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Amphibole control on copper systematics in arcs: Insights from the analysis of global datasets

Nicholas D. Barber a,*, Marie Edmonds a, Frances Jenner b, Andreas Audétat c, Helen Williams a

a Department of Earth Sciences, University of Cambridge, United Kingdom
b School of Environment, Earth, and Ecosystem Sciences, The Open University, United Kingdom
c Bayerisches Geoinstitut, Universität Bayreuth, Germany

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Abstract

Copper, obtained from porphyry deposits formed in arc settings, is a critical resource, and is primarily sourced from magmas. However, the processes that shape the copper contents of arc magmas are up for debate. Existing models place emphasis on different petrological agents that explain large-scale trends in copper systematics. Previous studies have noted the ‘Cu paradox,’ where the magmas with high Sr/Y ratios, indicative of ore-forming potential, have the lowest copper concentrations. Here we compile a multidimensional database of volcanic whole rock compositions and couple it with simple petrological models to elucidate the controls on volcanic whole rock compositions with respect to Cu. We show that calc-alkaline, high Sr/Y magmas undergo major element modification caused by extensive amphibole and/or garnet fractionation, which promotes sulphide precipitation and copper depletion. We demonstrate the importance of amphibole fractionation as a globally important process that promotes both calc-alkaline differentiation and sulphide fractionation in arc magmas, as well as its role in signalling the right set of chemical conditions in magmas that ultimately feed copper porphyry deposits. This work also raises the possibility of amphibole as a geochemical and petrological indicator of potential porphyry-forming conditions in a magma, which we show should be readily detectable by a combination of different geochemical metrics. Despite their paucity in copper, high Sr/Y magmas are associated with porphyry deposits, implying that the propensity of magmas to form such deposits depends on factors other than a magma’s bulk copper content.

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Keywords: Copper systematics; Global datasets; Amphibole; Sulphide stability; GIS; Igneous differentiation; Subduction zones; Porphyry deposits

1. INTRODUCTION

Copper (Cu) is economically important owing to its role in the development of electrical components and its critical status in the transition to green energy (Arndt et al., 2017). Porphyry deposits, which are temporally and spatially associated with arc magmatism (Fig. 1), account for over 70% of global Cu ore production (Singer et al., 2005), and significant amounts of Au and Mo production (Singer et al., 2005). Great progress has been made in developing a general model of porphyry development (Richards, 2015; Chiaradia and Caricchi, 2017; Blundy et al., 2015; Wilkinson, 2013; Lee and Tang, 2020; Richards, 2003; Richards, 2011), yet more work needs to be done to understand how these processes are expressed in modern volcanic environments (Hedenquist et al., 1994). Prevailing magmatic models of Cu porphyry formation focus on two
important processes, which may promote Cu enrichment, transport, and deposition in and around porphyry stocks: (i) the saturation of the magma in sulphide (an Fe and S-bearing liquid or mineral phase), into which Cu partitions strongly (Kiseeva and Wood, 2013; Kiseeva et al., 2017; Patten et al., 2013; Jugo et al., 1999; Simon et al., 2006; Li and Audetat, 2015; Li and Audetat, 2013), a process which may deplete the magma Cu when sulphide fractionates, but may also enrich a magma if sulphide is remobilised by hotter or more oxidised magma (Wilkinson, 2013; Park et al., 2019; Jenner et al., 2010; Chiaradia, 2014); and (ii) the exsolution of a volatile phase, which may unmix to a brine and vapor at low pressure (Hedenquist et al., 1994), into which Cu and other metals partition and eventually precipitate from, leading to ore deposition (Edmonds and Mather, 2017; Blundy et al., 2015; Nadeau et al., 2010; Audetat et al., 2008).

A number of models have been proposed to explain how these processes may enhance a magma’s potential to eventually go on and form a mineralized deposit. Recently, it has been suggested that sulphide saturation may be avoided by simultaneous Fe depletion and auto-oxidation caused by garnet fractionation (Lee and Tang, 2020). Garnet’s preference for Fe$^{2+}$ leaves residual magmas enriched in Fe$^{3+}$ (Lee and Tang, 2020). This in turn produces a more oxidized and evolved melt (with higher $S^{6+}$/$S^{2-}$), which pushes the magma further away from sulphide saturation (Jugo et al., 2010) and may even trigger the dissolution of existing sulphides. It was proposed that this garnet-mediated process could enrich the melt in Cu, which could then be transferred to fluids when porphyry formation commences (Lee and Tang, 2020). Alternatively, others have argued that sulphide accumulation may be critical to later ore development by pre-concentrating a reservoir of localized sulphides at the base of a magma reservoir (Wilkinson, 2013), which is later remobilised by intruding hotter or more oxidised magmas, thereby enriching these magmas in Cu (Wilkinson, 2013).

However, other studies have de-emphasized the importance of magmatic Cu contents as an indicator of ore potential (Richards, 2015; Du and Audetat, 2020), promoting instead the importance of magmatic water content (Richards, 2011; Loucks, 2014; Rezeau and Jagoutz, 2020), time scales of magma differentiation (Chelle-Michou et al., 2017), and larger magma volumes (Steinberger et al., 2013) as some of the critical controls on the capacity of a magma to produce a porphyry. Nonetheless, nearly all models agree that the process of sulphide saturation is important for understanding eventual ore formation. Sulphide saturation is a complex process mediated by temperature, pressure, and silicate melt composition, and thanks to continued empirical work the factors leading to sulphide saturation in silicate melts are increasingly well understood (O’Neill, 2020; Smythe et al., 2017; Jugo et al., 2010; Kiseeva and Wood, 2015; Zajacz and Tsay, 2019; Nash et al., 2019). However, it is presently unclear (i) what petrological processes in arc magmas will promote, delay, or are simply associated with sulphide fractionation, and (ii) whether the early fractionation of a sulphide, and associated loss of Cu, is detrimental to later porphyry formation (Park et al., 2019).

One reason that these outstanding questions remain is the measure of sample bias in our existing understanding of Cu systematics. Crucially, we need to establish what generic petrological processes, if any, can explain the abundances of Cu in arc magmas in all major subduction zones as a function of sulphide stability. In order to address this need, we compiled a large global database of volcanic arc whole rock compositions (Fig. 1). The ArcMetals database ($N = 55,795$) contains data from 17 arcs, encompassing geochemical and contextual information such as major.
trace element and radiogenic isotope compositions, geology, location, and geologic age (see Methods), expanding on the approach taken by earlier studies (Richards, 2015; Chiaradia, 2014; Chiaradia, 2015; Farner and Lee, 2017; Lee and Tang, 2020; Rezeau and Jagoutz, 2020). Combined, these parameters allow us to explore Cu systematics in all arc settings. We interpret the generic features of ArcMetals with respect to Cu systematics using models of silicate melt differentiation (Ulmer et al., 2018) and sulphide saturation (O’Neill, 2020) based on recent high-quality experimental studies. This work pushes forward our understanding of key magmatic processes occurring in arcs that influence the capacity of magmas to develop ore deposits. Furthermore, this work provides a framework for further interrogation of major mineral controls on chalcophile metal behaviour in specific volcanic arcs.

2. METHODS

2.1. ArcMetals: data sources and compilation

2.1.1. Database design

This paper presents a new compilation of existing arc volcanic whole rock chemistry called ArcMetals (https://github.com/ndb38/arc-metals). This database was compiled with several crucial design distinctions in mind which distinguishes it from previous databases. First, we wanted the database to be fully integrated with the spatial dimension of the data. Hence, much of the compilation work takes place in a Geographic Information System (GIS) environment, where we can control and append geophysical and tectonic datasets. As discussed in greater detail both below and in the Supplement, this approach afford us several advantages over prior compilations. Here we take advantage of recent advances in the application of data science and Geographic Information Systems (GIS) in geochemistry, allowing us to build on prior studies that were based on simpler data compilation routines (Lee and Tang, 2020; Farner and Lee, 2017; Chiaradia, 2014; Chiaradia, 2015; Loucks, 2014; Richards, 2015) in the initial compilation and filtration of GeoRoc data, combining existing petrological datasets in ore systematic ways. Subsequently, we apply a range of geospatial techniques to append the maximum amount of geophysical data to this compilation without compromising the extent or quality of this data. This latter step is documented in detail in the Supplement, and is what sets this compilation apart from many previous datasets.

2.1.2. Geochemistry

Sample geochemical and analytical data were collected from the GeoRoc (Sarbas and Nohl, 2008) database. These data were compiled using open source python code, available on our GitHub. Initially, 19 arc magma datasets were included in the database, but the Kermadec and Banda files contained so few data upon filtering, that they were ultimately omitted (Fig. 1). Data mainly for arc volcanic rocks were compiled (see Supplement), but a companion plutonic dataset has also been generated (though not discussed in this study). Before filtering, the fully compiled database contained >200,000 records. In order to maximize the number of measurements per sample, six filters were applied to the initial compilation: (1) records with data obtained before 1960 C.E. were removed; (2) records with no recorded analytical technique were removed; (3) only those records pertaining to measurements by X-ray fluorescence (XRF), secondary ion mass spectrometry (SIMS), electron microprobe (EPMA), thermal ionization mass spectrometry (TIMS), inductively-coupled plasma mass spectrometry (ICP MS), laser ablation inductively-coupled plasma mass spectrometry (LA ICP MS), and Fourier-transform infrared (FTIR) spectroscopy were retained; (4) the database was reduced to individual records where sample name, material type (whole rock, glass, or inclusion), and analytical technique were the same (e.g. if 1 whole rock sample had 4 records in GeoRoc measured using XRF, this filter would reduce the 4 records to one average for XRF); (5) records with the same sample name and material type were averaged and collapsed into one record. This had the effect of, for example, combining a sample’s ICP MS measured trace elements with its XRF derived major elements; (6) samples that had the same element measured more than once using the same technique were removed. This filter only affected a small subset (a few hundred) samples, but having it in place makes it easier to quantify analytical errors (see Supplement). Before plotting, the final database was filtered to only include those magmas with a reported loss on ignition (LOI) less than 3.5 wt.%, following approaches used in similar databases (e.g. Loucks, 2014).

2.1.3. Geophysical parameters

Several global geophysical datasets were appended to the main database using the geospatial software QGIS 3.14. The data we appended included subducting slab surfaces and geometry generated from extensive seismic records (slab dip, depth to slab, slab thickness) (Hayes et al., 2018), crustal thickness (Szwillus et al., 2019), and subducting plate sediment cover thickness (Pasyanos et al., 2014). Every sample record in the database was linked to the geophysical datasets, which have good global coverage at a resolution of 10–100 km². A sub-population of database records had additional geophysical data appended based on their proximity to Holocene volcanoes (Syracuse and Abers, 2006). These data included convergence rate, slab thermal parameter, and slab age (Syracuse and Abers, 2006). The full QGIS methods and compilation scripts can be found in the Supplement.

3. RESULTS

We present summaries of global volcanic whole rock geochemical data in Figs. 2, 3, 4, 5 and 9. These are discussed and interpreted in Section 4, in light of modeling done to validate these trends (Figs. 6–8). Fig. 2a shows volcanic whole rock Sr/Y versus SiO₂, colour-coded for the different arcs (Fig. 2b), Cu content (Fig. 2c) and crustal thickness (Fig. 2d). It has been shown that magmas fertile for porphyry Cu deposit formation have high whole rock Sr/Y ratios at intermediate to felsic magma compositions (Fig. 2, Richards, 2011; Loucks, 2014; Chiaradia, 2015).
The Sr/Y ratio, which compares the Large Ion Lithophile Element (LILE) Sr to the high field strength element (HFSE) Y, is widely regarded as a proxy for high pressure fractionation of hydrous arc magmas (Richards, 2011; Chiaradia, 2014; Loucks, 2014). Strontium abundances during fractionation are primarily controlled by plagioclase (Ewart and Griffin, 1994), whereas Y abundances are controlled primarily by amphibole and garnet, as well as some minor phases like titanite (Green, 1994). The ratio of plagioclase to amphibole crystallized in a fractionating arc magma is decreased under conditions of high H$_2$O activity (Grove et al., 2002; Stolper, 1982), which simultaneously stabilizes amphibole phases which incorporate H$_2$O into their structure (Foden and Green, 1992). Thus, a hydrous magma should see abundant amphibole fractionation early in its differentiation in the mid to deep crust (up to 40 km., (Ridolfi et al., 2010; Santana et al., 2020; Alonso-Perez et al., 2009) and late-stage plagioclase crystallization at or near volatile saturation in the upper crust (Loucks, 2014). This fractionation sequence will result in an elevated Sr/Y ratio in deeply derived andesitic to dacitic magmas with significant amphibole (Loucks, 2014).

Fig. 2. Sr/Y vs. SiO$_2$ plots, colored for different features. Plots (b), (c), and (d) are both sub-sampled to only display 300 (b) and 1000 (c) and (d) samples for visual clarity. The black line in plot (a) differentiates “high” from “low” ore potential in arc magmas, as defined in the literature (Loucks, 2014). Magmas sitting above the black line have a greater association with ore-forming deposits. Plot (a) is colored by the density of points in the total dataset, and contours for sample density are included in all subsequent plots. Plot (b) shows a sub-sample of arc magmas colored and symbolized by arc; notice how ore-producing arcs (e.g. the Andes) are the only ones that tend to proliferate above the high ore potential line. Plot (c) shows a sub-sample of the database colored for Cu, where high and low ore potential magmas have mean [Cu] of 42 and 60 ppm respectively. Similarly, plot (d) shows that high ore potential magmas have thicker crust, on average, than low ore potential magmas.

Fig. 3 shows MgO versus total FeO plots for global volcanic arc whole rocks, colour-coded for Cu contents (Fig. 3a and c), Sr/Y (Fig. 3b) and crustal thickness (CT; Fig. 3d). These plots show that whole rocks with high Cu contents lie along a tholeiitic trend (with Fe-enrichment), whereas those whole rocks that lie along the calc-alkaline
trajectory (with Fe depletion) are copper-poor. These calcalkaline rocks also have the highest Sr/Y (Loucks, 2014). These observations are consistent with previous work (Richards, 2015; Chiaradia, 2014; Lee et al., 2012; Lee and Tang, 2020; Jenner, 2017).

We identify the principal processes responsible for the geochemical trends shown in Figs. 2 and 3 using rare earth element (REE) concentrations (Figs. 4 and 5). Fig. 4 shows data sorted so that the highest [Cu] and Sr/Y points are placed "on top" of the data cloud to ensure the reader is not missing any of the highest [Cu] and Sr/Y points. However, there are many more low [Cu] and Sr/Y points, as shown in the Supplement via 3D scatter plots set up using the same axes and plot design (Fig. S10). The schematics in Figs. 4 and 5 show vectors for the fractionation of garnet, amphibole, olivine, plagioclase, orthopyroxene and clinopyroxene using an index of REE plot curvature, Dy/Dy* (Davidson et al., 2013). Essentially, Dy/Dy* estimates the relative depletion of the middle rare earth (MREE) Dy in relation to its light (LREE) and heavy (HREE) counterparts. The Dy/Dy* metric makes a weighted determination of the slope and shape of an REE spider diagram with respect to Dy, as:

$$\frac{Dy}{Dy^*} = \frac{Dy_N}{La_N^{4/13} + Yb_N^{7/13}}$$

Dy/Dy* is of particular use for tracking amphibole/cpx and garnet fractionation (Davidson et al., 2013; Lee and Tang, 2020; Chen et al., 2019; Cox et al., 2019) in rock suites. Additionally, the trajectories for melting in the garnet source field are plotted following (Davidson et al., 2013). The ratio Dy/Dy* tends to be lowered by amphibole and clinopyroxene fractionation. These phases will deplete Dy relative to Yb. Olivine, plagioclase, and orthopyroxene will drive Dy/Dy* towards higher values, as these phases do not incorporate Dy into their structure and thus Dy will be enhanced relative to light (LREE) and heavy (HREE) rare earth elements. Garnet fractionation will move Dy/Yb to higher values during fractionation (i.e. deplete Yb relative to Dy) while simultaneously increasing Dy/Dy*. Mantle melting in the presence of garnet will lead to more moderate values of both Dy/Yb and Dy/Dy* if a garnet rich source is extensively melted.

The REE systematics of the global database can be further explored using a statistical approach (O’Neill, 2016), which compares parameters describing the shape of chondrite-normalized multi-REE plots (Fig. 5). The REE polynomials, symbolized by $\lambda$, describe the shape of REE curves (O’Neill, 2016) based on multivariate statistics across all of the REEs (O’Neill, 2016). The polynomials are determined from the following calculation in orthogonal form:

$$\ln([\text{REE}]/[\text{REE}]_{\text{Ch}}) = \lambda_0 + \lambda_1 f_1^{\text{orth}} + \lambda_2 f_2^{\text{orth}} + \ldots$$

Where the $f$ variables represents polynomials of REE atomic radius ($r_{\text{REE}}$), chosen to avoid co-correlation of the
4. DISCUSSION

4.1. Geochemical Characteristics of High Sr/Y Magmas

We follow the example of Loucks (2014), in recognizing the close association between high Sr/Y magmas and porphyry mineralization (Fig. 2a (Loucks, 2014)). Following this approach, this work shows that high Sr/Y magmas show an association with continental arcs such as Mexico, the Andes and the Cascades (Fig. 2b and Supplementary Material), a low mean whole rock Cu concentration (< 50 ppm, Fig. 2c) and thicker crust (mean of ~40 km, Fig. 2d); these features are consistent with previous studies (Chiaradia, 2014; Chiaradia, 2015; Richards, 2015). These same magmas also sit in the high Dy/Yb, low Dy/Dy* quadrant of Fig. 4, and the high $k_2$, high $k_1$ quadrant of Fig. 5. To confirm whether the mean Cu and crustal thickness differences in Fig. 2 between high and low Sr/Y magmas is statistically robust, these values were compared using a two-way analysis of variance (ANOVA) hypothesis test, and a subsequent Tukey’s highly significant difference (HSD) test. The null hypothesis tested in all cases was that the mean of a given measure is the same between two groups. The likelihood that this hypothesis is true due to random chance is calculated using an F statistic, given by:
Fig. 5. REE behaviour as described by λ spider-plot shape parameters O’Neill (2016). Top panels the right-hand panel is a schematic, showing how λ1 vs. λ2 plots describe mineralogical controls on REEs during differentiation, following O’Neill (2016). While λ1 describes slope, it is calculated according to the radius of ordered REEs. Hence, a negative λ1 corresponds to a positively sloped REE spider profile. (a) and (b) plot λ1 vs. λ2 colored for Cu and Sr/Y respectively. Grey points show the entire global database. Colored points are those that plot in the high ore potential field of Fig. 3. Mean [Cu] of high Sr/Y field given in a). Empirical results from Ulmer et al. (2018) are shown in the orange shaded area in Figure (a), where the arrow denotes the evolving REE contents of empirical products in λ space. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 6. Compilation of melt compositions as compared to the crystallization time and chemistry of amphibole in the empirical products analyzed in this study (Ulmer et al., 2018). Plot (a) shows the melt Mg# as a function of temperature - amphibole stabilizes at 1050 °C for both FC ba AuPd and FC Mb AuPd. Plot (b) shows the FeO content of amphibole plotted against the FeO content of the corresponding silicate melt. Panel (c) shows the amphibole’s Mg# plotted against silicate melt FeO.

The supplement for information on the applicability of this t-test, which uses the t-statistic rather than the F for hypothesis testing, but is applied much the same way as an ANOVA. After determining the p-value, which is a simple but easily misinterpreted measure of the likelihood of difference between the means occurring due to random chance, the difference between the different treatments (e.g. different arcs, different Sr/Y groups) is compared using a Tukey HSD test (see method: https://www.statsmodels.org/stable/generated/statsmodels.stats.multicomp.pairwise_tukeyhsd.html), which calculates the following test statistic:

\[ q_s = \frac{Y_s - Y_R}{SE} \]
where $Y_A$ is the larger of the two means, $Y_B$ the smaller, and SE the standard error of the sum of the means. See the Supplement for detailed plots comparing the test statistics, tables with statistical outputs and constraints, and the code used in these analyses. The mean (1) Cu and (2) crustal thickness of the high and low Sr/Y groups in Figs. 2 c and d have been compared, respectively, using t-tests (high or low Sr/Y) t-tests, and these differences have been shown to be statistically significant ($p < 0.005$). Additional examples of statistical analyses on ArcMetals are shown in the example code.

The low mean Cu concentrations ([Cu]) in whole rocks associated with higher Sr/Y (Fig. 2c) highlights the so-called “Cu-paradox” (Lee and Tang, 2020), where Cu is present in low abundance in the magmas that appear to be most capable of forming ore deposits. Observations such as these have been used to support porphyry formation models where crystallization of sulphide removes Cu from the silicate melts, to be later remobilized by one of several petrological processes (Lee and Tang, 2020; Wilkinson, 2013; Jenner et al., 2010; Park et al., 2019). However, it is also possible that melt [Cu] depletion may have little bearing on whether a magma goes on to form an ore deposit (Richards, 2015; Du and Audetat, 2020). While this initial analysis confirms the findings of prior studies that magmatic [Cu] is significantly lower in high Sr/Y magmas on a global scale (Richards, 2015; Lee and Tang, 2020; Chiaradia, 2014), the petrological processes driving this association have not yet been resolved. To address this, empirical datasets are combined with simple trace element partitioning and sulphide saturation models to better understand what petrological processes are associated with high Sr/Y and low Cu in magmas.

4.2. Amphibole vs. Garnet Signatures

Globally, it can be seen that both low Cu (Fig. 3a) and high Sr/Y (Fig. 3b) volcanic whole rocks follow a calc-alkaline path, showing consistent Fe loss with decreasing [MgO] (paralleling the high Sr/Y ellipse in Fig. 3d). Similar
is both HREE depletion and overall enrichment in the LREE (Fig. 4b). Fig. 4b shows a strong preference for high Sr/Y magmas to sit in the bottom right "garnet-influenced" quadrant (high Dy/Yb, low Dy/Dy*), and Fig. 5a and b show many high Sr/Y whole rocks sitting near the amphibole fractionation and garnet source field in $\lambda_1$ vs. $\lambda_2$ space (higher $\lambda_1$, higher $\lambda_2$). While there is a clear association between high Sr/Y, low Cu magmas and the amphibole fractionation field, the location of high Sr/Y magmas at low Dy/Dy* and moderate Dy/Yb can also be explained through a magma formed in the "melting of mantle garnet" field (bottom right quadrant) which subsequently experienced (1) garnet fractionation at pressures around 1.2 GPa (Alonso-Perez et al., 2009), followed by (2) amphibole +/- clinopyroxene fractionation at lower pressures (Alonso-Perez et al., 2009). Whatever the exact sequence of processes, a magma plotting in the bottom-right quadrant of Fig. 4 must have experienced some HREE depletion and/or LREE enrichment to match the predicted patterns of this metric (Davidson et al., 2013). Additionally, all magmas have a starting REE profile that may not begin at the "crosshair" intersection of the two bold black lines (defined for chondritic-derived melt (Davidson et al., 2013)). Rather, each magma is likely derived from a mantle source with a unique REE profile defined by prior melt extraction, metasomatism, and other pre- and syn-melting features. Correcting for such source features is beyond the scope of this work, but would be an interesting area of ArcMetals to explore more fully. The complexities of the petrogenesis leading to changes in Dy/Dy* should make us cautious about using the Dy/Dy* systematics alone to diagnose the petrogenesis of high Sr/Y magmas. Rather, any petrogenetic process proposed here must be validated by other independent metrics, empirical observations, and/or modeling.

This garnet to amphibole/cpx sequence may only be piecemeal at shallower pressures, where amphibole/cpx will dominate as a fractionating phase outside the stability field for garnet (Alonso-Perez et al., 2009). Thus, while the importance of amphibole in these systems will be demonstrated further, garnet likely also plays an important role, especially in the deep roots of arc magmatic systems (Alonso-Perez et al., 2009; Lee and Tang, 2020; Cox et al., 2020). By the same logic, the influence of clinopyroxene cannot be ruled out, as cpx will produce a similar compound fractionation trend as suggested by the blue line for amphibole-garnet in Fig. 4a, albeit with a steeper slope. Clinopyroxene is one of the most common minerals in arc volcanic rocks, and empirical determinations of pyroxene stability indicate that amphibole-friendly magmatic conditions are comparatively favourable for clinopyroxene (Ulmer et al., 2018; Alonso-Perez et al., 2009). However, amphibole is favoured over clinopyroxene at higher magmatic water contents (Alonso-Perez et al., 2009) and lower temperatures (Alonso-Perez et al., 2009; Ulmer et al., 2018) i.e. conditions more common in andesites stored in the shallow to mid-crust (aligning with the conditions of magmatic storage prior to porphyry formation (Chiaradia and Caricchi, 2017)). Furthermore, the $\lambda$ plots show less ambiguously that high Sr/Y, low Cu magmas often have a much
steeper slope, consistent with amphibole control as opposed to cpx (Fig. 5a,b). This emphasizes the importance of combining geochemical metrics to get the most out of whole rock data.

Garnet fractionation, as discussed in Section 1, has gained popularity in the recent literature as a potential ore fertility mechanism (Chen et al., 2019; Lee and Tang, 2020; Cox et al., 2020). Experiments have shown that garnet is stabilized as a fractionating phase at pressures above 0.8 GPa (approximately 24 km. depth) (Alonso-Perez et al., 2009; Greene et al., 2006) and where melt water contents are high (above 4 wt.%) (Alonso-Perez et al., 2009). Direct evidence of garnet in arc magmas is rarely found in modern volcanics, but it has been found commonly in fossil arc systems (Bissig et al., 2017). Whether ancient or modern, where garnet can be seen widely in arc systems is in the lower crust cumulate lithologies of exhumed “arc roots,” sections like the type section in Kohistan, Pakistan (Petter, 2019; Jan and Howie, 1981; Ducea et al., 2015). Models developed for the mantle wedge underlying Central America found evidence for the presence of mantle heterogeneities rich in garnet-peridotite or garnet pyroxenite lithologies (Feigenson and Carr, 1993). Melting of such mantle garnet “veins” would impart a signature with higher Dy/Yb than the arc array, as discussed in detail in previous work (Davidson et al., 2013; Feigenson and Carr, 1993).

While the role and importance of garnet fractionation is not disputed in explaining the occurrence of some magmas with low [Cu] in thicker-crust arcs (Cox et al., 2020; Lee and Tang, 2020), this analysis suggests that amphibole is also an important candidate for moderating global arc Cu systematics as shown in Figs. 4 and 5. The importance of amphibole has been previously proposed in the context of porphyry deposits (Loucks, 2014) and the broader controls on magmatic Fe (Richards, 2015), although in both of those cases REEs were not used as a metric to measure amphibole’s presence. Amphibole fractionates in many magmatic environments at moderate (15–40 km.) depths. For many of the reasons the garnet hypothesis is forgoing the methods of Shaw 2006 (Shaw et al., 2006): Starting abundances of Ni, Cu, Sr, Y, and all REEs were taken from the trace element compositions of the relevant starting materials (Baker et al., 1994; Hurlimann et al., 2016; Grove et al., 2002), and used to model the partitioning (reconstructing Cu, Ni, Sr, Y, and REE systematic) (4.3.1) and sulphide saturation (4.3.2) model to estimate the impact of amphibole on melt chemistry and subsequent sulphide systematics. Special attention is paid to amphibole’s method of changing melt chemistry, with regards to the calc-alkaline differentiation trend discussed in Fig. 3. All of these empirical and modeled observations are synthesized together by a simple mass balance model to link the fractionation of amphibole with the stabilization of sulphide and subsequent depletion of [Cu].

### 4.3. Amphibole control on melt chemistry and sulphide stability

To understand how amphibole is able to affect these global Cu trends, it is necessary to show how amphibole can provide a link between high Sr/Y, calc-alkaline, potentially porphyry-developing magmas, and their low Cu contents. This is first done by analyzing the empirical results of a well-constrained series of isobaric fractional crystallization experiments (Ulmer et al., 2018) in the context of the low Cu, high Sr/Y magmas centered in this work (Section 4.3.1). These empirical datasets are integrated with a trace element partitioning (reconstructing Cu, Ni, Sr, Y, and REE systematic) (4.3.1) and sulphide saturation (4.3.2) model to provide a link between high Sr/Y, calc-alkaline, potentially porphyry-developing magmas, and their low Cu contents. This is first done by analyzing the empirical results of a well-constrained series of isobaric fractional crystallization experiments (Ulmer et al., 2018) in the context of the low Cu, high Sr/Y magmas centered in this work (Section 4.3.1). These empirical datasets are integrated with a trace element partitioning (reconstructing Cu, Ni, Sr, Y, and REE systematic) (4.3.1) and sulphide saturation (4.3.2) model to provide a link between high Sr/Y, calc-alkaline, potentially porphyry-developing magmas, and their low Cu contents. This is first done by analyzing the empirical results of a well-constrained series of isobaric fractional crystallization experiments (Ulmer et al., 2018) in the context of the low Cu, high Sr/Y magmas centered in this work (Section 4.3.1).

#### 4.3.1. Analysis of empirical datasets

Trace element abundances of experimental results from an existing empirical study (Ulmer et al., 2018) were modeled to constrain the effect of amphibole on the liquid line of descent in arc magmas, in order to understand how amphibole fractionation may connect major and trace element systematics under high Sr/Y-favoring conditions. The experimental study in question ran isobaric experiments under equilibrium (EC) and fractional crystallization (FC) conditions at 1.0 GPa and from 1200 to 720 °C on hydrous oxidized basaltic (Hurlimann et al., 2016) (called “FC Mb AuPd” in the original work) and oxidized andesitic (Baker et al., 1994) (called “FC ba AuPd” in the original work) composition melts.

The experimental data used in this work’s model (Ulmer et al., 2018) were prepared by transcribing Table 1 (starting conditions), Table 2 (modal abundances of minerals at each experimental temperature step), and Table 3 (glass and amphibole compositions, determined by EPMA) into Supplemental Data Table 2 of this work. In Ulmer et al. 2018, major elements were the only measured chemical species. Starting abundances of Ni, Cu, Sr, Y, and all REEs were taken from the trace element compositions of the relevant starting materials (Baker et al., 1994; Hurlimann et al., 2016; Grove et al., 2002), and used to model the partitioning of trace elements into fractionating mineral phases following the methods of Shaw 2006 (Shaw et al., 2006):

\[
\frac{c_t}{c_0} = F^{D_{-1}}
\]

where \( t \) = timestep \( *t^* \) in a given experiment, corresponding to a specific set of \( T, P, \) and \( X \) conditions. Also called “run number” in Supplementary Table 2 our tab; \( c_t = con-
centration of an element in the residual liquid; $c_i = \text{initial concentration of an element in bulk liquid, before fractionation; } F^i = \text{fraction of residual liquid } i$; $D^i = \text{"Bulk } D^i\text{" or weighted sum of whole rock partition coefficients; }$

$$D^i = \sum_{i=0}^{X_i} X_i \cdot D_{i-1} \quad \text{(6)}$$

where $X_i = \text{mass fraction of mineral } i\text{ in accumulated solid fraction, and } D_{i-1}^i = \text{partition coefficient (or } K_d) \text{ between mineral } i\text{ and liquid } l$. Final form of Eq. (5) requires that $c_i$ is solved at each experimental step in the differentiation sequences:

$$c_i = c_o \cdot F^{D-1} \quad \text{(7)}$$

The effect of these trace element models on elements like Cu and Ni can be found in the Supplement (Fig. S11). Note that of the three FC experiments total in the experimental database, only two are focused on in this work (i.e. the oxidized runs). The results from Ulmer et al. (2018) experienced some empirical complications around 900 °C, and the remaining melt fraction (F) and resultant melt chemistry was reversed from what one would expect to occur naturally (decreasing F with decreasing T). In order to make modeling these results as straightforward as possible, runs 10–12 and 9–11 were discarded from the oxidized andesite and oxidized basalt experimental runs respectively. One of the most remarkable features of these empirical results is the well constrained liquid line of descent (LLD) for calc-alkaline, hydrous, moderately oxidized magmas, and the resulting association between LLD and major mineral phases like amphibole. Also remarkable is this empirical datasets constraint of the changes of $\text{FO}_2$ with magmatic differentiation (Ulmer et al., 2018).

4.3.2. Comments on amphibole’s effect on Mg#

Experimental work (Alonso-Perez et al., 2009; Ridolfi et al., 2010; Nandedkar et al., 2016; Ulmer et al., 2018) has already implicated amphibole as one of the most important phases controlling major element characteristics in high pressure calc-alkaline magmas. However, invoking amphibole as an important agent of melt chemistry change, as suggested from Figs. 4 and 5, raises the question of how amphibole, with an Fe/Mg exchange coefficient ratio of around 0.3 (Nandedkar et al., 2016), can maintain the consistent melt Mg# trend observed in calc-alkaline suites (see Fig. 3, high Sr/Y trend). The calc-alkaline trend observed in high Sr/Y magmas sees a constant decrease of FeO with MgO, meaning that equal proportions of both elements have to be leaving the melt to explain the changes in melt chemistry. Despite the higher exchange coefficient of Mg in amphibole as compared to Fe (Ridolfi et al., 2010), some empirical work has noted that the majority of major element differentiation in metaluminous/peraluminous magmas should be controlled by amphibole at the relevant pressure and temperature conditions (Alonso-Perez et al., 2009).

To test this, the relationship between amphibole, melt Mg#, and FeO contents from the oxidized FC experiments (Ulmer et al., 2018) has been plotted in Fig. 6. Where amphibole stabilizes at 1050 °C (Fig. 6a), the Mg# of the melt does not see the predicated rapid Mg# drop off as a function of either temperature (Fig. 6a) or melt FeO wt.% (Fig. 6b and c). Instead, melt Mg# decreases smoothly through the amphibole stability field. Two major factors help to explain this trend: (1) coexisting phases crystallizing at the same time as amphibole, and (2) amphibole’s increasing FeO content and correspondingly lower Mg# as the magmas cool. Starting with the former, phases including clinopyroxene, garnet, plagioclase, ilmenite, and magnetite (in order of appearance) contribute to changes in melt chemistry in the andesite, and in the basalt, this series of other minerals also includes spinel. By mineral chemistry, phases like ilmenite, magnetite, spinel, garnet (Fe/Mg exchange coefficient of around 0.7, Alonso-Perez et al., 2009), and even clinopyroxene contain enough iron to explain the consistent decrease in FeO seen in the empirical data modeled here. It should be noted however that amphibole is one of the dominant minerals in the fractionating sequence following its initial stabilization at 1050 °C - thus, a good portion of the changes in melt chemistry must be attributed to amphibole. Amphibole stability may in fact be a marker all on its own of the right conditions contributing to calc-alkaline differentiation; contrast the trends seen in Fig. 3 for the oxidized andesite and oxidized basalt (Fig. 3e) with that of the reduced basalt (FC ba Pt-C), where the lack of amphibole contributes to the more tholeiitic pattern of differentiation and a lack of FeO depletion.

Furthermore as the empirical (Ulmer et al., 2018) melts’ reach temperatures below 900 °C, Fig. 6b and c show that amphibole itself begins to take on more iron into its structure. The stable amphibole species steps down from paragasis at temperatures >900 °C, to tschermakite (900 °C), tschermak hornblende (800 °C), and finally magnesiohornblende (720 °C) at the end of each fractional crystallization experiment (amphibole names follow established nomenclature (Leake et al., 1997)). Not only does the FeO wt.% of amphibole increase with greater degrees of differentiation, but Fig. 6c sees a decrease in the Mg# of fractionating amphibole’s as well. Fig. 6 shows that, consistent with the literature (Alonso-Perez et al., 2009; Ridolfi et al., 2010; Nandedkar et al., 2016; Ulmer et al., 2018), amphibole is not only associated with fractionating assemblages moving an arc magma towards calc-alkaline differentiation, but it can indeed alter the FeO content of a melt, in spite of its general preference for Mg cations over Fe. This extended discussion of amphibole’s impact on Fe is pertinent to the crucial element of modeling done in this study, namely Fe’s impact on sulphide saturation.

4.3.3. Integrating sulphide models into empirical results

To see whether major and trace element changes in a silicate melt affect the stability of sulphides in the melt requires an empirical model of sulphide saturation. Sulphide saturation is described by the experimental parameter, “sulphur content at sulphide saturation,” or SCSS (O’Neill and Mavrogenes, 2002). Generally, SCSS is negatively correlated with pressure (O’Neill and Mavrogenes, 2002) and positively correlated with temperature (O’Neill and Mavrogenes, 2002; Nash et al., 2019;
For mid-ocean ridge basalts, melt FeO content, oxygen fugacity ($f_{O_2}$), temperature, and pressure are the main drivers of sulphide saturation (Kiseeva et al., 2017). In arc magmas, higher water and sulphur contents (Wallace and Edmonds, 2011; Edmonds and Mather, 2017) coupled with higher oxidation state (Richards, 2015; Jugo et al., 2010) cause the SCSS to respond differently as compared to MORB, though major elements like FeO still play a major role. For most arc magmas, volatile and oxidation conditions lie outside the range for which many SCSS models are calibrated (O’Neill and Mavrogenes, 2002), with some exceptions (Jugo et al., 2010; Fortin et al., 2015). Since most arc magmas have a considerable fraction of sulphur present as $S^{6+}$, they may instead saturate in anhydrite, which is much more soluble in silicate melts (Zajacz and Tsay, 2019).

SCSS was modeled for the oxidized basalt and oxidized andesite runs (Figs. 3, 5, 6, 7, 8), using the major elements (Ulmer et al., 2018) and modeled trace elements (Cu, Ni) of the individual products as inputs using a new SCSS parameterization (O’Neill, 2020):

$$\text{SCSS} = \Delta G^0_{\text{FeO+Rb}} / RT + \ln C_{S^{6+}} / \ln \phi_{S^{6+}}$$

This SCSS method builds on the work of (Smythe et al., 2017), and is very sensitive to the bulk silicate melt composition, the sulphide composition, Fe-Ni-Cu partitioning into that sulphide, and P and T at ranges appropriate for these models (O’Neill, 2020). For all experiments, SCSS decreases with FeO and decreasing temperature (Fig. 7a and b respectively) (O’Neill and Mavrogenes, 2002). The SCSS values are initially calculated assuming the redox state of the magma will favor mainly $S^{2-}$ (O’Neill, 2020). To account for the likely abundance of $S^{6+}$, SCSS was calculated for a range of potential $S^{6+}$ speciation end members based on the ANNO buffer values for $f_{O_2}$ measured in the modeled experiments. These values, 0.5, 1.0, and 1.5 ANNO, correspond to AQFM values of 0.78, 1.28, and 1.78 respectively (ANNO to AQFM conversion accomplished using the Excel tool “Calc-fO2-buffer” from http://www.kaylaiacovino.com/tools-for-petrologists/, using data from previous work (Frost et al., 2001)). These $f_{O_2}$ values translate to $\phi_{S^{6+}}$ values of 0.2, 0.75, and 0.95 following a simple correction scheme developed for oxidized magmas (Wieser et al., 2020):

$$\text{SCSS}_{\text{tot}} = \frac{\text{SCSS}^{\text{FeS}}}{1 - \phi_{S^{6+}}}$$

This in turn is based on an earlier parameterization of $S^{6+}$ as a function of AQFM buffer (Jugo et al., 2010):

$$\frac{S^{6+}}{S^{2-}} = \frac{1}{1 + 10^{3.33 - 2\Delta RM^O}}$$

Fig. 7 shows the results of modeling the different $S^{6+}$ abundances in the magmas by using different symbols denoting the different oxidation state of $S$ in each empirical dataset. The onset of amphibole crystallization is shown both as function of FeO (Fig. 7a) and temperature (Fig. 7b). To aid the reader in understanding what major minerals are fractionating during the experiments, a schematic blue line has been added to Fig. 7a showing the major controls on SCSS changes at different stages in the model. Where amphibole was the dominant fractionating phase in these experiments, there was only secondary involvement of minerals like garnet, magnetite, ilmenite, clinoxyroxene, plagioclase, and spinel, though amphibole become less abundant as temperature continued to decrease (Ulmer et al., 2018). High-temperature (>1050 °C) FeO loss is attributed to clinopyroxene and to a lesser extent orthopyroxene and olivine (only in the basalt), followed by a much more amphibole (as well as garnet) mediated lowering of SCSS at temperatures below 1050 °C. These two FeO loss trends, clinopyroxene vs. amphibole mediated, are separated by a small but abrupt decrease in SCSS, which is a function of temperature change as clinopyroxene abundance decreases and amphibole starts to appear (the “drop-off” on the blue curve in Fig. 7a).

The onset of amphibole fractionation is associated with a continued lowering of the SCSS, but no clear dramatic decrease (Fig. 7a, b). However, if we consider conditions relevant to porphyry development, clinopyroxene in these experiments only fractionates at temperatures greater than 1050 °C, which should be less common in the sort of mid-crust, high water, low Cu and high Sr/Y magmas presumed to be important for developing porphyry systems. The continued drop in SCSS as a function of FeO (Fig. 7a) can be attributed to other phases stabilizing alongside amphibole, and the increasing FeO and decreasing Mg# trends seen in amphiboles from Fig. 6b and c. Since arc magmas usually contain bulk sulphur contents between 1000 and 2000 ppm (red box in Fig. 7a and b) (Wallace and Edmonds, 2011), these models confirm that most hydrous arc magmas are at or near sulphide saturation (Richards, 2015; Du and Audetat, 2020) during most of their differentiation, and as a result will become depleted in Cu as sulphides are removed: this hypothesis can be verified by further modeling (see Section 4.3.4). Furthermore, these magmas demonstrate that even with $\phi_{S^{6+}}$ values approaching 0.75 at AQFM values greater than 1.2 (diamond and circle curves for andesite and basalt, respectively), sulphide fractionation can be pervasive.

The noticeable but not necessarily dramatic decrease in SCSS at amphibole-in is mediated by several competing factors: (1) As should be obvious from the SCSS parameterization above, FeO is the crucial oxide driving much of the change in melt SCSS (O’Neill and Mavrogenes, 2002; O’Neill, 2020). Of particular importance is the activity of FeO in the melt, $a_{FeO}$, which is affected by the activity coefficient of FeO $\gamma_{FeO}$ according to Eq. (8) (Eqn. 46 in original source (O’Neill, 2020)). (2) Different cations have a competing effect in this $\gamma$ term. For example, MgO has a strong negative correlation with $\gamma_{FeO}$. This means that higher MgO will lower the activity coefficient of FeO, mitigating
the drop in SCSS as would be expected from FeO loss alone. Should olivine or another mafic phase precede amphibole fractionation as is the case for the oxidized basalt, the comparatively lower MgO will lead to a strong increase in $\gamma_{FeO}$, magnifying the SCSS decrease. (3) By the same logic, the strong positive correlation between Na, K, and $\gamma_{FeO}$ implies that a loss of alkali elements will enhance SCSS reduction during amphibole fractionation. So the SCSS-buffering effect is mitigated in part by a strong decrease in the alkali content of the oxidized andesite melt, promoting the continued SCSS decrease seen in Fig. 7. (4) The In $\gamma_{FeO}$ term, or the activity of the sulphide itself, has a large impact on SCSS. Related work has shown that the interplay between the Fe, Cu, and Ni abundances in sulphide is a strong control on the SCSS (Wallace and Edmonds, 2011). Hence, the models of Cu and Ni abundance in the trace element partitioning models included a small weight fraction (0.001) of sulphide fractionating at each step, assuming the sulphide in question is a sulphide melt (see next section for details), and using partition coefficients for Cu and Ni between sulphide melt and silicate melt from (Li et al., 2012). These Kd's are 1070 for Cu, and 490 for Ni, taken from experimental run LYO4 (Li et al., 2012). The consistent decrease of Cu and Ni in the silicate melt leads to a concurrent decrease in the sulphide Fe/(Fe + Cu + Ni) content in the hypothetical sulphide, and a corresponding decrease in the SCSS. Finally, (5) the continual decrease in temperature at constant pressure as performed in these experiments strongly affects the SCSS in the preferred SCSS model (O’Neill, 2020). This approach to modeling SCSS alongside calc-alkaline differentiated magmas shows the potential for amphibole to contribute to the modification of wholesale melt chemistry.

4.3.4. Mass balance of amphibole’s effect on S

Fig. 7 shows the almost wholly linear decrease of modeled SCSS as the oxidized andesite and basalt runs progress. However, SCSS alone only tells half the story - to explain how amphibole fits into the story of the low Cu, high Sr/Y magmas concerned in this work, there needs to be a compelling link between this decrease in SCSS and the melt’s sulphur content [S]. One way to demonstrate this is to take a similarly simple FC modeling approach as was done for the trace elements during the crystallization of the oxidized andesite and basalt. This is made possible if (1) there is a [S] imposed on each empirical melt, (2) the SCSS can be related to the [S] in such a way as to estimate the mass of sulphide fractionating at each step, and (3) the melt [S] can subsequently have been shown to have been perturbed by the modeled decrease in SCSS as a function of amphibole.

The results of these models are shown in Fig. 8. Fig. 8a shows the wt% of amphibole crystallizing in each experimental step (Ulmer et al., 2018). Amphibole stabilizes at 35% crystallization (or $F = 0.65$) in the oxidized andesite (FC ba AuPd), and 63.70 % crystallization (F = 0.363) for the oxidized basalt (FC Mb AuPd) (Ulmer et al., 2018). S contents in the melt were estimated based on inventories of [S] in the literature (Wallace and Edmonds, 2011), as the starting experimental materials did not have reported measurements of volatile trace elements like sulphur (Baker et al., 1994; Hurlimann et al., 2016; Ulmer et al., 2018). The modeling done here assumed an oxidized arc basalt to have a starting [S] of 2000 ppm, or 0.2 wt.% (Wallace and Edmonds, 2011). Andesites should have already experienced some [S] loss, either as a consequence of degassing or as Fig. 3c shows as a result of basaltic differentiation and subsequent sulphide fractionation - thus, the oxidized andesite series was modeled as having a starting melt [S] of 1000 ppm (corresponding roughly to the point the empirical basalt reached andesite-like melt compositions). Next, a simple mass balance was adapted from earlier work (Kiseeva and Wood, 2015) on sulphide’s effect on [Cu] in MORB composition magmas:

$$X_\text{ulf} = \frac{S_\text{sulf} - (SCSS)(X_\text{melt})}{S_\text{ulf}}$$

where $X_\text{ulf} =$ the mass fraction of sulphide produced at each experimental step; $S_\text{sulf} =$ the starting [S] in each step (initially 2000 and 1000 ppm respectively for the basalt and andesite); SCSS =$ the modeled SCSS value (O’Neill, 2020); $X_\text{melt} =$ the remaining fraction of melt, or F and; $S_\text{ulf} =$ the wt.% concentration of sulphur in the fractionating sulphide phase.

The goal of using this equation is to estimate the abundance of sulphide leaving the system once SCSS and [S] become equivalent, and by extension to model the decrease in [S] as a product of calc-alkaline differentiation. While the prior modeling in this work coupled with the original empirical results provide most of these parameters, this equation requires a fit to a particular sulphide composition, $S_\text{ulf}$. A suitable magmatic sulphide is required, one which was in equilibrium with basaltic to andesitic composition magmas, and which can also house significant amounts of Cu and Ni. The sulphide chosen was an average sulphide melt modeled to be the parent of lower-temperature sulphides at Merapi volcano, Java, Indonesia (Nadeau et al., 2010). Sulphide melts were prefered here over other sulphide phases like pyrrhotite or cubanite because they are often primary to the original melt, and only decompose into other, crystalline sulphide phases at lower temperatures (Nadeau et al., 2010). The average sulphide melt from Merapi was estimated as having 38.2 wt.% S, 52000 ppm Cu, and 2300 ppm Ni (Nadeau et al., 2010). Fitting this value as $S_\text{ulf}$ into Eqn. 11 allows us to construct a mass balance of sulphur in the empirical silicate melts analyzed in this study. The model results are presented as an Excel worksheet as Supplementary Table 5. As previously discussed, the silicate melts used here (Ulmer et al., 2018) correspond well to magmas stored at moderate depth under high H$_2$O, moderate to high pressure conditions. Thus, these results should be generalizable to many other hydroxide arc andesites and basals analyzed from ArcMetals and other compilations.

Fig. 8b and c shows that amphibole crystallization corresponds to almost all of the sulphide fractionation and subsequent [S] loss in andesites. Starting from a basalt, amphibole stabilizes too late, such that amphibole plays only a marginal role in adjusting melt chemistry and corresponding sulphur and sulphide changes. Thus, conditions that favor olivine and pyroxene stability are conducive to
widespread sulphide fractionation in the most primitive arc basalts, supporting our contention that most arc magmas experience pervasive sulphide saturation. However, this work has been much more concerned with the kind of andesitic, high Sr/Y magmas that would predominate in the mid- to lower crust in porphyry-friendly settings. In andesites, amphibole is the only major phase crystallizing when sulphides stabilize (Fig. 8c). These coinciding trends are the combination of a lower S, (due to earlier basaltic sulphide fractionation) combined with the earlier stabilization of amphibole and subsequent amphibole-mediation of melt chemistry changes (e.g. Fig. 6). Thus it seems where an andesite begins to fractionate, amphibole is the dominant mediator of sulphide fractionation. This fits with general observations that those high Sr/Y, low Cu magmas (with measurable Dy/Dy* and λ deviations consistent with amphibole) are likely intermediate composition trapped at depth. These results also suggest that it is amphibole-mediated changes in andesite melt chemistry, in turn a function of melt temperature, that lowers SCSS enough to cause sulphide precipitation as the [S] in the melt increases slowly as a an incompatible element until reaching the SCSS (Fig. 8c). This again highlights the importance of amphibole as both a mediator of melt chemistry changes during calc-alkaline differentiation, and a signal of the optimal conditions for calc-alkaline and low Cu conditions in a melt. However, as Fig. 7 showed, amphibole isn’t radically changing melt chemistry all on its own. Rather, the conditions associated with amphibole crystallization (P, T, H₂O, fO₂, melt chemistry) are conducive to a noticeable if modest decreases in SCSS, FeO (and other major elements), [S] and thus mass of sulphide fractionated and concurrent decrease in melt [Cu], where some but not all of these changes are directly the effect of amphibole. Equally important but not considered in depth in this work is the role of S degassing; should a magma degas significant volumes of sulphur, then is direct evidence linking MSS fractionation with Cu depletion in arc magmatic sequences. Such a connection would not only emphasize the pervasiveness of sulphide fractionation in arc magmas, but would also further implicate minerals like amphibole, clinopyroxene, and garnet as controls sulphide stability. This study attempts to provide such evidence by analyzing those whole rock compositions in ArcMetals that can be shown to have fractionated MSS sulphide. Fig. 9 plots whole rock Cu/Ag vs. MgO, coloured for both Gd/Yb (Fig. 9a), Dy/Dy* (Fig. 9b), and for crustal thickness (CT) (Fig. 9c). The motivation behind constructing such plots stems from the fact that Cu fractionates more strongly into MSS than Ag (Patten et al., 2013; Li et al., 2012) (whereas the opposite is true in sulphide melts: sulphide melt fractionation should promote higher Cu/Ag), and such a ratio gives us the benefit of sensitively detecting the presence of a fractionating MSS at sulphide saturation (Jenner, 2017). A low Cu/Ag ratio, below average mid-ocean ridge basalt (MORB) (Jenner et al., 2012; Jenner, 2017) and continental crust (HHolland and Turekian, 2004), is consistent with crystalline sulphide fractionation and consequent Cu removal from the silicate melt (Jenner, 2017) at lower temperatures than were modeled in Fig. 7.

Gd/Yb (Fig. 9a) is a proxy for garnet involvement in petrogenesis because Gd partitions less strongly into garnet than Yb (Jenner et al., 2012), and has been used to infer the prevalence of garnet fractionation in the Andes (Cox et al., 2020). The highest Gd/Yb ratios are associated with whole
rocks with the lowest Cu/Ag for a given MgO content (Fig. 6a), suggesting a direct correlation between the proportion of garnet fractionation and the proportion of crystalline sulfide fractionation. Owing to the complexities in analysing whole rocks for Ag, Cu/Ag datasets are rare and are currently biased towards the Andes data, though there are some measurements in other transitional arcs (Fig. 9 symbols). There is no clear link between Cu/Ag in the whole rock and crustal thickness plot, but the very thickest crust (>40 km) is associated with evolved volcanic rocks, with a higher Gd/Yb (indicating garnet involvement) and low Cu/Ag. Fig. 6b shows that lower Dy/Dy*, indicative of amphibole, is also associated with low Cu/Ag, evolved magmas. Thus, both garnet and amphibole are implicated in the petrogenesis of magmas that have experienced the most crystalline sulphide fractionation. One drawback to using Cu/Ag as a proxy for sulphides in the global database is the dearth of available Ag and other chalcophile data in the literature. Only in the past 5–10 years have Ag and other difficult to measure elements become easily measurable using ICPMS (Jenner, 2017; Jenner and Arevalo, 2016; Cox et al., 2019; Cox et al., 2020; Wieser et al., 2020). As studies reporting suites of chalcophile elements in magmatic systems grow, future iterations of our ArcMetals database may yet be able to make broader, more detailed analysis of Cu/Ag in arc systems. Furthermore, our earlier modeling (Figs. 7 and 8) suggest that a proxy like Cu/Ag is not necessary to confirm sulphide fractionation, as sulphide fractionation should be a ubiquitous phenomenon in arc magmas, thanks to changes in SCSS mediated by phases like amphibole. However, natural data, without the benefit of the kind of thermodynamic knowledge available in empirical datasets, can still benefit form chalcophile trace element ratios like Cu/Ag.

It is worth noting that the Gd/Yb content of a magma could be affected by both garnet fractionation, or melt derivation from a garnet rich source rock. If the latter, one would expect small fractional melts to have high Gd/Yb followed by progressively lower Gd/Yb as more and more Gd poor minerals melted out. To pick apart these competing REE trends (shown schematically in Dy/Dy* space in Fig. 4, and in λ space in Fig. 5) is beyond the scope of this article.
of this work. Prior work on Cu/Ag establishes that particular volcanoes in the Andes likely experienced garnet fractionation as opposed to a mantle garnet signature (Cox et al., 2020). One way to assess how widespread mantle-garnet melting is as reflected in whole-rock geochemical records would be to compare a suite of garnet-fractionating rocks to a suite of candidate garnet-melting rocks, like those analyzed for REEs in Central America (Feigenson and Carr, 1993). A few well constrained cogenetic suites of rocks, analyzed for REEs and plotted as a liquid line of descent in Dy/Dy* and λ space would show what the relevant differences, if any, there are between garnet fractionation and garnet melting in complex natural magmas.

Our schematic model (Fig. 10) emphasizes the primary importance of amphibole fractionation and the related importance of garnet/chloropyroxene fractionation (Lee and Tang, 2020). This work emphasizes the importance of amphibole as a potentially widespread petrological-control mechanism that explains many of the generic features of high Sr/Y magmas in arcs. The global applicability of this amphibole-centred model complements garnet fractionation models, which work best in arcs with mature magmatic columns, deep brittle-ductile transitions (which promote longer residence times of magma in crust (Lee and Tang, 2020)), and thicker crusts (Lee and Tang, 2020; Cox et al., 2020). This amphibole model also complements the more immature arc focused magnetite fractionation models, which work best in young island arcs and/or back-arc basins (Jenner et al., 2010; Jenner et al., 2012). Our amphibole-centered model should be viewed on a continuum with these other petrological models, where water-rich, calc-alkaline trending arc magmas stored in the mid to deep crust (15–40 km.) will fractionate amphibole and stabilize sulphide in “semi-mature” arcs (called “immature continental arcs” as proposed in Lee et al. 2020, Fig. 6b (Lee and Tang, 2020)). The exact depth of storage, temperature, /O2, and amphibole stability in these magmas will depend on many geological, chemical, and physical factors, as elaborated on in relevant experimental results (Ulmer et al., 2018; Alonso-Perez et al., 2009; Nandedkar et al., 2016; Ridolfi et al., 2010). While the specific depth at which amphibole-mediated SCSS reduction will occur is variable, the important feature to note about this model is that amphibole represents a bridge between the magnetite-dominated and garnet-dominated petrological models proposed in recent years. As these semi-mature arcs evolve, garnet will fill much the same role as amphibole in controlling Fe and stabilizing sulphide. This allows for a certain degree of temporal evolution in the primary petrological vector controlling bulk [Cu] in arc magmas, where one would expect a young, thermally immature arc to modulate its Cu via magnetite (Jenner et al., 2010), evolving to amphibole at a moderate maturity and thickness, culminating in garnet-dominated Cu control by the time the arc reaches maturity (Lee and Tang, 2020).

One point of departure with the analysis in this study as compared to previous studies relates the importance and timing of sulphide saturation. Some studies (Lee and Tang, 2020; Park et al., 2019) have assumed that early sulphide saturation is detrimental to eventual porphyry copper deposit development, articulating the ‘Cu paradox,’ of low magmatic [Cu] being associated with ore deposits (Lee and Tang, 2020). Studies using platinum group elements as proxies for sulphide saturation in arc systems make this assumption more explicit, arguing that ore development requires late sulphide saturation (Park et al., 2019). However, as suggested in recent work (Du and Audetat, 2020), porphyry systems that have experienced both prolonged amphibole fractionation and early sulphide crystallization (and subsequent Cu loss) seem perfectly capable of developing porphyry deposits later in their lifetime (Du and Audetat, 2020). The global analysis shown here appears to confirm that melt Cu concentration does not act as a primary control on the potential of a magma to go on and make a porphyry deposit. Similarly, this amphibole-mediated SCSS reduction model does not imply that amphibole fractionation is a critical missing link leading to porphyry development. Rather, widespread amphibole
fractionation (as reflected by the petrography and/or whole-rock chemistry of a magma) is a consequence of particular magmatic conditions (P, T, /O2) that might, under the right geological conditions, signal the right environment to develop a porphyry. Thus, amphibole is seen as a driver of melt chemistry and SCSS changes, but more and an indicator of, rather than a driver of, porphyry development. In agreement with the observation that high Sr/Y magmas have lower Cu concentrations (Richards, 2015; Lee and Tang, 2020) (Fig. 3c), this analysis finds that combined amphibole and/or garnet fractionation of whole rock chemistry (Fig. 4, 5) can lead to early sulphide saturation (Figs. 7 and 8). The low magmatic Cu concentrations observed in the high Sr/Y magmas in this database are fundamentally the consequence of the deep, hydrous evolution of arc magmas magmas. This crustal processing can lead to higher the consequence of the deep, hydrous evolution of arc mag-

...ation, and subsequently increases the likelihood of sulphide...
AUTHOR CONTRIBUTIONS

NDB conceived the project with advice and comments from ME and FJ. NDB compiled the database, wrote the Python code, and performed the analysis. ME, FJ, AA, and HW provided equal contributions regarding data interpretations and significant references. NDB wrote the manuscript with equal editing contributions from ME, FJ, AA, and HW.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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APPENDIX A. SUPPLEMENTARY MATERIAL

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.gca.2021.05.034.

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