this technique has the potential to date old calcite from a variety of settings with relatively low HREE concentrations. In-situ Lu–Hf dating of calcite can be regarded as a complimentary, and in some cases alternative, technique to carbonate U–Pb dating, where Lu–Hf dating is well suited for older samples, or to obtain primary precipitation ages for systems affected by Pb mobility. Coupling in-situ Lu–Hf dating with other isotopic systems (U-Th-Pb, C, O, Sr, Nd) may be particularly powerful for constraining the origin, nature and redox conditions of the fluids or melts from which the calcite precipitated.

### Author contributions

Acknowledgments
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Figures and Tables
Figure 1: A combination of SEM mineral maps (A,C,D,E) and photos of analysed samples. (A) P01 (Phalaborwa Carbonatite) shows calcite in petrogenetic context to chalcopyrite. (B) Photo of LC 1 hand sample. (C) Photo of the ME 1 sample in outcrop, with inset showing the mineralogy of the analysed sample. (D) calcite from ME 2 (Mt Isa) in contact with hematite, pyrrhotite and andradite, with inset showing hand sample (D) OL-MB (Otter Lake), showing analysed calcite with associated minerals, with inset showing relationship between apatite (Ap) and calcite (Cal) in hand sample. (E) FF014 (Flin Flon deposit) shows calcite vein in chlorite matrix with disseminated pyrite, with inset showing analysed block (dark coloured matrix is composed of chlorite). Black circles represent laser spot locations. Mineral abbreviations: Cal: Calcite, Cpy: chalcopyrite, Py: pyrite, Mag: magnetite, Cu: cubanite, Di: diopside, Scp: scapolite, An: andradite. Larger size sample images are included in appendix C.

Figure 2: Time resolved signals for $^{175}\text{Lu}/^{(176)}\text{Hf (+82)}$, $^{43}\text{Ca}$, $^{175}\text{Lu}$, and $^{(176)}\text{Hf (+82)}$ demonstrating the effects of plasma loading on the signal intensities (i.e. dip in signal intensities at ~10-15s ablation), but not for the $^{176}\text{Hf}/^{175}\text{Lu}$ ratio, which remains constant downhole. The time resolved intensity of each analyte has been offset in the graph for better comparison, therefore the Y axis scale is not continuous. Green horizontal lines show the scale for $^{176}\text{Hf (+82)}$, blue horizontal lines show the scale for $^{175}\text{Lu}$, red horizontal lines show the scale for $^{43}\text{Ca}$, and black horizontal lines show the scale for the $^{176}\text{Hf (+82)}/^{175}\text{Lu}$ ratio. Presented data is from an analysis of MKED calcite.
Figure 3: Anchored inverse isochron and weighted average ‘single spot’ ages for analysed samples, corrected for matrix-induced fractionation against MKED1 calcite. Isochrons have been anchored to an initial $^{177}\text{Hf}/^{176}\text{Hf}$ ratio of $3.55 \pm 0.07$. Ellipses represent data points and $2\sigma$ uncertainty. Uncoloured ellipses represent data points excluded from the regression calculation. Weighted average ages are corrected for common-Hf where relevant (see Table 1 and text). Blue bars represent $2\sigma$ uncertainties. Black lines represent weighted average ages, with grey boxes representing the 95% confidence interval uncertainty.

![Figure 3: Anchored inverse isochron and weighted average ‘single spot’ ages for analysed samples, corrected for matrix-induced fractionation against MKED1 calcite.](image)

Figure 4: Lu ppm vs $2\sigma$ uncertainty for each calcite analysis. The grey curve shows a function fitted to the data from samples with ages between 1500 and 1540 Ma (samples ME1, ME2, and MKED, with symbols outlined in black). Only data points with similar ages were used to construct this guiding curve as the obtained precision is age-dependant. The Lu-Hf ages for older samples (e.g. P01 and FF014) are more precise relative to younger samples for a given Lu concentration (assuming no common Hf). Note: MKED is the calcite Lu-Hf standard used to correct the analysed samples. All data for MKED is included in supplementary file 1.

![Figure 4: Lu ppm vs $2\sigma$ uncertainty for each calcite analysis.](image)
TABLE 1: AGES AND LU AND HF CONCENTRATION INFORMATION FOR THE ANALYSED SAMPLES.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Age (Ma)</th>
<th>95% CI</th>
<th>n</th>
<th>Min. % Hf corr</th>
<th>Max. % Hf corr</th>
<th>Min Lu ppm</th>
<th>Max Lu ppm</th>
<th>Min Hf* ppb</th>
<th>Av Hf* ppb</th>
<th>Max Hf* ppb</th>
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<tr>
<td>P01</td>
<td>2054</td>
<td>1.5%</td>
<td>36</td>
<td>0%</td>
<td>2.65%</td>
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<td>270</td>
<td>505</td>
<td>0.03</td>
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<td>LC 1</td>
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<td>19</td>
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<td>0.46%</td>
<td>1.25%</td>
<td>1900</td>
<td>1600</td>
<td>3600</td>
<td>0.011</td>
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<tr>
<td>ME 1</td>
<td>1540</td>
<td>0.6%</td>
<td>79</td>
<td>0%</td>
<td>0.28%</td>
<td>2.4%</td>
<td>4300</td>
<td>5325</td>
<td>6300</td>
<td>0.002</td>
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<tr>
<td>ME 2</td>
<td>1500</td>
<td>0.9%</td>
<td>29</td>
<td>0.16%</td>
<td>0.58%</td>
<td>3.04%</td>
<td>700</td>
<td>3159</td>
<td>5500</td>
<td>0.002</td>
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<tr>
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<td>30</td>
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Note: 95% CI refers to the 95% confidence interval uncertainty on the calculated age. n refers to the number of analyses used for the age calculation. % Hf corr refers to the average % decrease in age due to the common Hf correction. Hf* concentrations have been calculated from $^{178}$Hf and assume no radiogenic ingrowth of $^{176}$Hf and thus represent the ‘common’ Hf concentration for each sample.

Appendix A: MKED calcite sample description

A sample of orange-pink calcite associated with the MKED1 titanite U-Pb standard (1517.32 ± 0.32 Ma, U-Pb TIMS; Spandler et al., 2016) was analysed as a matrix matched secondary standard in order to correct unknown samples for matrix related analytical offsets, such as laser induced elemental fractionation and plasma loading effects. The calcite was sampled from the same drill core from which the titanite standard was taken (full details can be found in Spandler et al., 2016). The sample consists of massive calcite surrounding large (~8cm) euhedral titanite crystals. The titanite is interpreted to have grown in the same fluid as the calcite, but just prior to calcite crystallization. The average age across all 4 analytic sessions is 1560 ± 10 Ma (fig. A2), suggesting that matrix fractionation during laser ablation produces ages that are systematically approximately 3% too old.
Figure A1: images of MKED calcite. A shows calcite chip from where the analysed sample was taken. B shows underside of the same chip, where a large titanite crystal has been removed. Red box shows remnant fragments of titanite.
Figure A2: Demonstration of the systematic analytical offset observed for calcite Lu-Hf ages. Green rectangles are 95% confidence intervals around weighted mean ages for each session, with session number in the bottom right corner of each rectangle. Horizontal black line shows the weighted average age of all analytical sessions, with grey rectangle showing 95% confidence interval uncertainty. The combined weighted average age for all analytical sessions is shown in the top right corner. The expected age is from Spandler et al. (2016). Weighted mean ages were calculated using ISOPLOTR (Vermeesch, 2018).

Appendix B: Mt Elliott Andradite U-Pb data:

Cogenetic andradite was analysed from the Mt Elliott 2 calcite sample (fig. 1). The sample was analysed using the same laser system as used for Lu-Hf analysis, but coupled with an Agilent 7900 quadrupole mass spectrometer. As the University of Adelaide does not currently possess an andradite U-Pb standard, U-Pb and Pb-Pb ratios were corrected to NIST610 SRM, using ratios from (REF). A large aspect ratio ablation spot (120 microns in diameter, drilling approximately 30 microns deep)
was used to minimise the effects of downhole fractionation (REF – Sylvester?) however, it is possible that calculate U-Pb ages are inaccurate due to the lack of matrix matched primary standard. As the data appears to be concordant, however, a weighted average age can be calculated from the $^{207}\text{Pb}/^{206}\text{Pb}$ ratios (fig. B1), which should not be significantly affected by laser induced matrix fractionation (REF). As such, the calculated age is considered accurate within uncertainty.

Figure B1: U-Pb age of ME 2 andradite. Left panel shows the weighted average $^{207}\text{Pb}/^{206}\text{Pb}$ age and the right panel shows the concordia age on a Tera-Wasserburg Concordia plot. For the weighted average, vertical rectangles are $2\sigma$ uncertainties around calculated single spot ages, with the black bar showing calculated weighted mean age, and the grey rectangle showing associated 95% confidence interval uncertainty. For the concordia plot, each ellipse shows the $2\sigma$ uncertainty around each analysis, with white ellipse representing 95% confidence interval uncertainty around the calculated concordia age. Weighted mean age and concordia age were calculated using ISOPLOTR (Vermeesch, P., 2018).

Appendix C: Large sample images

The following are larger versions of the sample images from figure 1. Mineral abbreviations are: Cal: Calcite, Cpy: chalcopyrite, Py: pyrite, Mag: magnetite, Cu: cubanite, Di: diopside, Scp: scapolite, An: andradite
Figure C1: Large image of sample P01 from Phalaborwa carbonatite, South Africa. Large images shows SEM mineral map. Inset shows hand sample photo. Cpy: Chalcopyrite, Cu: Cubanite, Cal: calcite, Mag: Magnetite.
Figure C2: Large image of sample LC1 from Lime Creek, Mt Isa region, Australia. Image shows hand sample. Di: diopside, Cal: calcite.
Figure C3: Large image of sample ME1 from Mt Elliott, Mt Isa region, Australia. Large images shows sample location and inset shows hand sample. Di: diopside, Scp: scapolite, Cal: calcite, Mag: magnetite
Figure C4: Large image of sample ME2 from Mt Elliott, Mt Isa region, Australia. Large images shows SEM mineral map with black circles showing laser spot locations. Inset shows hand sample photo. Cpy: Chalcopyrite, Py: pyrite, Cal: calcite, An: andradite.
Figure C5: Large image of sample OL-MB from the Yates mine, Canada. Large images shows SEM mineral map with black circles showing laser spot locations. Inset shows hand sample photo. Ap: apatite, Cal: calcite.
Figure C6: Large image of sample FF014 from Flin Flon, Manitoba and Saskatchewan, Canada. Large images shows SEM mineral map with black circles showing laser spot locations. Inset shows hand sample photo. Py: pyrite, Cal: calcite.
### APPENDIX TABLE 1: ANALYSIS AND LA-ICP-MS/MS TUNING PARAMETERS

#### plasma parameters

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#### Q1 parameters

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