

Application of LA ICP-MS sulphide analysis and methodology for deciphering elemental paragenesis and associations in addition to multi-stage processes in metamorphic gold settings

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- 1 Title: Application of LA ICP-MS sulphide analysis and methodology for deciphering
- elemental paragenesis and associations in addition to multi-stage processes in metamorphic
 gold settings
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- 9

10 Abstract

Quantitative laser ablation inductively coupled plasma-mass spectrometry (LA ICP-MS) 11 12 element distribution analysis represents a relatively new and powerful approach to assess gold mineralizing processes. Although there is a trend towards imaging in LA ICP-MS, this article 13 demonstrates that the compilation of the individual time-slice datasets (TSD) from LA analyses 14 provides greater potential to explore and better understand ore systems. Based on several 15 analyses of gold-bearing sulphides from a variety of lode gold mineralized settings, this study 16 introduces new diagrams and geostatistical techniques as recommended procedures to be applied 17 to any gold deposit environment to better understand its genesis with possible exploration 18 implications. The nature of the mineralizing fluid is investigated by studying Ni and Co 19 20 variations in the sulphide grains in environments ranging from ultramafic to felsic, whereas upgrading of gold tenor within an ore deposit is identified using Au versus Ag plots grouped by 21 element association and elemental geochemical affinity. Furthermore, the textural transformation 22 23 and coupled dissolution/precipitation (CDP) in pyrite is documented using Ag versus Ni plots with associated defined fields (i.e., diagenetic versus metamorphic). Lastly, standard 24

geostatistical tests are used to assess element associations through agglomerative hierarchical
 clustering and exploratory statistical methods. Meanwhile, a multidimensional scaling approach
 can evaluate the geochemical affinity of the mineralizing fluids.

Keywords: laser-ablation ICP-MS, sulphides, gold, trace elements, procedure, and statisticalanalysis

30 1. Introduction

Quantitative laser ablation inductively coupled plasma-mass spectrometry (LA ICP-MS) 31 element distribution maps and profiles of sulphides are relatively new and powerful geochemical 32 techniques that provide important insight into the identification and interpretation of multi-33 34 dimensional element coupling and decoupling, elemental paragenesis and mineralizing events. These methods have been applied to a wide variety of gold deposit types (e.g., Large et al., 2007, 35 2009, 2011; Ulrich et al., 2009; Cook et al., 2013; Bull and Large, 2014; Gao et al., 2015; 36 37 Lawley et al., 2015; Kontak 2015). The application to multiple gold-associated sulphide phases 38 (e.g., diagenetic and later pyrite generations, metamorphic pyrrhotite and hydrothermal 39 arsenopyrite) has provided concentration data and the spatial context of various elements in the 40 host sulphide phase (e.g., primary grains, recrystallized grains, growth patterns, fracture 41 networks, inclusions) at a multi-micron scale (i.e., 10s µm). Importantly, this method also 42 minimizes the averaging effect of conventional bulk solution-based ICP-MS analyses, which commonly masks important information that relates to elemental associations and their 43 44 paragenesis (e.g., Ulrich et al., 2009; Warren et al., 2015; Jenner and Arevalo, 2016). 45 Most previously published laser ablation-ICP-MS element distribution maps were created by 46 stitching together many time-resolved traverses or spots. This approach has visual appeal, but

47 may not reveal subtle element associations. Alternatively, or in tandem, time-resolved traverses can be used as "time-slice datasets" (TSD) where each sweep of the MS is treated as a separate 48 data point providing the means to evaluate relationships between elements at a much finer scale. 49 The strength of this latter approach are rarely exploited, but a few recent examples of this type of 50 51 data processing generated at our institution have been presented (e.g., Dostal et al., 2015; Genna 52 and Gaboury, 2015; McDivitt, 2016; Neyedley et al., 2016; Gourcerol et al., 2016a). The advantage of this approach is that it provides a means to identify elemental correlations not 53 readily observed from element distribution mapping, as is illustrated below, and thus enables one 54 55 to establish an elemental paragenesis for the phase analyzed and thereby the mineralized system. Therefore, the latter supplements information provided by the element distribution maps. In the 56 case of gold deposit studies, integration of those maps and TSD treatment allows one to identify 57 the true element association of deposits which is not easily discernable solely using the 58 conventional technique of element distribution maps or traverses. 59

The purpose of this article is to present and discuss a routine means of integrating the 60 elemental distribution maps produced from LA ICP-MS analysis of sulphides with TSD in order 61 to optimize data interpretation in the context of : 1) elemental paragenesis, 2) deposit genesis, 62 and by extension 3) exploration. With these goals, a selection of samples from multiple deposits 63 are used herein to demonstrate the utility of these novel methods. In addition, several elemental 64 plots for sulphide phases are introduced (e.g., Ag versus Ni; Co versus Co/Ni; Au versus Ag) to 65 understand and portray the complexities of gold mineralizing events and the potential influence 66 of host stratigraphy on fluid and hence sulphide chemistry. We also use geostatistical methods 67 (e.g., multidimensional scaling and discriminant analysis) to: 1) confirm the element associations 68 suggested from the maps and binary element plots; 2) confirm the number of gold mineralizing 69

| 70 | events identified with TSD; and 3) address the nature of sulphide-transporting fluids. In order to |
|----|--|
| 71 | demonstrate the effectiveness of the TSD method, we use sulphide datasets gathered from |
| 72 | several Canadian orogenic gold deposits with which we are familiar based on our previous field |
| 73 | and petrological studies. These include the Archean banded iron formation (BIF) - hosted |
| 74 | Meliadine (Nunavut; e.g., Lawley et al., 2015; Gourcerol et al., 2016a, b), Meadowbank |
| 75 | (Nunavut; e.g., Janvier et al., 2015; Gourcerol et al., 2016a, b) and Musselwhite (Ontario; e.g., |
| 76 | Oswald et al., 2015; Gao et al., 2015; Gourcerol et al., 2016a, b) deposits consisting of sulphide |
| 77 | (i.e., arsenopyrite, pyrite and pyrrhotite, respectively) replacement of Fe-rich material (e.g., |
| 78 | magnetite), as well as the Phanerozoic metaturbidite-hosted Beaver Dam deposit (Nova Scotia) |
| 79 | in which sulphides such as arsenopyrite occur mainly as quartz vein filling (e.g., Kontak et al., |
| 80 | 1993; Sangster and Smith, 2007; Kontak et al., 2013). |

81 2. **Analytical methods**

82

2.1 LA ICP-MS sulphide trace element chemistry

The trace element chemistry of several sulphides were determined by LA ICP-MS in 83 Laurentian University's Chemical Fingerprinting laboratory. Measurements were made by 84 ablating sulphides with a Resonetics (now Australian Scientific Instruments) RESOlution M-50 85 193 nm, 20 ns pulse duration ArF excimer laser ablation microprobe coupled to a Thermo X-86 Series II quadrupole ICP-MS. The data reported herein were acquired in several different 87 88 sessions (of several hours) spanning approximately two years. For all sessions, ablation took place in ultra-pure helium flowing at a rate of 650 ml/min in a Laurin Technic two-volume 89 90 sample cell that has exceptionally fast wash-out (i.e., < 1.5 s for 99% signal washout), thus 91 affording good spatial resolution (Müller et al. 2009). The vaporized material and helium were combined with argon (750-800 ml/min) and a small amount of nitrogen (6 ml/min), for enhanced 92

93 sensitivity (e.g., Hu et al., 2012), outside of the sample cell and approximately 2.5 meters before
94 entering the torch. Ablation and plasma conditions varied from session to session and sample to
95 sample, and are summarized in Table 1.

96 Quantification of the raw data was conducted with the trace element data reduction scheme in Iolite 3 (Paton et al. 2011) using NIST610 (Jochum et al., 2011) and Fe (i.e., ⁵⁶Fe or 97 57 Fe since it varied from experiment to experiment) as the external and internal references, 98 respectively. Moreover, periodic measurements of BHVO2g and Po725 were used as reference 99 materials. Note that Fe is a major element in the sulphide grains and as such, its concentration is 100 101 significantly stoichiometrically controlled. Thus, all trace element concentrations which hinge on EDS/EMP analyses of Fe in individual sulphide grains are representative of the analyzed grains. 102 103 In order to avoid significantly overestimating concentrations in fractures and inclusions, the internal standard was not forced on a point by point basis, but was instead applied to a 104 105 representative section of the target mineral. In this way, the data are considered quantitative in 106 the target mineral (e.g. arsenopyrite, pyrite, pyrrhotite) and semi-quantitative in other phases (e.g., inclusions and/or surrounding material) and along fractures, etc. Despite differences in 107 ablation between NIST610 (silicate glass) and sulphides contributing to melting and/or 108 109 fractionation, similar measurements conducted by Wohlgemuth-Ueberwasser et al. (2015) yielded concentrations within 20% of accepted values when calibrating sulphides with NIST610. 110 We note that a glass reference material is not ideal, however, glasses are often used because there 111 are few if any widely available non-glass reference materials that are satisfactorily homogeneous 112 113 and cover a wide range of trace elements. The USGS synthetic doped basalt glasses (e.g., GSD-114 1G) are recommended for similar studies in the future due to their higher Fe content that more closely resembles the sulphides. Fig. 2 demonstrates this to some degree, even for individual 115

116 time-slices, as the time-resolved Au concentration from Po725 (Sylvester et al., 2005) calibrated 117 in the way described above are generally within 20% of the accepted value. In the present study, we are interested not in absolute concentrations of a given element, but rather the relationships 118 119 between elements.

120 Before conducting the LA ICP-MS analyses, photomicrographs, and domain descriptions 121 (e.g., fracture locations, zoning) were collected as they are required to properly interpret the geochemical data. It is important to make such observations prior to ablation, as LA ICP-MS is a 122 123 micro-destructive technique that may obliterate or obscure important morphological features.

124

2.2 Generation of elemental maps and time-slice datasets

The present study uses sulphide element maps and TSD to better understand Au mineralizing 125 126 processes. For maps, a series of parallel and equal length time-resolved traverses were merged 127 into images using Iolite (e.g., Woodhead et al., 2007) and further subjected to bilinear interpolation between adjacent traverses and 3x3 pixel smoothing to improve the data 128 129 visualization (see, e.g., Rittner and Müller, 2012). The color scaling of geochemical maps can 130 highlight different features and can have a significant influence on their interpretation. For instance, Fig. 1 shows the Co, Sb, and Ag element maps for an arsenopyrite grain from the 131 Dufferin deposit in the Meguma Terrane (Nova Scotia, Canada) produced using: 1) logarithmic 132 133 scaling, 2) empirical cumulative density function scaling, 3) linear scaling with ± -3 standard 134 deviation extrema, and 4) linear scaling with minimum/maximum extrema (note that this causes color saturation at the low and high ends of the spectrum). In addition, note that linear scales 135 tend to obscure low abundances. This figure demonstrates that color scaling alone may highlight 136 137 various aspects of the element distribution in the sulphide lattice and structure and, therefore, all options must be considered. 138

139 In terms of the TSD, each traverse (whether individual or as part of a map) consists of a series of individual time-slice measurements made during one cycle of the quadrupole. In this 140 study, rather than integrating data into a single number, we leave it in its time resolved form 141 142 consisting of measurements every 0.3 to 1.2 seconds (depending primarily on the number of analytes and dwell time per analyte). In doing so, a higher degree of spatial resolution (compared 143 144 to conventional spot-based geochemical mapping) is maintained, thus permitting the evaluation of domains such as zoning, fractures and inclusions (Fig. 2). Despite being time-consuming 145 (hours of work), this latter method thus provides the means to treat all the traverse data from a 146 147 single grain as analogous spot data, but is typically much less time intensive than constructing a map from spots (e.g., Gao et al., 2014; Lawley et al. 2015). 148 Moreover, rather than comparing TSD data to spot analyses, we compare the TSD data to the 149 accepted values for reference materials. Note that a spot analysis does not have the spatial 150 resolution for some of the features we describe if a large spot size is used. If instead, a small spot 151 size is used, there are down-hole effects (i.e., fractionation of elements based on their volatility). 152 153 There is fundamentally more noise within individual time-slices but the individual slices are typically within 20% of the accepted values and in a study of this type, the interest is in 154 155 variations on the scale of orders of magnitude given the extensive use of log based plots.

3.

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Treatment of data and applications

This section documents a step-wise procedure which should be considered as routine for the treatment and interpretation of LA ICP-MS multi-element analyses. The section is sub-divided to deal separately with elemental paragenesis, the nature of the mineralizing fluid(s), processes that may relate to upgrading of the gold tenor of the samples being studied and hence the deposit they come from, and the use of pyrite geochemistry to discriminate diagenetic versus metamorphic/hydrothermal origins. Later in this section a variety of statistical techniques foridentifying element associations are utilized.

164 **3.1 Elemental paragenesis**

Quantitative element distribution maps and traverses performed on sulphide grains, combined with detailed petrographic and/or scanning electron microscopy – energy dispersive X-ray spectrometry (SEM–EDS) studies, provide the basis to establish an elemental paragenesis which is specific to the studied sulphide grains. Therefore, this crucial step underlies the identification of element associations as well as their spatial distribution in the sulphide grain (e.g., core versus rim, fractures, and overgrowths).

Regarding element distribution maps, the intensity of colors highlights the concentration and 171 172 loci of elements, thus, showing their distribution throughout the grain. Fig. 3 illustrates an example of selected LA ICP-MS element distribution maps produced for an arsenopyrite grain 173 from the Beaver Dam gold deposit and the corresponding elemental paragenesis derived from 174 175 this is shown in Fig. 4. In this sample, Co mimics the growth zoning which suggests this element 176 has a primary origin. Other elements which are co-spatial with Co are similarly interpreted. Thus, in this case, Ni, Mo Se, Sb and to a lesser extent Au show similar patterns and are 177 therefore also inferred to locally mimic the morphology of the growth zoning. In some cases, 178 179 elements are seen to overprint the zoning and thus possibly reflect a relatively late primary 180 feature (i.e., Mo, Sb, Se). The uniformly distributed, albeit low concentrations of Ag and Zn suggest they are also in the sulphide lattice either as background concentrations or as local 181 micro-inclusions suggesting a primary origin. Bismuth, Pb and Ag are mainly associated with 182 183 early fractures (F1) whereas Al, Zn and Mn appear to be relatively late and are associated with later crosscutting fractures (F2), as deduced from petrographic study. Finally, Au seems to also 184

be broadly associated with both the F1 and F2 fractures as micro-inclusions (Fig. 3), and Ni and
Se seem to be reworked given their locally variable concentrations along fractures in addition to
Mo and Sb which may reflect remobilization along F1. Thus, based on the interpretation of the
elemental distribution maps, four element associations are discerned for this sample; 1) Co-Ni; 2)
Mo-Se-Sb (Au-Ag-Zn); 3) Bi-Pb-Ag (Au); and 4) Al-Zn-Mn (Au). We are unaware of any other
method whereby this information has previously been generated for sulphides from a mineralized
setting.

In contrast to elemental maps, elemental traverses only provide concentrations along selected 192 193 lines (Fig. 5). Therefore, identification of fractures, growth zoning (e.g., SEM imaging), and 194 inclusions prior to doing the analysis is indispensable to extract information from the dataset (Fig. 5A). An example of a LA ICP-MS traverse done on an arsenopyrite grain from the 195 Meliadine gold district is shown in Fig. 5. The following points are highlighted: 1) Bi, Pb and Co 196 197 show irregular (i.e., bumpy) patterns, which may reflect both oscillatory zoning and a larger 198 scale zoning, the latter reflected by lower elemental abundances at the top of the sample (to fracture 3) versus below fracture 5 at the bottom; 2) Bi, Pb, and Co are depleted between 199 200 fractures 3 and 4 and subsequent enrichment in fractures 4 to 5. We suggest that these features 201 may reflect the role of fluids in processes of coupled dissolution and precipitation (i.e., CDP), which is generally discussed more in the context of silicates (e.g., Putnis, 2002; Putnis and 202 Putnis, 2007; Rubatto et al., 2008; Fu et al., 2009; Martin et al., 2011) whereby elements are 203 204 redistributed; and 3) in contrast, Sb and Mo, which also show some variation in the first part of 205 the traverse (i.e., 25 secs), appear to have been depleted between fractures 4 to 6. Based on these 206 observations, two element associations are noted: 1) Bi-Pb-Co; and 2) Sb-Mo. The latter indicates therefore that, with careful observation, LA ICP-MS traverse data can provide 207

| 208 | information equivalent to element distribution maps but with a much shorter acquisition |
|-----|---|
| 209 | timeframe (e.g., Morey et al., 2008; Gourcerol et al. 2016a). Importantly, we also note that spot |
| 210 | analyses cannot provide such meaningful and detailed information and may in fact average out |
| 211 | some or all of the details noted here. |

It is worth noting that, in a gold exploration context, identifying the sulphide species hosting the gold mineralization must be done prior to any LA ICP-MS map analysis either by spot or traverse mode in order to optimize time and get relevant mapping. Subsequently, it is important to acquire at least one LA ICP-MS map on a host sulphide phase prior to any traverse analysis in order to get a 2D interpretation of zoning and fracture element associations. Moreover, several grains should be analyzed in order to get a qualitative average of element associations.

218 **3.2 Discriminant geochemical plots**

In the preceding section, TSD-based analysis yielded element associations for mineralizing events. Given the existence of the event-specific element associations, the next logical avenue of enquiry is the use of the TSD technique in constraining the nature of the mineralizing fluids. In the following section, several plots specifically designed as part of this study are used to address and potentially constrain the nature of the mineralizing fluid, the stages (i.e., number of events) involved in the mineralizing processes, and the origin of elements (i.e., orogenic versus diagenetic).

3.2.1 Nature of the mineralizing fluid

Examination of LA ICP-MS element distribution maps from several studies in orogenic (e.g.,
Large et al., 2007, 2009, 2011; Lawley et al., 2015) and intrusion-related (e.g., Malartic deposit;
Gao et al., 2015) gold deposits indicates that Co and Ni are always present in the sulphide lattice

where they substitute for Fe, thus representing a primary origin reflecting mineral growth (i.e.,
zonation). When the contents of these latter elements are quantified by element distribution
maps, they potentially offer a proxy that can be used to constrain the source and/or nature of the
fluid; this aspect is further explored below.

234 Compilation of the trace element data for several rock types (i.e., felsic to ultramafic) from the Superior Province (Canada) (cf. caption to Fig. 6 for references) is used to construct the 235 series of diagrams - Co/Ni versus Ni, Co versus Co/Ni, and Ni versus Co - (Fig. 6). The 236 distribution of data in these plots clearly indicates, as expected, variations in the Co and Ni 237 238 contents for different rock suites with the felsic versus ultramafic rocks defining the extreme 239 depletion and enrichment data for the intermediate/lamprophyre rocks defining an intermediary 240 field. These plots are used with the TSD data from the element distribution maps and traverses to constrain the nature of the mineralizing fluids and detect fluid:rock interaction along the fluid 241 242 pathway.

243 The application and use of the Co and Ni data is illustrated in Fig. 7 using data from several pyrrhotite (n = 10) and arsenopyrite (n = 3) samples from, respectively, the Musselwhite deposit 244 and the Meliadine gold district (Gourcerol et al., 2016a), as well as arsenopyrite from the Beaver 245 Dam deposit (Fig. 3). Importantly, these data cannot be compared directly to each other as they 246 247 are for different sulphide phases (i.e., pyrrhotite versus arsenopyrite). However, these samples suggest that the fluids responsible for pyrrhotite deposition at Musselwhite were likely derived 248 from or interacted with ultramafic rocks having intrinsically variable contents of Ni and Co, an 249 250 interpretation considered necessary in order to explain the four distinctly different fields defined 251 in these plots. The latter is of course based on the assumption that factors controlling element partitioning between fluid and sulphide remain constant. A similar result occurs for the 252

253 arsenopyrite data from the Meliadine gold district where two distinct fields are defined. More 254 detailed work at this deposit suggests that the two populations may reflect an early event associated with arsenopyrite crystallization and a second, later event related to the overprinting 255 256 effects of the Trans-Hudson orogeny (Lawley et al., 2015; Gourcerol et al., 2016). In the last example, arsenopyrite data from Beaver Dam defines two distinct fields which suggest that fluids 257 258 must have interacted with different reservoirs, one that is relatively mafic/ultramafic and a 259 second of an overall intermediate nature. Given that the former is dominant, the data suggests the fluids interacted more with a mafic reservoir. In conclusion, we note that without use of the 260 261 approach provided here with the TSD, the distinct fields defined for each of these samples and the interpretations based on this would not have been possible. 262

3.2.2 Upgrading of gold tenor through ore system reactivation

An important aspect of hydrothermal ore deposits is the longevity of some hydrothermal 264 265 mineralizing systems, such as is suggested for porphyry deposits (e.g., Sillitoe, 2010), and the consequent complexity of mineral geochemistry and mineral paragenesis that may result. The 266 occurrence of substantial upgrading of gold tenor in some gold systems (e.g., Wagner et al., 267 2007) is considered to be a result of overprinting events (e.g., Tomkins and Mavrogenes, 2001; 268 Morey et al., 2008; Thomas et al., 2011; Cook et al., 2013; Lawley et al., 2015). Given the 269 importance of recognizing this process in gold mineralized settings, we explore here the use of 270 271 binary Au versus Ag diagrams for an analyzed sulphide color coded by single elements and element associations (Fig. 8) in order to identify spatial distribution and relationship of each 272 other in the Au:Ag sulphide evolution. 273

The TSD for the arsenopyrite from Beaver Dam (Figs. 3, 4), for which elemental distribution maps suggest upgrading of Au as well as Ag and Zn along F1 and F2 fracture sets (Fig. 3), is

| 276 | used to define elemental associations. Note that these element associations cannot be defined |
|-----|---|
| 277 | with spot analyses. These plots show that Au and Ag contents have a consistent relationship with |
| 278 | an Au:Ag ratio of 10:1 and that Co, Ni and Mo-Sb-Se show Au and Ag concentration variations, |
| 279 | which confirm their primary origin (Fig. 8A, B, C). In contrast, two elemental remobilizations |
| 280 | are suggested for Au and Ag contents: 1) the F1 association Bi-Pb-(Cd-In); and 2) the F2 |
| 281 | association Al-Ti-Mn-(V-Cu-Sn-Cr-W)) aligned along F1 and F2 structures (Fig. 8F and G, |
| 282 | respectively). Thus, F1, which is defined as a Bi-Pb-In-Cd association, also shows Ag |
| 283 | enrichment that may be due to its remobilization and/or introduction of an Ag-rich fluid. The |
| 284 | distribution of Se (Fig. 8D) suggests it may also be involved with this remobilization event along |
| 285 | F1, but to a lesser extent. The F2 element association of Al-Ti-Mn-V-Cu-Sn-Cr-W (Fig. 8G) |
| 286 | collectively shows a high aggregate concentration, which is coincident with the main Au versus |
| 287 | Ag trend and is therefore suggestive of upgrading of the precious metals. In Fig. 9, it is apparent |
| 288 | that this elemental association is part of the arsenopyrite (i.e., stoichiometric values) rather than |
| 289 | due to micro-inclusions. This observation is therefore attributed to the coupled dissolution of |
| 290 | earlier arsenopyrite along fractures and precipitation (i.e., CDP process) of a second generation |
| 291 | of arsenopyrite which incorporates these new elements. This is a micro-scale zone refining |
| 292 | process. Also in regards to Fig. 9, the trend of lower Fe values may represent |
| 293 | element remobilization along the F2 fractures, as suggested in Fig. 9H. |

Regarding gold remobilization, the presence of the Ag-, Au-Ag- and Au-rich outliers, which are clearly identified in the dataset, provide evidence for the presence of micro-inclusions in the host sulphide (Fig. 8H). The origin of these Au-rich and Ag-rich micro-inclusions, as shown in Fig. 8H, may involve zone refining for the Au- and Ag-rich grains, as present on the F1 and F2 fractures, or a continuation of the primary mineralizing system such that the Au:Ag ratio is retained. Thus, the use of TSD allows distinction of primary versus remobilization/upgrading ofmineralization in sulphide grains.

It should also be noted in regard to Figure 8 that the continuation of the Au-Ag contents in 301 these plots may be more apparent than "real". For instance, variable size micro-inclusions of 302 303 gold having the same Au: Ag ratio as the invisible Au in the sulphide may contribute to this trend. An example of this would be an inclusion having 1/3 the radius of the laser beam (e.g., 5 μ m 304 versus 15 μ m) would contribute 1/10 of the value of a gold particle the size of the laser beam and 305 for 1/5 would contribute 1/20 of the value. Thus, it is possible that variable sizes of micro-306 307 inclusions can account for the distribution of Au and Ag in the binary plot. Important, however, 308 is that the Au-rich, Ag-poor data points require gold particles with a different Au:Ag ratio and hence origin. 309

310 **3.2.3.** Assessment of element enrichment within pyrite from sediment-hosted gold deposits

Large et al. (2009) used LA ICP-MS generated data from pyrite to assess the origin of four sediment-hosted gold deposits (i.e., Sukhoi Log, Bendigo, Spanish Mountain, and North Carlin Trend). Using these data, they used the chemical signature of pyrite to discriminate early diagenetic versus metamorphic/hydrothermal origin within orogenic- and Carlin-type gold deposits. Based on these latter data and the assigned origin of the various pyrite grains analyzed, Gourcerol et al. (2016a) defined discrimination fields in a binary Ag versus Ni plot (Fig. 10) to distinguish the origin of the analyzed pyrites as well as their associated trace elements.

In order to illustrate the application of this approach and the discrimination diagram (Ag versus Ni), we use TSD for one pyrite grain from the Meadowbank deposit, which is more fully discussed in Gourcerol et al. (2016a). As the various elemental plots show (Fig. 11), this pyrite 321 records a two-stage growth history involving an early diagenetic stage that was modified, 322 probably via coupled dissolution/precipitation processes, during a later metamorphic- (i.e., orogenic-) related event. As seen in Fig. 10, integration of the TSD with the discriminant fields 323 324 suggest the early diagenetic pyrite is enriched in Co as well as Bi, Pb and to some extent Mo, whereas the later metamorphic (i.e., orogenic) stage is characterized by enrichment in Au, Se and 325 Mo. Consequently, this approach provides an important means to ascertain the geochemical 326 nature of gold event(s) and establish the origin of the associated metals during pyrite growth in 327 sedimentary rocks. 328

329

3.3 Geostatistical analysis

A geostatistical-based set of procedures was applied to compilations of individual TSD for 330 the arsenopyrite grain from the Beaver Dam deposit (Fig. 3). In the following sections, the 331 results of this approach are presented to illustrate how this approach can both complement and 332 333 confirm the conclusions reached using the elemental distribution maps and discriminant diagrams. Thus, this methodology allows identification and evaluation of elemental affinities, as 334 well as classification of the affinity groups and their relationship to gold mineralizing processes, 335 and, in doing so, may provide and/or confirm the number and character of gold mineralizing 336 events and their respective element associations. Note that the statistical software XLSTAT 337 (2014, Addinsoft, Inc., Brooklyn, NY, USA) was used as a seamless add-on to MS Excel[®]. 338

339

3.3.1. Agglomerative hierarchical clustering

In order to discriminate groups that define element associations in a given sulphide grain, an 340 exploratory agglomerative hierarchical clustering (AHC) can be performed. In this approach, 341 342 usually presented as a dendrogram plot (Fig. 12), data processing leads to a progressive clustering of elements by a single linkage based on their degrees of similarity (i.e., 343

concentration) calculated after normalizing the data (i.e., using the same scale). The results are
presented as a matrix of element(s) versus element Pearson correlation coefficients (McInnes et
al., 2008), in this case using the TSD as seen below.

Using the results for the single arsenopyrite grain identified above, the resultant dendrogram plot shows three distinct elemental groups: 1) those associated with the F2 fracture set (see paragenetic sequence in Fig. 4) including Zn-Ag-Au-Cu-Sn-Cr-Al-Mn-W-Ti-V; 2) a primary association or group which includes Co-As-Fe-Ni; and 3) those associated with the F1 fracture set including Mo-Sb-Se-Bi-Pb-Cd-In.

It is noted, however, that this approach gives an overview of element associations and must 352 be used with caution as, at a more refined scale, it is limited in that only the closest relationship 353 354 between elements is illustrated. The consequence of this latter limitation is that minor associations may not be represented, depending on the dataset. Thus, only the highest similarities 355 (i.e., highest correlation values) can be considered as "true" similarities, which for the present 356 357 case (Fig. 12) would include Ag-Au-(Zn), Mo-Sb, Cd-In, Al-Mn and Ti-V associations. In this regard, it is possible that the later remobilization weakens some of the associations, such as Mo-358 Sb-Se (Fig. 4), due to Se being remobilized along F1 to join the Bi-Pb-Cd-In grouping. 359 Similarly, the Au-Ag-Zn element association reflects remobilization along the late F2 set from 360 the primary crystallization. 361

362

3.3.2. Exploratory statistical methods

Exploratory statistical methods are another approach commonly used in geostatistical studies to analyze data sets and define variables into statistical factors with visual methods. Multidimensional scaling (MDS), principal component analysis (PCA), and linear discriminant 366 analysis (LDA) can be applied to the datasets in order to extract a potential series of factorial 367 variables governing distribution of elements within the sulphides as part of the lattice, as microinclusions, or focused along fractures. These methods are performed after a normalization of the 368 369 data (i.e., using the same scale) based on a matrix of elements versus element Pearson correlation coefficient (McInnes et al., 2008) for the TSD. These methods allow improvement over the main 370 limitation of the AHC as several dimensions are graphically represented. In the following 371 discussion, the TSD generated from the elemental mapping of the same arsenopyrite grain (i.e., 372 from Beaver Dam deposit; Figs. 3, 4) is used to illustrate these various statistical means of data 373 374 treatment.

375 3.3.2.1. Multidimensional scaling

The MDS method (Wilkinson et al., 1992; Borg and Groenen, 1997, pp. 1-14), similar to 376 the AHC, uses a distance measurement determined from a matrix of Pearson correlation 377 378 coefficient. Distances are calculated for all the element pairs using standardized data (z-score transformed dataset) to provide a graphical representation of the proximity of elements to each 379 other, using a linear Kruskal loss function reflecting the spatial configuration. Thus, the main 380 factor of discrimination is termed Dimension 1 and the second, Dimension 2. Moreover, 381 elements plotting close together will show a positive correlation which reflects their similar 382 behavior, whereas those elements on opposite sides will show a negative correlation due to 383 dissimilar behavior (e.g., Greenough et al., 2007). 384

The dataset for the selected arsenopyrite defines four distinct groups using MDS (Fig. 13A) which are again interpreted in the context of the elemental mapping and paragenesis (Figs. 3, 4) : 1) a primary association defined by Co-Fe-Ni-As; 2) a second primary association of Mo-Sb-Se; 3) a secondary association related to the F1 fracture event enriched in Pb-In-Cd-Bi; and 389 4) another secondary association related to F2 fractures and composed of W-Ti-V-Al-Mn-Sn-Cu-390 Zn-Ag-Au. Further interpretation of the MDS suggests Dimension 1 may reflect compositions related to fractures whereas Dimension 2 may reflect remobilization/precipitation into micro-391 392 inclusions (Fig. 13A). In addition, as pointed out by Greenough et al. (2007), elements tend to organise following Goldschmidt's geochemical classification (Fig. 13B), thus the MDS method 393 can be used to qualitatively identify various fluids equilibrated with the sulphide grain 394 (Greenough et al., 2007). Using this approach, the data suggest a chalcophile affinity for the fluid 395 giving rise to the elements associated with the F1 fracture set versus a lithophile affinity for 396 397 elements associated with the F2 fracture set. Moreover, the paragenesis of Au can be assessed with MDS combined with LA ICP-MS maps as Au shows a pattern similar to Mo (Fig. 3) along 398 with a similar Goldschmidt classification (Fig. 13B); hence, this infers that these elements share 399 a common temporal relationship and were introduced into this arsenopyrite grain at the same 400 time. The MDS also shows that Au was subsequently remobilized by F1 and F2 events which led 401 to its precipitation and upgrading as invisible to visible micro-inclusions along F1 and F2 (Figs. 402 3L, 13B). 403

404 3.3.2.2. Principal Component Analysis

Like the MDS, the PCA method is a dimension-reduction method applied on a correlation matrix (Pearson coefficient; Grunsky and Smee, 1999) resulting in two-dimensional plots, also referred to as a *correlation circle*. In this plot, elements are distributed as a function of their affinities and dissimilarities (i.e., positive and negative correlations) with respect to two main "factors" which best explain the variability (i.e., PC1 and PC2). These factors represent the main variables within the dataset. In the case of variables positioned close to the center of the circle, it is necessary to check whether the proximities correspond to a "true similarity" with the 412 hypersphere by looking at the correlation circles generated by other factor pairs (i.e., PC1 and 413 PC3). Note that like for the MDS, the PCA method minimizes dimensions but preserves covariance of data, whereas MDS preserves distance between data points (Ghodsi, 2006). 414 The TSD for the same arsenopyrite again shows three to four principal components that 415 416 together account for the variability of the entire dataset; these are named Primary, F1, F2 and F2' (Fig. 14). Based on the defined grouping, we interpret the associations as follows. The PC1 on 417 the positive side of this axis has two groupings, mainly Bi, In, Pb, Cd and Se and to a lesser 418 extent Fe, Sb and Mo. These associations are attributed to element associations which 419 420 characterize the F1 fracture event with slight to strong remobilization along it (Fig. 14A). The 421 best example of this F1 association is Se, and to some extent Ni, which is considered as a primary trace element in the arsenopyrite structure (Figs. 3, 4) that was later remobilized along 422 the F1 vector in the correlation circle. Hence, the PC1 negative axis shows F2 element 423 424 association affinity (e.g., Al, Mn, and V) as they are relatively late in the history of the grain. The 425 positive PC2 axis is composed of the F1 and F2 element associations as well as Ag, Au, Zn and minor Ni, whereas the negative axis is composed of the primary element associations (Fig. 4) 426 427 which suggests PC2 represents the influence of F2 (Fig. 14A). Finally, the positive PC3 axis is composed mainly of Ag, Au and Zn (F2') and suggests that invisible- to visible micro-inclusions 428 occur along the fracture sets (Fig. 14B). Importantly, the position of Ag, Au and Zn along PC1 429 and PC3 suggests they were subsequently remobilized from F1 and F2 as previously suggested 430 (Figs. 3, 4). 431

432 3.3.2.3. Linear Discriminant Analysis

Finally, the LDA method (Fisher, 1936) uses an approach similar to the PCA technique but
varies in the sense that it classifies a dataset into predefined groups such as ranges of

concentration (e.g., Venables and Ripley, 2002, Grunsky et al., 2013) prior to any treatment.
Thus, elements are plotted in two dimensional diagrams based on their discrimination scores
(i.e., F1 and F2; F1 and F3) as a function of their behavior with an explanatory variable such as
Au concentration. Therefore, categorization of Au values as a function of selected elements can
provide insight into which elements are closely associated or unrelated to Au mineralization.
This approach can also be applied to the element associations related to subsequent Au
remobilization (e.g., Wagner et al., 2007).

Thus, application of the LDA approach to the TSD for the same arsenopyrite grain (Fig. 15) 442 shows that Au is strongly associated with Ag and Zn (Fig. 15A, B; note that the latter plot 443 444 superposes Au values on this vector). In addition, it is also seen that Au is highly associated with 445 Al, Mn, Cu, amongst other elements. and weakly associated with Co, Mo and Sb (Fig. 15C, D). Examining the element associations (Fig. 16), it is clear that elevated Au values (i.e., > 100 ppm) 446 447 are most strongly associated with the F2 and F1 features whereas Primary' and Primary element associations, that is Mo-Sb-Se and Co-Ni-Fe-As respectively, show a much weaker Au tenor 448 (Fig. 16; from b.d.l. to <100 ppm). 449

In summary, the LDA plots clearly demonstrate that the gold mineralization was likely initially concentrated as invisible Au in the arsenopyrite (i.e., within its lattice) and subsequently remobilized during F1 and F2 events along with Ag and Zn resulting in Au upgrading as microinclusions along the fracture sets.

4. Limitations of LA ICP-MS element distribution analyses and TSD compilation

The data analysis approach described herein, although demonstrably powerful, is not without
limitations. These limitations broadly fall into one of two categories: technological and
methodological, and are discussed below with possible solutions.

Technically, as noted earlier by Ulrich et al. (2009), a main limitation of all LA ICP-MS 458 element distribution analyses (e.g., maps or traverses) is the beam size (typically > 5 um), which, 459 depending on the geometry of the sample and its internal chemical and physical structure, 460 inevitably results in mixed signals. Chemical and physical variation can also occur in the third 461 dimension, down into the specimen, as the particular laser configuration and target mineral will 462 463 yield a certain depth of ablation. The latter arise, for example, from low- and high concentration zones and inclusion(s). This issue cannot be forestalled because the size and shape of such 464 features are variable and not known *a priori*. The only possible solution is to remap fine-scale 465 466 features using a smaller beam size, but even so, the resolution is limited by the smallest beam sizes achievable and the sensitivity and acquisition speed are reduced accordingly. Additionally, 467 the sequential nature of common quadrupole ICP-MS instruments means that the proposed data 468 analysis procedures compare elements that can be derived from different (albeit very close) 469 physical locations. For example, suppose Co and Au are being compared and they are analyzed 470 250 ms apart in the quadrupole cycle; if the laser is firing at 8 Hz (i.e. every 125 ms) the 471 472 corresponding Co and Au measurements are related to different material. Often this is insignificant because the comparison is being made from locations belonging to the same feature 473 474 or domain (e.g., growth zones). However, for micro-inclusions, it can be problematic. This issue affects all instruments having a duty cycle comparable to the laser pulsing period, and is best 475 solved with simultaneous or near-simultaneous detection instruments such as the SpectroMS 476

(Ardelt et al., 2013) or Tofwerks icpTOF (Gundlach-Graham et al., 2015; Burger et al., 2015).
Additional technological advancements that will benefit the type of data analysis conducted in
this study include: faster/more sensitive quadrupole ICP-MS instruments (e.g., the latest
generation of Thermo and Agilent instruments can collect data ~ 100 times faster than the
instrument employed in this study), improved cell designs (i.e. faster wash in/out), and rapid
aerosol transport (e.g., Teledyne's Aris), which all contribute to improved spatial resolution and
analysis time.

Limitations arising from the analytical method employed here (i.e., traverses) include 484 resampling previously ablated material and the accuracy and statistical significance of individual 485 486 time-slices of data. The issue of resampling previously ablated material (i.e. as ejecta or 487 condensate) is particularly prevalent when dealing with features with strong concentration contrasts. A similar phenomenon is encountered at grain boundaries between minerals with 488 489 different major element compositions. This issue can be avoided by mapping with a series of adjacent spots (e.g., Woodhead et al., 2007; Lawley et al., 2015) that can be "pre-cleaned" by a 490 pulse or two of the laser, but this method is typically more time consuming and costly than 491 492 rastering. Individual traverses of a rastered map can also be pre-cleaned, but this only eliminates contamination from adjacent traverses, not material redeposited ahead of the laser on the current 493 path. There are several issues relating to the accuracy of data acquired and processed as 494 described herein. Firstly, in our opinion, the quantification method employed here is an 495 improvement over simply forcing one internal standard everywhere, but still leaves certain parts 496 497 of the map as semi-quantitative (e.g., fractures/cracks, inclusions, other minerals). This could be improved by analyzing all the major elements expected in the mapped area and normalizing to 498 100% as an internal reference (Gagnon et al., 2008; Liu et al., 2008) or by using Iolite's 499

"MinMapping" feature (Paul et al., 2014). Both options improve the accuracy in other phases, 500 501 but do not necessarily improve the accuracy for inclusions and other features that are smaller than the beam, which can be underestimated or overestimated depending on the ablation 502 503 efficiency and geometry. Moreover, normalizing to 100% requires that all the major elements be known and analyzed prior to analyzing samples, which can be problematic when keeping the 504 suite of analytes as small as possible. Secondly, the use of a silicate glass (NIST610) as a 505 reference material for sulphide analyses can be problematic as melting and fractionation are 506 expected to limit the accuracy. One study found that the use of NIST610 as an external reference 507 508 for sulphide analyses with a 193 nm wavelength laser resulted in concentrations generally within 20% of the accepted value (Wohlgemuth-Ueberwasser and Jochum, 2015), which is commonly 509 satisfactory for the type of study conducted here. Unfortunately, there are no widely available 510 511 sulphide reference materials that contain an appropriate suite of trace elements that can be used as an alternative at present. New reference materials may one day become available to alleviate 512 this problem, but in the meantime, the use of fs-lasers could minimize the ablation differences 513 between silicate glass and sulphides (Wohlgemuth-Ueberwasser et al., 2015). 514

Although there are some limitations to the current approach as discussed above, the examples used throughout this study demonstrate that, despite these limitations, the expected element associations are easily identified, suggesting that the data analysis procedure is effective. Different samples, containing different internal chemical and physical structures may be less or more troublesome, and therefore, selection of appropriate samples is a crucial step in this technique. In the event of the latter, there are several technological advances available now that could improve this procedure in terms of quality and speed, and it is expected that once that technology is more widely adopted among the geoanalytical community, this type of dataanalysis will be even more powerful.

524 **5.** Conclusions

We have shown in this study that, using LA ICP-MS data generated from both elemental 525 526 mapping and traverses of sulphide grains, considerable insight can be gained regarding the 527 development of the chemical domains in the host and in particular the nature of gold 528 mineralization. By processing these data in the multiple ways presented here, it is possible to 529 enhance our understanding of the nature and origin of gold mineralizing events and, to this end, 530 we have introduced several new concepts to be considered in such studies. The first step, where 531 possible, is to use elemental maps to establish both elemental paragenesis and to identify 532 elemental associations. Given that the map acquisition is both time and cost intensive and that all grains are suitable for such study, we have also explored and shown that traverse data can also 533 provide insight into the nature of gold mineralization. In order to further understand the 534 chemistry of the analyzed material, we also propose innovative procedures for the treatment of 535 time-resolved LA ICP-MS data (i.e., time-slice datasets (TSD)). The use of the TSD in a variety 536 537 of new discriminant plots (i.e., Co versus Co/Ni; Au versus Ag; Ag versus Ni) and treatment in three-dimensional element plots versus normal binary plots provides possible insight into the 538 origin of the mineralizing fluids and timing of gold mineralization (i.e., diagenetic versus 539 540 metamorphic/orogenic).

Lastly, conventional geostatistical tools (e.g., MDS, PCA, and DA), which are routinely
applied to other geochemical datasets, have been introduced and their application to TSD
illustrated. The latter provides further insight into the nature of gold events and also validation of

the conclusions inferred from the other methods used. Thus, we suggest that the integration of the proposed protocol of treating LA ICP-MS data on carefully selected sulphide grains with other geochemical tools commonly applied to gold deposit studies and gold exploration (e.g., fluid inclusions, isotopes) provides the means to better understand the nature and origin of mineralized settings and better discriminate gold mineralizing events and systems.

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| 923 | 8. Figure Captions |
| 924 | Fig.1. LA ICP-MS element maps for an arsenopyrite grain from the Dufferin deposit, Meguma |
| 925 | Terrane (Nova Scotia, Canada). The maps show the distribution of Co (left column), Sb (middle |
| 926 | column) and Ag (right column) with four different color scales: 1) logarithmic concentration |

scale between the minimum and maximum values (A, B, C); 2) empirical cumulative distribution

928 function scale between the minimum and maximum values (D, E, F); 3) linear concentration

| 929 | scale between the data median +/- 3 standard deviations (G, H, I); and 4) linear concentration |
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| 930 | scale between minimum/maximum values (J, K, L). Note that straight lines, which are mainly |
| 931 | seen in the Sb maps, correspond to earlier LA traverses done on the grain. The X and Y axis |
| 932 | scale is given in microns. |
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| 939 | Fig. 2. Concentrations (in ppm) for selected elements of varying contents plotted against elapsed |
| 940 | time in multiples of single cycles (i.e., 0.3 seconds) of the quadrupole (i.e., 0.3, 1.2, 3 and 5 |
| 941 | seconds) for traverses. As discussed in the text, the time for a single cycle depends on the |
| 942 | number of analytes and dwell time per analyte. A) to C) Element concentrations of Al, Ni and Co |
| 943 | in the BHVO2G standard. D) Element concentration of Au in the Po725 standard (Sylvester et |
| 944 | al. 2005). Note the dashed black line and darker grey field represent the expected values (i.e., Al |
| 945 | in BHVO2G 72000 \pm 1000 ppm; Ni in BHVO2G 116 ± 14 ppm; Co in BHVO2G 44 \pm 4 ppm; Au |
| 946 | in Po725 44.5 \pm 2.5 ppm), whereas lighter grey fields refer to \pm 20% of accepted values. Thus, |
| 947 | this figure illustrates that the shorter time of acquisition is the best option to identify micro- |
| 948 | inclusions of, for example, sulphides. Note that the concentration values for individual time |
| 949 | slices are usually within 20% of the accepted value. |

951 Fig. 3. LA ICP-MS element maps showing the distribution of selected elements in part of an arsenopyrite grain from the Beaver Dam deposit, Meguma Terrane (Nova Scotia, Canada). A) to 952 J) Elemental maps plotted as concentrations scaled between the data median +/- 3 standard 953 954 deviations (in ppm). K, L) Element maps plotted as logarithmic concentrations scaled between the minimum and maximum values (in weight %). Note the following features illustrated in the 955 956 maps: 1) the black dashed lines shown in A represent primary growth zoning; 2) the white lines 957 represent two sets of fractures referred to as F1 (dashed lines) or F2 (dotted lines). Note that the straight lines, which are mainly seen in the Sb maps, correspond to earlier AL traverses done on 958 the grain. Scale is given in microns. 959

| 962 | Fig. 4. An elemental paragenesis for the arsenopyrite grain from the Beaver Dam deposit (Nova |
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| 963 | Scotia, Canada) shown in Fig. 3. Note that the elemental paragenesis was derived based on the |
| 964 | distribution of the selected elements in the elemental maps for this grain in addition to the |
| 965 | presence of fractures (F1, F2). |
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| 968 | Fig. 5. Quantitative LA ICP-MS elemental profiles (in ppm) acquired from a traverse across an |
| 969 | arsenopyrite grain from the Meliadine gold district (Nunavut, Canada). A) Reflected light |
| 970 | photomicrograph of the analyzed arsenopyrite grain. The solid black line represents the traverse |
| 971 | whereas the numbered (1 to 6) dashed black lines represent micro-fractures identified based on |
| 972 | reflect light observations. B) concentration profiles for selected elements along the traverse |
| 973 | shown in image A. Note that the dashed black lines refer to the micro-fractures identified in |
| 974 | image A. Abbreviations: Aspy: Arsenopyrite; Po: Pyrrhotite. |
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| 978 | |
| 979 | Fig. 6. Binary plots showing the distribution of Co, Ni and Co/Ni for various rock types from the |
| 980 | Archean Superior Province (Canada): A) Co/Ni versus Ni; B) Co versus Co/Ni; and C) Ni versus |
| 981 | Co. The distribution of the elements and elemental ratios in these plots define the corresponding |

982 compositional fields: pink corresponds to felsic, light green to mafic, light blue to ultramafic, 983 light purple to lamprophyre and the dashed field to intermediate rocks. Data used to define the fields are from the following sources: Arndt and Nesbitt, 1982; Arndt, 1986; Barrie et al., 1991a, 984 b, 1999; Beswick, 1983; Canil, 1987; Dinel et al., 2008; Dostal and Muller, 2013; Fan and 985 Kerrich, 1997; Feng and Kerrich, 1992; Finamore et al., 2008; Fralick et al., 2009; Gaboury and 986 Pearson, 2008; Goldstein and Francis, 2008; Hollings and Wyman, 1999; Hollings and Kerrich, 987 1999a, b; Hollings et al., 2000; Hollings and Kerrich, 2000; Innes, 1978; Kerrich et al., 1999, 988 2008; Kitayama and Francis, 2014; La Fleche et al., 1992; La Fleche and Dupuy, 1992; Lafrance 989 990 et al., 2000; Lahaye et al., 1995; Lahaye and Arndt, 1996; Maurice et al., 2003, 2009; Merci-Langevin et al., 2007; Nesbitt et al., 2009; Parks et al., 2014; Picard and Piboule, 1986; Piercey 991 et al., 2008; Polat et al., 1999, 2012; Polat and Kerrich, 1999; Pollat and Münker, 2004; Pollat, 992 993 2009; Rinne and Hollings, 2013; Sproule et al., 2002; Stone et al., 1993, 1995, 1996; Stone and Stone, 2000; Thurston and Fryer, 1983; Ujike and Goodwin, 1987; Ujike et al., 2007; Wyman 994 and Kerrich, 1993; Wyman and Kerrich, 1993, 2009; Wyman and Hollings, 1998; Wyman, 1999; 995 Wyman et al., 2000, 2002. 996

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Fig. 7. Distribution of elements (Co, Ni) and elemental ratios (Co/Ni) based on TSD from LA
ICP-MS analyses for both traverse and mapping modes of sulphides from the Musselwhite,
Meliadine and Beaver Dam deposits. The data are shown in the same discriminant diagrams as
seen in the Figure 6. A) Co/Ni versus Ni; B) Co versus Co/Ni; and C) Ni versus Co. The data
plotted are based on 10 pyrrhotite samples from the Musselwhite deposit, 3 arsenopyrite grains

1004 from the Meliadine gold district, and a single element distribution map for arsenopyrite from the1005 Beaver Dam deposit.

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Fig. 8. Binary element plots (in ppm) of Au versus Ag based on the TSD for the mapped 1008 1009 arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada); there are approximately 1010 39800 data points represented in this diagram. Note that in Figures A to G the data points are 1011 colour coded (i.e., cold to hot) in order to reflect enrichment in either an element or elemental 1012 association; see text for further discussion. Figure H is a representative summary plot showing an 1013 interpretation of the elemental distributions. Note the dashed red line in panels A to G represents 1014 the detection limit for Au that the F1 element association consists of Bi-Pb-In-Cd, and the F2 1015 element association consists of Al-Ti-Mn-V-Cu-Sn-Cr-W.

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Fig. 9. Binary elements plots of Au (ppm) versus Fe (wt.%) based on the TSD for the mapped arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada) and colored by element associations. These diagrams demonstrate that element associations are characterized mainly by stoichiometric values of Au and Fe suggesting they are part of the arsenopyrite grain and not related to late remobilization or inclusions. Note a small group of points shows low Fe and high Fe element association contents which may illustrate remobilization along F2 fractures.

| 1026 | Fig. 10. Binary plot of Ag versus Ni (in ppm) in pyrite grains of both early diagenetic and |
|------|--|
| 1027 | metamorphic/hydrothermal origin from four sediment-hosted gold deposits reported by Large et |
| 1028 | al. (2009) - Sukhoi Log (Russia), Bendigo (Australia), Spanish Mountain (Canada) and North |
| 1029 | Carlin Trend (USA). Based on the distribution of Ag and Au in this diagram, discrimination |
| 1030 | fields have been defined for early diagenetic, metamorphic/hydrothermal orogenic, and Carlin- |
| 1031 | type (see Gourcerol et al., 2016a for further discussion). |
| 1032 | |
| 1033 | |
| 1034 | Fig. 11. Selected multi-element plots of Ag versus Ni (in ppm) plus an additional element for LA |
| 1035 | ICP-MS TSD for pyrite from the Meadowbank deposit (Nunavut, Canada). Note that additional |
| 1036 | elemental data are plotted using color codes for various concentrations (i.e., cold to hot) and that |
| 1037 | these define two elemental groupings, as discussed in the text. The data are also plotted in the |
| 1038 | context of discriminant fields (see Fig. 10) for different sediment-hosted gold deposits (i.e., |

1039 diagenetic versus metamorphic/hydrothermal orogenic versus Carlin-type).

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| 1044 | Fig. 12. A | dendogram | plot for | agglomerative | hierarchical | clustering | (AHC) | statistical | analysis | of |
|------|------------|-----------|----------|---------------|--------------|------------|-------|-------------|----------|----|
|------|------------|-----------|----------|---------------|--------------|------------|-------|-------------|----------|----|

- selected elements for TSD for the arsenopyrite grain analyzed from the Beaver Dam deposit
- 1046 (Nova Scotia, Canada). Note the three distinct colors defined the three elemental groupings
- 1047 based on their element affinities.

1050

1051 Fig. 13. Results of multidimensional scaling (MDS) analysis of selected elements for TSD of an 1052 arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada). The plots show the 1053 relationship between selected elements (A) and their respective Goldschmidt classification (B). 1054 Note that Dimension 1 and Dimension 2 have been interpreted, respectively, in the context of 1055 fracture controlled enrichment and remobilization due to coupled dissolution/precipitation. Elements are written in color as defined by the dendrograms in the AHC diagram (Fig. 12) 1056 whereas field colors correspond to the element association defined from the paragenetic 1057 sequence and element map (Figs. 2, 3). For instance, Mo is part of the Primary' event according 1058 1059 to the MDS diagram but refers also to F1 element association by the dendrogram as it was slightly remobilized along the fracture set. Note Primary' and Primary refer to the paragenesis in 1060 1061 Figure 3.

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Fig. 14. Principal component analysis (PCA) diagrams of selected elements based on TSD for an arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada) as a function of PC1 and PC2 (A) and PC1 and PC3 (B). Note the elements are written in the color as defined by the dendrograms in the AHC diagram (Fig. 12) whereas field colors correspond to the element associations defined from the paragenetic sequence and element map (Figs. 2, 3).

| 1070 | Fig. 15. Linear discriminant analysis (LDA) diagrams of selected elements for the TSD for an |
|------|---|
| 1071 | arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada). Note the elemental |
| 1072 | associations with Au are discriminated by Au concentration (in ppm) as a function of factors 1 |
| 1073 | and 2 in plots A and B and factors 1 and 3 in plots C and D. See the text for further details. Also |
| 1074 | note that F1 refers to the factor explaining 44.92% of the trace element variation and F2 explains |
| 1075 | 37.07% of the trace element variation. The plots in A) and C) illustrate the variable and vector |
| 1076 | correlation circles whereas B) and D) represent Au concentration groups as a function of the |
| 1077 | element vectors (e.g., higher Au concentration associated with Ag and Zn). |

| 1081 | Fig. 16. Linear discriminant analysis (LDA) diagrams of selected element associations for TSD |
|------|--|
| 1082 | for the arsenopyrite grain from the Beaver Dam deposit (Nova Scotia, Canada). Note that the |
| 1083 | data are discriminated by Au concentration (in ppm) for the function factors 1 and 2 (in plots A |
| 1084 | and B) and factors 1 and 3 in plots C) and D. In addition, plots A and C illustrate the variable and |
| 1085 | vector correlation circles whereas plots B and D represent Au concentration groups as a function |
| 1086 | of the element association vectors (e.g., higher Au concentration associated with factors F1 and |
| 1087 | F2). Note the element associations are based on the paragenesis sequence (Fig. 3) and "Primary" |
| 1088 | refers to the Co-Ni association whereas, "Primary" " refers to the Mo-Se-Sb element association. |

| Laser ablation system | |
|---------------------------------|---|
| Instrument | Resonetics RESOlution M-50 |
| Laser type | ArF excimer |
| Wavelength | 193 nm |
| Pulse duration | 20 ns |
| Repetition rate | 5 - 7 Hz |
| Spot diameter | 9 - 48 um |
| Speed | 3 - 18 um/s |
| | (typically $1/3$ to $1/2$ the beam diameter/s) |
| Energy density | $4 - 5 \text{ J/cm}^2$ (mesured) |
| Carrier gas (He) | 650 ml/min |
| Additional gases | 6 ml/min N ₂ |
| Primary reference material | NIST610 |
| Secondary reference material(s) | NIST612, BHVO2g, Po725 |
| • | (some combination thereof) |
| Mass spectrometer | |
| Instrument | Thermo X-Series II |
| Plasma RF power | 1440 - 1490 W |
| Plasma gas flow | 730 - 800 ml/min Ar |
| Isotopes analyzed for D4-A and | ²⁷ Al. ⁴⁷ Ti. ⁵¹ V. ⁵² Cr. ⁵⁵ Mn. ^{56, 57} Fe. ⁵⁹ Co. ⁶⁰ Ni. ⁶⁵ Cu. ⁶⁶ Zn. ⁷⁵ As |
| LC-12-16 maps | ⁸² Se, ⁹⁵ Mo, ¹⁰⁷ Ag, ¹¹¹ Cd, ¹¹³ In, ¹¹⁸ Sn, ¹²¹ Sb, ¹⁸² W, ¹⁹⁷ Au, |
| | ²⁰⁸ Pb, ²⁰⁹ Bi |
| | 10 ms dwell time per analyte |
| | total duty cycle of 276 ms |
| Isotopes analyzed for other | ³³ S, ⁴⁵ Sc, ⁴⁷ Ti, ⁵⁵ Mn, ⁵⁶ Fe, ⁵⁹ Co, ⁶⁰ Ni, ⁶⁵ Cu, ⁶⁶ Zn, ⁷⁵ As, ⁸² Se, |
| traverses and maps | ⁹⁵ Mo, ¹⁰¹ Ru, ¹⁰³ Rh, ¹⁰⁵ Pd, ¹⁰⁷ Ag, ¹¹¹ Cd, ¹¹⁵ In, ¹¹⁸ Sn, ¹²¹ Sb, |
| | ¹²⁵ Te, ¹⁸² W, ¹⁸⁹ Os, ¹⁹³ Ir, ¹⁹⁵ Pt, ¹⁹⁷ Au, ²⁰² Hg, ²⁰⁵ Tl, ²⁰⁸ Pb, ²⁰⁹ Bi |
| | ²³⁸ |
| | 10 ms dwell time per analyte except Au, which was 30 ms |
| | total duty cycle of 403 ms |
| | |

Table 1: LA-ICP-MS Instrumentation and Acquisition Parameters

 $\frac{\text{U}^{*}/\text{Th}^{+}}{\text{ThO}^{*}/\text{Th}^{+} \text{ and } \text{U}^{*}/\text{Th}^{+} \text{ were determined on NIST612 during instrument tuning}}$

< 0.6 %

 $\mathrm{ThO}^{+}/\mathrm{Th}^{+}$









- Major - Minor Trace





























