2	Distal Pb-Zn-Ag veins associated with the world-class Donggou porphyry
3	Mo deposit, southern North China craton
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#### 21 Abstract

The southern North China craton hosts numerous world-class porphyry Mo and Pb-Zn-Ag vein deposits. Whether or not the Pb-Zn-Ag veins are genetically associated with the porphyry Mo system remains contentious. Here we focus on the genetic relationships between the Sanyuangou Pb-Zn-Ag vein deposit and the world-class Donggou porphyry Mo deposit, and discuss the potential implications from the spatial and temporal relationships between porphyry and vein systems in the southern North China craton.

28 At Sanyuangou, vein-hosted sulfide mineralization mainly comprises pyrite, sphalerite, and 29 galena, with minor chalcopyrite, pyrrhotite, bornite, tetrahedrite, covellite, polybasite and argentite. 30 The mineralization is hosted by a quartz diorite stock, which has a zircon U-Pb age of  $1756 \pm 9$  Ma. 31 However, sericite from alteration selvages of Pb-Zn-Ag sulfide mineralization yields a well-defined 32  $^{40}$ Ar/ $^{39}$ Ar plateau age of 115.9 ± 0.9 Ma. Although nominally younger, the sericite  $^{40}$ Ar/ $^{39}$ Ar age is 33 similar to the age of the nearby Donggou porphyry Mo deposit (zircon U-Pb age of  $117.8 \pm 0.9$ ; 34 molybdenite Re-Os ages of  $117.5 \pm 0.8$  Ma and  $116.4 \pm 0.6$  Ma). Pyrite from Donggou has elevated 35 contents of Mo and Bi, whereas pyrite from Sanyuangou is enriched in Cu, Zn, Pb, Ag, Au, and As. 36 This trace element pattern is consistent with metal zonation typically observed in porphyry related 37 metallogenic systems. Pyrite grains from Sanyuangou have lead isotopes overlapping those from Donggou (17.273-17.495 vs. 17.328-17.517 for 206Pb/204Pb, 15.431-15.566 vs. 15.408-15.551 for 38 39 <sup>207</sup>Pb/<sup>204</sup>Pb, and 37.991-38.337 vs. 38.080-38.436 for <sup>208</sup>Pb/<sup>204</sup>Pb). Collectively, the geological, 40 geochronological, and geochemical data support a magmatic-hydrothermal origin for the Sanyuangou 41 Pb-Zn-Ag deposit and confirm that the Pb-Zn-Ag veins and the Donggou Mo deposit form a porphyry-42 related magmatic-hydrothermal system.

Given the widespread Pb-Zn-Ag veins and Mo mineralized porphyries in many districts of the southern North China craton, the model derived from this study has broad implications for further exploration of Mo and Pb-Zn-Ag resources in the area.

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47 Key words: porphyry Mo deposit; Pb-Zn-Ag veins; geochronology; trace elements; *in-situ* lead
48 isotopes; Donggou deposit

#### 49 Introduction

50 Porphyry deposits are the world's most important repositories of Cu and Mo (Sillitoe, 2000). They 51 are commonly generated above subduction zones at convergent margins and are associated with 52 contemporaneous calc-alkaline magmas (Richards 2003; Cooke et al., 2005; Seedorff et al., 2005). 53 Some deposits, especially porphyry Mo deposits in continental collision setting, are associated with 54 high K calc-alkaline to shoshonite magmas (e.g., Chen et al., 2004, 2016; Li et al., 2007; Mao et al., 55 2010). Porphyry-related base metal vein and replacement mineralization may form economically 56 important Zn, Pb, Cu, Ag, and Au deposits proximal or distal to porphyry Cu and Mo deposits (Lang 57 and Eastoe, 1988; Sillitoe, 2010; Catchpole et al., 2015). A continuum of porphyry-related 58 mineralization styles was proposed by Sillitoe (1973) to explain the close spatial relationship of various 59 hydrothermal deposit types (porphyry, skarn, base and precious metal veins, and epithermal deposits) 60 in many porphyry systems. Geochemical footprints of the porphyry system commonly comprise a 61 district-scale metal zonation of Cu-Mo through Zn-Pb to Ag-Au (Emmons, 1927; Jones, 1992; Dilles 62 and Einaudi, 1992; Babcock et al., 1995; Seedorff et al., 2005; Sillitoe, 2010). Such metal zonation, 63 when properly recognized, may provide useful exploration vectors to end-member mineralization styles 64 in a porphyry related metallogenic system. A comprehensive geochronological and geochemical study 65 is critically important in constraining a possible genetic link between porphyry and peripheral Zn-Pb-66 Cu-Ag-Au mineralization (e.g., Lawley et al., 2010; Schütte et al., 2012; Catchpole et al., 2015).

67 Large areas of the southern North China craton (NCC) host several giant mineral concentrations 68 (Fig. 1). The Xiaoqinling district has been China's second largest gold producer in the last three 69 decades (Mao et al., 2002; Li et al., 2012a, 2012b). To the east, the Xiong'ershan district has 70 traditionally been an important gold producer (Chen and Fu, 1992; Fan et al., 2000), but has also 71 recently become the most important silver producer and a significant Zn-Pb source along the southern 72 NCC (Chen et al., 2004; Mao et al., 2006; Li et al., 2010, 2013b). The Luonan-Lushi, Luanchuan and 73 Waifangshan districts host several world-class porphyry Mo deposits (Fig. 1; Li et al., 2005; Ye et al., 74 2006; Mao et al., 2011). Molybdenite Re-Os dating has revealed two pulses of porphyry intrusion and 75 porphyry-skarn Mo mineralization at 148-138 Ma and 131-112 Ma, respectively (Mao et al., 2008). 76 These Mo deposits are unrelated to a convergent plate margin; rather they are located in 77 intracontinental settings and are associated with high K calc-alkaline or alkaline felsic magmas induced by thinning and destruction of lithosphere beneath this craton during the Late Jurassic to Early
Cretaceous (Mao et al., 2008, 2010, 2011; Pirajno and Zhou, 2015).

80 In the last two decades, numerous Pb-Zn-Ag veins have been discovered in some of the Mo-81 mineralized districts, such as the Lengshuibeigou deposit in the Luanchuan ore field (Fig. 1; Yan, 2004; 82 Qi et al., 2007; Wang et al., 2013). Genesis of these Pb-Zn-Ag veins is debated (e.g., Chen and Fu, 83 1992; Chen et al., 2004; Mao et al., 2006, 2011; Yao et al., 2008; Li et al., 2013b). Previous studies 84 interpret the Pb-Zn-Ag veins to be a porphyry-related (Ye, 2006; Mao et al., 2006, 2009, 2011; Wang 85 et al., 2013), or an orogenic-type system associated with the Triassic orogenic deformation involving 86 the continental collision between the North China and Yangtze cratons (Chen et al., 2004; Qi et al., 87 2007; Yao et al., 2008).

88 The Fudian ore field in southern NCC (Fig. 1) hosts the giant Donggou porphyry Mo deposit and 89 numerous Pb-Zn-Ag vein deposits including Sanyuangou, Laodaizhanggou, and Wangpingxigou (Fig. 90 2). Current geochronologies for the Donggou porphyry Mo deposit are non-conclusive based on a 91 SHRIMP zircon U-Pb age of  $112 \pm 1$  Ma for the Donggou porphyry and molybdenite Re-Os ages of 92  $116.5 \pm 1.7$  to  $114.1 \pm 1.4$  Ma for Mo-bearing quartz and K-feldspar veins (Ye et al., 2006; Mao et al., 93 2008). In addition to the Donggou porphyry, several large granitoid intrusions and a number of diorite 94 dikes occur in the Fudian ore field (Fig. 2). Most Pb-Zn-Ag veins are in close proximity of or hosted by 95 these intrusions (Fig. 2), but the genetic relationship to the magmatic events has not been determined. 96 Alteration selvage sericite associated with a Pb-Zn-Ag vein of the Sanyuangou deposit (Fig. 2) has an 97  $^{40}$ Ar/ $^{39}$ Ar plateau age of 110.1 ± 9.2 Ma (Ye, 2006), which is roughly consistent with a sphalerite Rb-Sr 98 isochron age of  $117 \pm 27$  Ma for the Wangpingxigou Pb-Zn-Ag deposit (Fig. 2; Yao et al., 2010). The 99 large uncertainties of the <sup>40</sup>Ar/<sup>39</sup>Ar and Rb-Sr ages for the Pb-Zn-Ag deposits and the discordance 100 between the U-Pb and Re-Os ages for the Donggou porphyry Mo deposit necessitates further precise 101 geochronological coupled with geochemical studies to evaluate a possible genetic link between 102 porphyry Mo and vein Pb-Zn-Ag mineralization.

Here we firstly present a comparative geochronology study (U-Pb, Re-Os, and <sup>40</sup>Ar/<sup>39</sup>Ar) for the Donggou porphyry Mo deposit and the Sanyuangou Pb-Zn-Ag vein deposit to assess their temporal relationship. We further analyze trace elements of pyrite from both deposits using laser ablation ICP-MS to investigate any geochemical affiliation between the two systems. Lastly, we use *in-situ* lead isotopes of pyrite to evaluate any genetic link between the Pb-Zn-Ag and porphyry Mo systems.

Synthesis of existing geological and geochronological data provides a metallogenic model that may prove a useful exploration guide for porphyry Mo and Pb-Zn-Ag veins in the whole southern NCC.

#### 110 Geologic setting

111 The Fudian ore field is located in the Waifangshan district of the southern NCC (Fig. 1). The 112 NCC consists of the Western and Eastern Blocks that are separated by the Trans-North China Orogen 113 (TNCO) formed during collision between the two blocks at ca. 1.85 Ga (Fig. 1). This collision resulted 114 in the final amalgamation and stabilization of the craton (Zhao et al., 2001). The basement of the NCC 115 is dominated by Archean to Paleoproterozoic high-grade metamorphic rocks that are variably overlain 116 by Mesoproterozoic to Late Paleozoic unmetamorphosed marine sedimentary rocks (Zhao et al., 2000, 117 and references therein). Although, the NCC behaved as a coherent, stable continental block from the 118 Late Paleoproterozoic to the Late Paleozoic (Yang et al., 1986; Zhai, 2010), its margins were 119 repeatedly affected by Late Paleozoic to Early Mesozoic subduction and orogenesis (Chen and Fu, 120 1992). During the Late Mesozoic, the NCC witnessed extensive magmatism (Wu et al., 2005; Mao et 121 al., 2010), exhumation of numerous metamorphic core complexes (Fig. 1; Wang and Zhang, 1999; 122 Zhang and Zheng, 1999; Shi et al., 2004), and development of intracontinental rift-basins (Ren et al., 123 2002). These tectonic signatures are interpreted to be a consequence of tectonic reactivation or 124 lithospheric destruction of the eastern NCC, induced by the westerly subduction of the paleo-Pacific 125 plate in Early Cretaceous (e.g., Zhu et al., 2012).

126 The southern NCC is dominated by the Late Archean to Early Paleoproterozoic Taihua and 127 Xiong'er Groups (Fig. 1). The Taihua Group consists of amphibolite facies metamorphic rocks, mainly 128 including amphibolite, felsic gneiss, khondalite, migmatite, and metasedimentary rocks intercalated 129 with mafic to ultramafic rocks (Hu et al., 1988; Chen and Fu, 1992). The amphibolites have a whole-130 rock Sm-Nd isochron age of  $2,542 \pm 57$  Ma (Zhou et al., 1998), whereas the felsic gneisses and 131 khondalite series have zircon U-Pb ages of 2.6-2.3 and 2.2-2.18 Ga, respectively (Xu et al., 2009; Li et 132 al., 2015). The Xiong'er Group is distributed widely in the Xiaoshan, Xiong'ershan, and Waifangshan 133 districts. It is composed of volcanic rocks ranging in composition from basalt to rhyolite, but 134 dominated by andesite, with minor intercalations of clastic rocks (Zhao et al., 2002; Peng et al., 2008). 135 In-situ zircon U-Pb dating indicates that rocks of the Xiong'er Group mainly erupted between 1.8 and 136 1.75 Ga (Zhao et al., 2004; He et al., 2009; Zhao et al., 2009). Mesozoic and Cenozoic sedimentary
137 rocks are locally present in the area (Fig. 1).

138 To the south, the southern NCC is separated from the North Qinling Terrane by the NW-oriented 139 Luanchuan Fault (Fig. 1), which was generated during the Mesozoic continental collision between the 140 Yangtze and North China cratons forming the Qinling Orogen (Dong et al., 2011). The Machaoying 141 Fault is an important north-dipping regional structure in the southern NCC, extending for more than 142 200 km along an approximately easterly strike (Fig. 1). Interpretation of geophysical data indicates that 143 the Machaoving Fault is a translithospheric structure (Hu et al., 1988). Secondary structures affiliated 144 with the Machaoying Fault are well developed and have been important in the formation and 145 distribution of polymetallic deposits in the area (Fig. 1; Chen and Fu, 1992; Yan et al., 2000; Ma et al., 146 2006).

147 Mesozoic granitoid intrusions are widespread in the southern NCC (Fig. 1). To the west of the 148 Waifangshan district, the Wuzhangshan and Huashan plutons intruded intermediate to acidic rocks of 149 the Xiong'er Group at  $157 \pm 1$  Ma and  $132 \pm 2$  Ma, respectively (Mao et al., 2010). To the south of the 150 district, the Heyu pluton consists of biotite monzonite and granite porphyry formed by multiphase 151 magmatism between 148 and 127 Ma (Gao et al., 2010b; Mao et al., 2010; Li et al., 2012c, 2013a). 152 Adjacent to the Heyu intrusive complex, the Taishanmiao pluton consists of coarse- to medium-grained 153 K-feldspar granite, fine- to medium-grained syenogranite, and fine-grained granite (Qi, 2014), which 154 were emplaced episodically between 125 and 115 Ma (Ye et al., 2008; Qi, 2014; Gao et al., 2014). 155 Compositionally, the Taishanmiao granites are classified as K-rich, aluminous to peraluminous granite 156 and aluminous A-type granite (Ye et al., 2008). A number of Mo-mineralized porphyries occur over the 157 Waifangshan district and adjacent areas (Fig. 1). These porphyries have been dated at 158 to 112 Ma 158 using in-situ zircon U-Pb geochronology (Li et al., 2006; Mao et al., 2010).

159 Geology of the Fudian ore field

160 The Fudian ore field is located in the eastern portion of the Waifangshan district (Fig. 1), 161 immediately to the northeast of the Taishanmiao intrusive complex (Fig. 2). The ore field is dominated 162 by basaltic andesite, andesite, dacite and rhyolites of the Paleoproterozoic Xiong'er Group, which are 163 locally covered by Cenozoic unconsolidated sediments (Fig. 2). Brittle faults are well developed in the

164 Fudian ore field (Fig. 2). The NW-striking Yangping-Wangping Fault (F1) and Jincun-Fudian Fault 165 (F2) are the main structures in the district, extending for more than 20 to 30 km along strike and 166 dipping 70°-80° to the southwest (Huang et al., 1992). Both faults are characterized by well-developed 167 tectonic breccias and cataclasite with intensive sericitization, chloritization, and silicification. 168 Numerous NE-striking faults displace and thus postdate the NW-striking structures (Fig. 2). There are 169 also some minor E-striking faults, which either crosscut or are displaced by the NE- or NW-oriented 170 faults. The E- and NE-striking faults are the major structural hosts of the Pb-Zn-Ag veins in the Fudian 171 ore field (Fig. 2; Huang et al., 1992; Ma et al., 2006).

172 Several granitoid intrusions were emplaced into the volcanic rocks of the Xiong'er Group (Fig. 2). 173 They are compositionally dominated by quartz diorite and quartz monzonite, with minor diorites (Fig. 174 2). The quartz monzonite at Wangpingxigou is enclosed by, rather than intrudes, volcanic rocks of the 175 Xiong'er Group. It intruded the Taihua Group and then covered by the Xiong'er Group. Ages of these 176 magmatic intrusions are undetermined, but a Mesoproterozoic age is speculated by local geologists 177 (HBGMR, 1989). The Donggou granite porphyry is the only Mesozoic intrusion at Fudian, but large 178 plutonic bodies of similar ages crop out to the south of the ore field (Figs. 1, 2). The Donggou porphyry 179 has an exposure of 0.01 km<sup>2</sup> (Fig. 3), but drill holes have revealed that it has a minimum lateral 180 dimension of 1550 m and a vertical extent of >850 m (Fig. 4; Ye et al., 2006). The porphyry has 181 phenocrysts dominated by perthite and quartz that account for 10-15 vol. % of the rocks, whereas the 182 matrix is composed principally of perthite (40-55 vol. %), quartz (20-30 vol. %), plagioclase (10-20 vol. 183 %) and albite (5-15 vol. %) with minor biotite. The rocks are weakly peraluminous with high Si, K, and 184 Ga/Al ratios, and are classified as aluminous A-type granites (Dai et al., 2009). Whole-rock Sr-Nd and 185 zircon Hf isotope data indicate that the porphyry was derived from partial melting of ancient lower 186 crust with minor input of mantle-derived mafic magmas (Dai et al., 2009). Regional gravity and 187 aeromagnetic data indicate a deep-seated plutonic body beneath the Donggou porphyry, likely 188 representing the northward extension of the Taishanmiao intrusion at depth (Ye et al., 2006).

#### 189 Mineralization

Mineralization in the Fudian ore field is largely represented by the Donggou porphyry Mo deposit
and several Pb-Zn-Ag vein deposits (Fig. 2). The Donggou Mo deposit was discovered in 1984 by the

No. 2 Team of Henan Bureau of Geology and Mineral Resources during a regional geological survey,
and has proven reserves of 0.63 Mt Mo at an average grade of 0.11 wt % (Ye et al., 2006). Spatially
associated with the Donggou deposit are several Pb-Zn-Ag vein deposits hosted by E- or NE-striking
faults, as best illustrated by the Sanyuangou, Wangpingxigou, and Laodaizhanggou deposits (Fig. 2).
The combined reserves of these deposits is 0.93 Mt Pb + Zn (Ye, 2006; Yao et al., 2008), whereas the
quantity of the Ag reserve is not published.

#### 198 Donggou porphyry Mo deposit

199 The Donggou porphyry Mo deposit (112°22'50" E, 33°57'03" N) is related to the Donggou porphyry, which intruded andesite and basaltic andesite of the Xiong'er Group. Unlike typical 200 201 porphyry Cu-Mo deposits where the ores are commonly localized in porphyritic intrusions, Mo 202 mineralization at Donggou is mostly hosted in the andesite of the Xiong'er Group up to 360 m from the 203 intrusive contact of the Donggou porphyry (Figs. 4, 5a, 5c). The ores in the volcanic rocks account for 204 ca. 98 % of the total reserves of the deposit (Ma et al., 2007). Minor Mo mineralization occur in the 205 uppermost zone (<70 m) of the Donggou porphyry (Figs. 4, 5b, 6a). The deposit consists of 19 206 orebodies, individually 47 to 254 m thick. Molybdenum mineralization in the Paleoproterozoic 207 volcanic rocks is largely controlled by abundant fractures in the rocks that provided high permeability 208 for magmatic-derived, Mo-bearing hydrothermal fluids (Yang et al., 2011, 2015).

209 Molybdenum mineralization is dominated by stockwork veinlets and veins bearing molybdenite, 210 with minor amounts of sulfide disseminations (pyrite, chalcopyrite, sphalerite, and galena; Figs. 5, 6b-c) 211 both within the Donggou porphyry and the proximal Paleoproterozoic volcanic rocks. They are largely 212 represented by fine-grained molybdenite disseminations (Fig. 5a), coarse-grained molybdenite 213 aggregates (Fig. 5b), and K-feldspar - molybdenite veins (Fig. 5c-d) that are commonly overprinted by 214 quartz – molybdenite veins (Fig. 5d). Polymetallic sulfide veins occur locally in the proximal volcanic 215 rocks. These veins are typically 10 to 50 cm wide and cut hydrothermally altered and Mo-mineralized 216 andesite (Fig. 5e-f). Pyrite, sphalerite, and galena are abundant in these veins, with minor molybdenite 217 and chalcopyrite (Figs. 5e-f, 6d-f).

Hydrothermal alteration is well developed both in the Donggou porphyry and the proximal andesite. The dominated alteration types are potassic (Fig. 5b-d) and silicic alteration (Fig. 5d-f).

220 Potassic alteration is largely represented by K-feldspar, locally associated with minor biotite, and 221 typically occurs as mm-to-cm-scale aggregates of K-feldspar (Figs. 5a-d). Silicic alteration occurs 222 mainly in the wall rocks surrounding the porphyry, shown by quartz with various amounts of sulfide 223 minerals (Figs. 5d-f). Molybdenite is present both in the potassic and silicic assemblages, occurring 224 mainly as aggregates (Fig. 5b-d). Less extensive alteration formed sericite, chlorite, and carbonate 225 aggregates or irregular veinlets. Based on field and petrographic observations, four paragenetic stages 226 are recognized at Donggou (Fig. 7): stage D1 (D is short for Donggou) K-feldspar - molybdenite veins; 227 stage D2 quartz – molybdenite veins; stage D3 quartz – polymetallic sulfide veins; and stage D4 quartz 228 - calcite veins. Molybdenite mainly occurs during stages D1 and D2.

#### 229 Sanyuangou Pb-Zn-Ag deposit

230 In the whole Fudian ore field, Pb-Zn-Ag vein deposits share common geological and 231 mineralization features. To understand their temporal and spatial relationships with the Donggou 232 porphyry, here we focus on the Sanyuangou Pb-Zn-Ag vein deposit. The Sanyuangou deposit 233 (112°22'19" E, 33°55'55" N) is located on the southern edge of the Fudian ore field, about 3 km to the 234 south of the Donggou porphyry Mo deposit (Fig. 2). Mineralization is hosted in a quartz diorite stock 235 elongated easterly that has been cut by numerous NW-, NE-, and E-oriented faults (Fig. 2). The Pb-Zn-236 Ag mineralization is largely represented by Pb-Zn-sulfide veins, which mainly occur in E-striking 237 faults (Figs. 2, 8a-b). These vein-hosting faults commonly dip 70°-85° to the north, but locally steeply 238 dip to the south. Individual veins are usually 300 to 2,000 m long, 0.5 to 5 m wide, and continuous for 239 several hundred meters down plunge. Quartz - pyrite veins are typically crosscut by Pb-Zn-sulfide 240 veins (e.g., Fig. 8a). Individual Pb-Zn-ore shoots are separated by less mineralized, subeconomic 241 segments along the veins (Fig. 8a-d). Massive sulfide ores (Fig. 8c-e) are common, and preferentially 242 localized in dilational jogs, splays, and bifurcations along the structures. Sulfide minerals in the Pb-Zn-243 Ag veins are dominated by pyrite, sphalerite, and galena, with minor amounts of chalcopyrite, 244 pyrrhotite, bornite, tetrahedrite, and covellite (Figs. 8a-e, 9a-d). Galena is the main silver-bearing 245 mineral, but minor polybasite and argentite are also present.

Hydrothermal alteration is well developed and ranges from cm-wide selvages to meter-wide halos
(e.g., Fig. 8a). The alteration assemblages consist mainly of quartz, sericite, siderite, ankerite, and

- 248 calcite, with minor chlorite (Figs. 8a, f, 9e-f). Field and textural relationships indicate three paragenetic
- 249 stages (Fig. 10): stage S1 (S is short for Sanyuangou) quartz pyrite veins; stage S2 quartz -
- polymetallic sulfide veins; and stage S3 quartz calcite veins. Pb-Zn-Ag mineralization is mainly
  associated with stage S2.

#### 252 Samples and Analytical Methods

#### 253 Geochronology

#### 254 Zircon U-Pb dating

255 The Donggou granite porphyry (DG01) hosting the Donggou porphyry Mo deposit and the quartz 256 diorite (SY01) and quartz monzonite (WP01) hosting the Sanyuangou and Wangpingxigou Pb-Zn-Ag 257 vein deposits, respectively, were collected for zircon U-Pb geochronology (Figs. 2, 3). U-Pb dating of 258 guartz diorite and guartz monzonite was aimed to test previous speculations that these host rocks are 259 temporarily and genetically related to the Pb-Zn-Ag mineralization. Zircon grains were handpicked 260 under a binocular microscope after conventional crushing, liquid and magnetic separation. 261 Representative grains were mounted in epoxy resin disks, and then polished, cleaned, and gold coated. 262 Prior to isotopic analysis, all grains were photographed under transmitted- and reflected-light 263 microscope and then imaged using the cathodoluminescence (CL) technique with a JEOL 8800 264 electron microprobe.

265 Zircon U-Pb dating was conducted using laser ablation inductively coupled plasma mass 266 spectroscopy (LA-ICP-MS) at the State Key Laboratory of Geological Processes and Mineral 267 Resources (GPMR), China University of Geosciences, Wuhan. Laser analysis was performed with a 268 GeoLas 2005. An Agilent 7500a ICP-MS instrument was used to acquire ion-signal intensities. Helium 269 was applied as a carrier gas. Argon was used as the make-up gas and mixed with the carrier gas via a 270 T-connector before entering the ICP. Nitrogen was added into the central gas flow (Ar + He) of the Ar 271 plasma to decrease the detection limit and improve precision (Hu et al., 2008). Each analysis 272 incorporated a background acquisition of approximately 20 to 30 s (gas blank) followed by 50 s data 273 acquisition from the sample. The Agilent Chemstation was utilized for the acquisition of each 274 individual analysis. Off-line selection and integration of background and analyte signals, and time-drift 275 correction and quantitative calibration for trace element analyses and U-Pb dating were performed by

*ICPMSDataCal* (Liu et al., 2008, 2010). Zircon 91500 was used as the external standard for U-Pb dating, and was analyzed twice every five sample analyses. Time-dependent drifts of U-Th-Pb isotopic ratios were corrected using a linear interpolation (with time) for every five analyses according to the variations of 91500 (Liu et al., 2010). Uncertainty of preferred values for the external standard 91500 was propagated to the ultimate results of the samples. Concordia diagrams and weighted mean calculations were made using Isoplot/Ex\_ver3 (Ludwig, 2003).

#### 282 Molybdenite Re-Os dating

283 Molybdenite samples representative of stage D2 quartz – molybdenite vein (DG03; Fig. 5d) and 284 stage D3 quartz - polymetallic sulfide vein (FDG08; Fig. 5f) of the Donggou porphyry Mo deposit 285 were collected to establish the timing of molybdenite mineralization using the Re-Os chronometer. The 286 Carius tube method was used for dissolution of molybdenite and equilibration of the sample with tracer 287 Re and Os (Selby and Creaser, 2001). Approximately 22-23 mg of molybdenite were dissolved and 288 equilibrated with a known amount of <sup>185</sup>Re and isotopically normal Os at 240 °C for 24 h. Solvent 289 extraction and microdistillation was used to separate Os, whereas anion exchange chromatography was 290 used to separate Re (Selby and Creaser, 2001). The concentrations of <sup>187</sup>Re and <sup>187</sup>Os were determined 291 at the Laboratory for Sulfide and Source Rock Geochronology and Geochemistry (a member of the 292 Durham Geochemistry Centre), Durham University, using isotope dilution negative thermal ionization 293 mass spectrometry (ID-NTIMS). Isolated and purified Re and Os solutions were loaded onto Ni and Pt 294 filaments, respectively, and analyzed using a Thermo Scientific TRITON mass-spectrometer with 295 Faraday collectors. Re-Os model ages were calculated by the equation,  $\ln ({}^{187}\text{Os}/{}^{187}\text{Re} + 1)/\lambda$ , where  $\lambda$ 296 denotes the decay constant for  $^{187}$ Re. The  $^{187}$ Re decay constant used is  $1.666 \times 10^{-11}$  year<sup>-1</sup>, with an 297 uncertainty of 0.31% (Smoliar et al., 1996; Selby et al., 2007). Uncertainties in the age calculations 298 include uncertainties associated with (1) <sup>185</sup>Re and <sup>190</sup>Os spike calibrations, (2) weighing the spikes, (3) 299 magnification with spiking, (4) mass spectrometric measurement of isotopic ratios, (5) blanks (Re = 2.4300 pg, Os = 0.1 pg,  ${}^{187}Os/{}^{188}Os = \sim 0.25$ ), and (6) the  ${}^{187}Re$  decay constant.

#### 301 Sericite <sup>40</sup>Ar/<sup>39</sup>Ar dating

302 Sample SY02 was taken from sericitic alteration assemblages associated with a stage S2 Pb-Zn-303 sulfide vein of Sanyuangou. The sample is dominated by sericite that is intergrown with sphalerite and 304 pyrite (Fig. 9f). After petrographic examination, suitable parts of the sample were crushed, repeatedly 305 sieved to obtain mineral grains as uniform as possible in size (0.5-2 mm), washed in distilled water in

an ultrasonic bath for 1 h, and dried. Sericite aggregates of 0.5 to 1.0 mm were screened under abinocular microscope.

308 Sericite mineral separates were irradiated along with the ZBH-25 biotite standard ( $132.7 \pm 1.2$  Ma 309 at  $1\sigma$ ; Wang, 1983) for 55 h in the Swimming Pool Reactor, Chinese Institute of Atomic Energy 310 (Beijing). After a three month cooling period, the samples were analyzed by the  ${}^{40}$ Ar/ ${}^{39}$ Ar stepwise 311 incremental heating method using a MM-1200B mass spectrometer at Institute of Geology, Chinese 312 Academy of Geological Sciences. The analytical procedures are detailed in Chen et al. (2002) and 313 summarized here. The Ar extraction system comprises an electron bombardment heated furnace in 314 which the samples were heated under vacuum. The released gases were admitted to a purification 315 system, with 30 min for heating-extraction for each temperature increment and 30 min for purification. 316 Purified Ar was trapped in activated charcoal finger at liquid-nitrogen temperature and then released 317 into the mass spectrometer for isotope analysis. Measured isotopic ratios were corrected for mass 318 discrimination, atmospheric Ar component, blanks, and irradiation-induced mass interference. The 319 correction factors of interfering isotopes produced during irradiation were determined by analysis of irradiated K<sub>2</sub>SO<sub>4</sub> and CaF<sub>4</sub> pure salts, and their values are  $({}^{36}\text{Ar}/{}^{37}\text{Ar})_{Ca} = 0.000240$ ,  $({}^{40}\text{Ar}/{}^{39}\text{Ar})_{K} =$ 320 0.004782, and  $({}^{39}Ar/{}^{37}Ar)_{Ca} = 0.000806$  (Chen et al., 2002). All dates are reported using  $5.543 \times 10^{-10}$  a 321 322 <sup>-1</sup> as the total decay constant for <sup>40</sup>K (Steiger and Jäger, 1977). The age uncertainties are reported at the 323 95% confidence level ( $2\sigma$ ), and include the uncertainties in irradiation correction factors and the J value, 324 but do not include the uncertainty in the potassium decay constant. The Ar-Ar data were calculated and 325 plotted using Isoplot/Ex\_ver3 (Ludwig, 2003).

#### 326 LA-ICP-MS analyses of pyrite

Two ore samples (FDG09 and FDG11) from stage D3 Pb-Zn-sulfide vein of the Donggou porphyry Mo deposit and two samples (SY03 and SY04) from stage S2 Pb-Zn-sulfide vein of the Sanyuangou Pb-Zn-Ag deposit were selected for pyrite trace element analysis. The analysis was conducted at the Centre of Excellence in Ore Deposits (CODES), University of Tasmania, Australia. Analytical instrumentation consists of a New Wave 213-nm solid-state laser microprobe and an Agilent 7500 Quadrupole ICP-MS. The operating conditions and procedures applied to pyrite have been detailed in Large et al. (2007) and Danyushevsky et al. (2011). 334 Spot analyses of pyrite were performed by laser-ablating spots of 35 um diameter. The repetition 335 rate was 5 Hz, and laser beam energy was maintained between 4 and 5 Jcm<sup>-2</sup>. Analysis time was 336 restricted to 90 s, consisting of 30-s background (laser off) and 60-s analysis (laser on). The primary 337 calibration standard (STDGL2b2) consists of a fused glass containing 25% Zn concentrate and 75% 338 pyrrhotite developed in-house (Danyushevsky et al., 2011). It was analyzed twice every 1.5 hours with 339 a 100-µm beam size at 5 Hz to correct for instrument drift. Data were reduced using SILLS software 340 following standard methods (Longerich et al., 1996). Iron was used as the internal standard, and the Fe 341 contents were determined by EMP analyses at the State Key Laboratory of GPMR, China University of 342 Geosciences, Wuhan.

#### 343 In-situ lead isotope analyses

344 Three ore samples (FDG09, FDG10, and FDG11) from stage D3 quartz - polymetallic sulfide 345 veins at Donggou and three samples (SY04, SY05, and SY06) from stage S2 Pb-Zn-sulfide veins at 346 Sanyuangou were selected for lead isotopic analyses to provide additional constraints on a possible 347 genetic link between these two deposits. In-situ lead isotope analyses on pyrite were performed on a 348 Neptune Plus MC-ICP-MS (Thermo Fisher Scientific, Dreieich, Germany) equipped with a Geolas 349 2005 excimer ArF laser ablation system (Lambda Physik, Göttingen, Germany) at the State Key 350 Laboratory of GPMR, China University of Geosciences, Wuhan. In the laser ablation system, helium 351 was used as the carrier gas for the ablation cell and was mixed with argon (make-up gas) after the 352 ablation cell. The spot diameter ranged from 44 to 90 µm dependent on Pb signal intensity. The pulse 353 frequency was from 4 to 10 Hz, but the laser fluence was kept constant at  $\sim 3$  J/cm<sup>2</sup>. A new signal-354 smoothing and mercury-removing device was used downstream from the sample cell to efficiently 355 eliminate the short term variation of the signal and remove the mercury from the background and 356 sample aerosol particles (Hu et al., 2014). The Neptune Plus was equipped with nine Faraday cups fitted with 10<sup>11</sup>  $\Omega$  resistors. Isotopes <sup>208</sup>Pb, <sup>207</sup>Pb, <sup>206</sup>Pb, <sup>206</sup>Pb, <sup>204</sup>Pb, <sup>205</sup>Tl, <sup>203</sup>Tl and <sup>202</sup>Hg were collected in 357 358 faraday cups within static mode. The mass discrimination factor for Pb was determined using a Tl 359 solution (NIST SRM 997) nebulized at the same time as the sample, using an Aridus II desolvating 360 nebulizer. The 202Hg signal was used to correct the remained 204Hg interference on 204Pb, using the 361 natural <sup>202</sup>Hg/<sup>204</sup>Hg ratio.

362 In this method, the natural Tl-isotopic composition was assumed. MASS-1, a sulfide standard 363 reference from USGS (Wilson et al., 2002), and two in-house standards PY-3 and Sph-2 were then run 364 to define the mathematical relationship between Tl and Pb mass bias (Woodhead, 2002). The Pb 365 isotopic compositions of those three sulfides samples were previously determined by solution MC-ICPMS at GPMR. MASS-1 was used to monitor the precision and accuracy of the measurements after 366 367 ten sample analysis, over the entire period of analysis. The obtained accuracy is estimated to be equal to or better than  $\pm 0.6$  % for <sup>208</sup>Pb/<sup>204</sup>Pb, <sup>207</sup>Pb/<sup>204</sup>Pb, and <sup>206</sup>Pb/<sup>204</sup>Pb compared to the solution value by 368 369 MC-ICP-MS, with a typical precision of  $0.4 \% (2\sigma)$ .

370 **Results** 

#### 371 Geochronological data

#### 372 Zircon U-Pb ages

373 The U-Pb isotope data are tabulated in Appendix 1 and illustrated in concordia diagrams (Fig. 374 11). Zircon grains from the Donggou granite porphyry are euhedral to subhedral, 30-200 µm long, with 375 aspect ratios of 1 to 2.5. Most zircon grains show oscillatory zoning in CL images (Fig. 11a), consistent 376 with a magmatic origin. These zircons have high Th (177-1911 ppm) and U (279-3722 ppm), with 377 Th/U ratios ranging from 0.43 to 1.71, which is typical of magmatic zircons (Claesson et al., 2000). A 378 total of thirteen spot analyses were made on twelve zircon grains for sample DG01. Eleven spots are 379 concordant, with the remaining two being slightly discordant (Fig. 11a). However, the discordant data 380 have <sup>206</sup>Pb/<sup>238</sup>Pb ages indistinguishable from the concordant ones (Appendix 1), indicating that the 381 discordance may reflect uncertainties related to <sup>207</sup>Pb measurement or common Pb correction, rather 382 than lead loss. All thirteen analyses have a weighted mean  $^{206}Pb/^{238}U$  age of 117.8  $\pm$  0.9 Ma (1 $\sigma$ , 383 MSWD = 0.1; Fig. 11a).

Zircon grains from the Sanyuangou quartz diorite are euhedral to subhedral, 50-200  $\mu$ m long, with aspect ratios of 1 to 2. In CL images, most zircon grains show oscillatory zoning (Fig. 11b). These grains have relatively high Th (358-2,403 ppm) and U (393-1,350 ppm), with Th/U ratios of 0.69 to 1.79 (Appendix 1). Fourteen spot analyses were made on 14 zircon grains for sample SY01. The spots are concordant and yield a weighted mean <sup>207</sup>Pb/<sup>206</sup>Pb age of 1,756 ± 9 Ma (1 $\sigma$ , MSWD = 1.1; Fig. 11b). Zircon grains from the Wangpingxigou quartz monzonite are euhedral and display welldeveloped prismatic and pyramidal faces. They are commonly 100-300  $\mu$ m long with length/width ratios of 1 to 3. Most zircon grains show spectacular oscillatory zoning in CL images (Fig. 11c). These grains contain 167-993 ppm Th and 214-885 ppm U, with Th/U ratios of 0.65 to 1.48 (Appendix 1). Fifteen spot analyses on 14 zircon grains from sample WP01 yield concordant ages with a weighted mean <sup>207</sup>Pb/<sup>206</sup>Pb age of 1,837 ± 11 Ma (1 $\sigma$ , MSWD = 1.1; Fig. 11c).

#### 396 Molybdenite Re-Os ages

Re-Os isotope data are presented in Table 1. All age uncertainties are quoted at the  $2\sigma$  level. Sample from the quartz – molybdenite vein (DG03) contains 2.6 ppm Re and 3.2 ppb <sup>187</sup>Os, and has a Re-Os model age of 117.5 ± 0.8 [0.9] Ma (bracketed value refers to errors including the decay constant uncertainty). Sample from the quartz – polymetallic sulfide vein (FDG08) contains 7.1 ppm Re and 8.6 ppb <sup>187</sup>Os, and has a Re-Os model age of 116.4 ± 0.6 [0.7] Ma. The age difference between sample DG03 and FDG08 is consistent with field relationship and paragenetic sequence between the stage D2 quartz – molybdenite vein and stage D3 quartz – polymetallic sulfide vein.

#### 404 Sericite <sup>40</sup>Ar/<sup>39</sup>Ar age

405 The <sup>40</sup>Ar/<sup>39</sup>Ar results of sample SY02 are summarized in Table 2 and the age spectrum and 406 inverse isochron are shown in Figure 12. All <sup>40</sup>Ar/<sup>39</sup>Ar ages are calculated using the decay constants of 407 Steiger and Jäger (1977) and the uncertainties are reported at the 95 % confidence level  $(2\sigma)$ . The 408 sample yields a well-defined plateau age of  $115.9 \pm 0.9$  Ma (MSWD = 0.7), consisting of seven 409 contiguous steps that account for 92.7 % of the total <sup>39</sup>Ar released (Fig. 12a). Isotopic results of these 410 steps formed a well-defined inverse isochron in the <sup>39</sup>Ar/<sup>40</sup>Ar versus <sup>36</sup>Ar/<sup>40</sup>Ar diagram (Fig. 12b), with 411 an inverse isochron age of  $116.0 \pm 1.2$  Ma (MSWD = 10.9) that is indistinguishable from the plateau 412 age. The inverse isochron plot has a  ${}^{40}$ Ar/ ${}^{36}$ Ar ratio of 299.0 ± 6.0, which is within uncertainty of the 413 accepted value of the present air (298.5  $\pm$  0.3; Renne et al., 2009) and thus suggests that no excessive 414 argon was incorporated into the mineral.

#### 415 Trace element compositions of pyrite

416 A total of 50 LA-ICP-MS spot analyses were made on pyrite grains from the Donggou porphyry
417 Mo and Sanyuangou Pb-Zn-Ag vein deposits. The trace element analyzed include Cr, Mn, Fe, Co, Ni,

418 Cu, Zn, Ga, As, Se, Mo, Ag, Cd, Sn, Sb, Te, Ba, Gd, Hf, Ta, W, Pt, Au, Tl, Pb and Bi. The full 419 analytical results are listed in Appendix 2, and the compositions of selected trace elements are 420 illustrated in Figure 13. Pyrite from the Donggou porphyry Mo deposit is depleted in most trace 421 elements, but contains higher Mo (up to 154 ppm) and Bi (29 ppm) compared to pyrite from the 422 Sanyuangou Pb-Zn-Ag vein deposit. Pyrite from Donggou also contains considerable Sn (0.10 to 6 423 ppm), Te (0.07 to 9.7 ppm), Pb (0.07 to 398 ppm), As (1.76 to 1821 ppm), and Sb (0.01 to 31 ppm). In 424 contrast, pyrite from Sanyuangou has elevated Cu (2.7 to 23,420 ppm), Zn (0.25 to 26,893 ppm), Pb 425 (0.02 to 637 ppm), Ag (up to 1091 ppm), Au (up to 21 ppm), and As (0.3 to 17913 ppm). Pyrites from 426 Donggou and Sanyuangou have similar Co (0.1-1224 vs. 0.5-1290 ppm) and Ni (0.1-116 vs. 0.3-300 427 ppm) contents and Co/Ni ratios (0.7-48 vs. 0.4-82; Fig. 14).

#### 428 Pyrite lead isotopic compositions

The *in-situ* lead isotope analyses of pyrite are listed in Table 3. Ten pyrite grains from Donggou have  ${}^{206}Pb/{}^{204}Pb$  ratios of 17.328 to 17.517,  ${}^{207}Pb/{}^{204}Pb$  of 15.408 to 15.551, and  ${}^{208}Pb/{}^{204}Pb$  of 38.080 to 38.436. Pyrite grains from Sanyuangou have Pb isotopic compositions that are very similar to that of Donggou ( ${}^{206}Pb/{}^{204}Pb$  = 17.273 to 17.495;  ${}^{207}Pb/{}^{204}Pb$  = 15.431 to 15.566;  ${}^{208}Pb/{}^{204}Pb$  = 37.991 to 38.337; n = 10).

- 434 Discussion
- 435 **Timing of mineralization**

#### 436 Donggou porphyry Mo deposit

437 Zircon grains from the Donggou granite porphyry show morphological and compositional features 438 typical of magmatic zircons (Claesson et al., 2000; Wu and Zheng, 2004; Appendix 1). The precise 439 LA-ICP-MS zircon U-Pb age (117.8  $\pm$  0.9 Ma) presented here provides a tight constraint on the 440 emplacement age of the Donggou porphyry. This age is consistent, within analytical uncertainty, with a 441 previous LA-ICP-MS zircon U-Pb age  $(117 \pm 1 \text{ Ma})$  for the Donggou porphyry (Dai et al., 2009). The 442 younger zircon U-Pb ages ( $112 \pm 1$  Ma by SHRIMP, Ye et al., 2006;  $114 \pm 1$  Ma by LA-ICP-MS, Dai 443 et al., 2009) either reflect multistage magmatism at Donggou or different analytical protocols used by 444 those authors. However, our field observations and drill core logging did not identify evidence for 445 multiple intrusions, nor did the authors who obtained the younger U-Pb ages presented evidence that 446 their samples were from intrusions other than Donggou porphyry. Thus it is suggested that the different 447 ages have more likely resulted from different analytical methods or from the analysis of zircon grains 448 that have experienced Pb loss.

449 The molybdenite Re-Os chronometer is remarkably robust due to the high closure temperature of 450 Re-Os isotopes in molybdenite (Suzuki et al., 1996; Stein et al., 2001; Selby et al., 2002; Chiaradia et 451 al., 2013). Thus, Re-Os dating of this mineral commonly provides reliable constraints on the timing of 452 ore formation (Selby et al., 2002). Molybdenite from the stages D2 and D3 veins of the Donggou Mo 453 deposit have overlapping Re-Os model ages of  $117.5 \pm 0.8$  Ma and  $116.4 \pm 0.6$  Ma (Table 1). The 454 molybdenite Re-Os ages are in excellent agreement with zircon U-Pb age of the Mo mineralized 455 porphyry, providing strong evidence for temporal and genetic relationship between magmatism and Mo 456 mineralization. The absolute value of molybdenite Re-Os age of the stage D3 quartz - polymetallic 457 sulfide vein (116.4  $\pm$  0.6 Ma) is slightly younger than that of the stage D2 quartz – molybdenite vein 458  $(117.5 \pm 0.8 \text{ Ma})$  and zircon U-Pb age  $(117.0 \pm 1 \text{ Ma})$  of the Donggou porphyry, but they are all 459 indistinguishable if the analytical uncertainties are considered. Based on these results, we propose that 460 the Donggou porphyry Mo deposit formed between  $117.5 \pm 0.8$  Ma and  $116.4 \pm 0.6$  Ma during the 461 Early Cretaceous, genetically associated with the Donggou porphyry.

#### 462 Sanyuangou Pb-Zn-Ag vein deposit

463 Field and petrographic relations (Fig. 9f) indicate that sericite is spatially and texturally associated 464 with stage S2 Pb-Zn sulfide veins at Sanyuangou, and thus <sup>40</sup>Ar/<sup>39</sup>Ar dating of sericite can provide 465 constraints on the timing of hydrothermal alteration and mineralization (Li et al., 2003; Chiaradia et al., 466 2013). One sericite sample has a  ${}^{40}$ Ar/ ${}^{39}$ Ar plateau age of 115.9 ± 0.9 Ma and an identical inverse 467 isochron age of  $116.0 \pm 1.2$  Ma (Fig. 12), which represent the time when sericite cooled below the 468 closure temperature of argon isotopes in mica. Formation of Pb-Zn-Ag veins at Sanyuangou has been 469 constrained at 206°-265°C based on the sulfur isotopic equilibration equations of pyrite-galena and 470 sphalerite-galena pairs from stage S2 veins (Li, 2013). This temperature range is lower than the argon 471 closure temperature in mica (300°-350°C; McDougall and Harrison, 1999), the present <sup>40</sup>Ar/<sup>39</sup>Ar age 472 can therefore be reliably interpreted as the timing of hydrothermal alteration and Pb-Zn-Ag 473 mineralization of the Sanyuangou deposit. Ye et al. (2006) reported a sericite <sup>40</sup>Ar/<sup>39</sup>Ar "plateau" age 474 of 110.1  $\pm$  9.2 Ma (24.2 % of the total <sup>39</sup>Ar released) for Sanyuangou, which overlaps with our new <sup>40</sup>Ar/<sup>39</sup>Ar data but has a significantly larger uncertainty. Our new <sup>40</sup>Ar/<sup>39</sup>Ar age therefore provides an
improved and tight constraint for the timing of mineralization at Sanyuangou.

477 Zircon grains from the Sanyuangou quartz diorite stock hosting the Pb-Zn-Ag mineralization have 478 a weighted mean  ${}^{207}\text{Pb}/{}^{206}\text{Pb}$  age of 1,756  $\pm$  9 Ma (Fig. 11b), which is considered to be the 479 emplacement age of this intrusion. To the east of the Sanyuangou quartz diorite, the Wangpingxigou quartz monzonite has a  ${}^{207}\text{Pb}/{}^{206}\text{Pb}$  age of 1,837 ± 11 Ma (Fig. 11c), which predates the Sanyuangou 480 quartz diorite stock by 80-100 m.y. Thus, these two intrusions are products of two episodes of granitoid 481 482 magmatism during the Late Paleoproterozoic. The present LA-ICP-MS zircon U-Pb and sericite 483 <sup>40</sup>Ar/<sup>39</sup>Ar dating results demonstrate that the Sanyuangou Pb-Zn-Ag deposits formed during the Early 484 Cretaceous and significantly postdate the host quartz diorite and quartz monzonite. Consequently, a 485 possible genetic link between the host intrusion and mineralization, as previously suggested (HBGMR, 486 1989), can be ruled out.

#### 487 A porphyry-related Mo-Pb-Zn-Ag metallogenic system

The Sanyuangou Pb-Zn-Ag vein deposit is located about 3 km south of the Donggou porphyry Mo deposit (Fig. 2). Sericite  ${}^{40}$ Ar/ ${}^{39}$ Ar dating of the alteration assemblages proximal to a Stage S2 Pb-Zn-sulfide vein (115.9 ± 0.9 Ma) at Sanyuangou and Re-Os dating of molybdenite from Stage D2 quartz – molybdenite vein (117.5 ± 0.8 Ma) and Stage D3 quartz – polymetallic sulfide vein (116.4 ± 0.6 Ma) at Donggou suggest that Pb-Zn-Ag mineralization at Sanyuangou was mutually consistent. Therefore, the close spatial-temporal relations between the Sanyuangou Pb-Zn-Ag and Donggou porphyry Mo deposits propose a genetic association.

495 Pyrite is an important scavenger of a variety of trace elements as Au, Ag, Cu, Pb, Zn, Co, Ni, As, 496 Sb, Se, Te, Hg, Tl, and Bi (Cook and Chryssoulis, 1990; Craig et al., 1998; Large et al., 2007; Reich et 497 al., 2013) and thereby can provide useful information on the composition and evolution of ore fluids 498 (Hawley and Nichol, 1961; Bralia et al., 1979; Craig et al., 1998; Morey et al., 2008; Large et al., 2007, 499 2009, 2011). Knowledge on the trace-element geochemistry of pyrite not only contributes to a better 500 understanding in the physicochemical parameters of ore fluids from which pyrite precipitated and thus 501 a more geologically reasonable metallogenic model for the pyrite-dominated deposits, but also 502 improves exploration strategies of these deposits (Heinrich et al., 2003; Danyushevsky et al., 2011).

503 Laser ablation ICP-MS spot analyses show that pyrite from Donggou is enriched in Mo and Bi (Fig. 504 13). In contrast, pyrite from Sanyuangou is depleted in Mo and Bi, but enriched in Cu, Zn, Pb, Ag, Au, 505 and As (Fig. 13). The trace element patterns of pyrite from these two deposits are consistent with metal 506 zonation typically observed in a porphyry-related hydrothermal system, commonly illustrated by Cu, 507 Mo  $\rightarrow$  Zn, Pb  $\rightarrow$  Ag, Au, As from porphyry ores to distal base-metal veins (Emmons, 1927; Jones, 508 1992; Seedorff et al., 2005; Sillitoe, 2010). On the other hand, pyrite from Sanyuangou has contents of 509 Sn, Sb, Co, and Ni similar to those from Donggou, with comparable Co/Ni ratios mostly higher than 1 510 (Figs. 13, 14). This observation also is supportive for a magmatic-hydrothermal origin for the 511 Sanyuangou Pb-Zn-Ag deposit (Bralia et al., 1979; Deol et al., 2012).

512 Pyrite may contain moderate to high quantities of Pb, but virtually lacks U, and therefore its Pb 513 isotopic compositions provide valuable information on the sources of lead and associated metals 514 (Meffre et al., 2008; Woodhead et al., 2009; Darling et al., 2012; Steadman et al., 2013). In-situ pyrite 515 isotopic compositions of Donggou and Sanyuangou are used to uncover a possible genetic link between 516 these two deposits. Pyrite from Sanyuangou has Pb isotopic compositions (<sup>206</sup>Pb/<sup>204</sup>Pb = 17.273-17.495,  $^{207}Pb/^{204}Pb = 15.431-15.566$ , and  $^{208}Pb/^{204}Pb = 37.991-38.337$ ) that are similar to values of pyrite from 517 518 Donggou (<sup>206</sup>Pb/<sup>204</sup>Pb = 17.328-17.517, <sup>207</sup>Pb/<sup>204</sup>Pb = 15.408-15.551, and <sup>208</sup>Pb/<sup>204</sup>Pb = 38.080-38.436). 519 Thorogenic (Fig. 15a) and uranogenic (Fig. 15b) Pb isotopic diagrams show that the pyrite samples 520 from the two deposits plot in the same area and display similar trends, indicating a common source of 521 the lead. Collectively, field, paragenetic, geochronological, geochemical, and isotopic data indicate that 522 the Pb-Zn-Ag mineralization at Sanyuangou is genetically related to the Donggou porphyry Mo system, 523 and can be best interpreted as a distal product of the Donggou magmatic-hydrothermal system.

524 There are additional Pb-Zn-Ag vein deposits surrounding the Donggou porphyry Mo deposit 525 (e.g., Wangpingxigou, Laodaizhanggou; Fig. 2). These deposits have geological and mineralization 526 characteristics consistent with the Sanyuangou deposit (Huang et al., 1992; Ye, 2006; Yao et al., 2008). 527 The Wangpingxigou deposit has a sphalerite Rb-Sr isochron age of  $117 \pm 27$  Ma (Yao et al., 2010). 528 This age, although having a large analytical uncertainty, indicates that Wangpingxigou formed in Early 529 Cretaceous, broadly synchronous with the Pb-Zn-Ag veins at Sanyuangou. However, ore-related 530 sericite from the Laodaizhanggou Pb-Zn-Ag vein deposit yields a well-defined <sup>40</sup>Ar/<sup>39</sup>Ar plateau age of 531 124.7  $\pm$  1.2 Ma (Li, 2013), which is about 7 m. y. older than Sanyuangou (115.9  $\pm$  0.9 Ma; Fig. 12a) 532 and Donggou ( $117.5 \pm 0.8$  Ma to  $116.4 \pm 0.6$  Ma; Table 1). As such, the age indicate that district-wide

533 Pb-Zn-Ag mineralization may have formed in association with multiple magmatic-hydrothermal534 activities.

535 Early Cretaceous magmatic activity is well-recognized at Waifangshan and surrounding districts. 536 The Taishanmiao pluton, with an exposure of ca. 290 km<sup>2</sup> in southwest of the Fudian ore field (Figs. 1, 537 2), formed by episodic magmatism from 125 to 115 Ma as revealed by zircon U-Pb dating (Ye et al., 538 2008; Qi, 2014; Gao et al., 2014). In the east of Taishanmiao, the Zhuyuangou quartz-molybdenite 539 veins (0.1 Mt Mo; Huang et al., 2010) are hosted in fine-grained granite and have molybdenite Re-Os 540 ages of  $122.2 \pm 2.3$  and  $119.6 \pm 2.2$  Ma (Huang et al., 2010). Taken together, we suggest that the Pb-541 Zn-Ag vein deposits in the Fudian ore field could have been products of multiple magmatic-542 hydrothermal events in the Early Cretaceous, with some (e.g., Sanyuangou and Wangpingxigou) being 543 related to the Donggou mineralized porphyry, but others (e.g., Laodaizhanggou) likely having an 544 association with earlier magmatism as represented by the Taishanmiao intrusive complex and 545 Zhuyuangou quartz-molybdenite veins (Huang et al., 2010).

#### 546 Implications for mineral exploration

547 Our study has shown that the Pb-Zn-Ag veins in the Fudian ore field are most likely distal 548 products of porphyry-related magmatic-hydrothermal systems. This view has significant implications 549 for Mo and Pb-Zn-Ag mineral exploration in the area, as these two mineralization styles can be used as 550 a vector for each other (Sillitoe, 2010). The Pb-Zn-Ag veins commonly occur in shallow and distal 551 parts of a porphyry system. The fault zones surrounding known porphyry Mo deposits are favorable 552 sites for localizing Pb-Zn-Ag veins (Seedorff et al., 2005; Sillitoe, 2010). Brittle faults with different 553 strikes are well developed in the Fudian ore field (Fig. 2), some of which are marked by variable 554 degrees of hydrothermal alteration. Such structures are good targets for Pb-Zn-Ag veins. Other districts 555 along the southern NCC also host giant porphyry and porphyry-skarn Mo deposits that formed during 556 the Late Jurassic to Early Cretaceous (Fig. 1), including Jinduicheng (Huang et al., 1994), Nannihu (Li 557 et al., 2003), Leimengou (Li et al., 2006), and Yuchiling (Li et al., 2012d, 2013a). Our model in the 558 Fudian ore field has broad implications for exploration of Pb-Zn-Ag deposits in those districts, where 559 fractures zones, particularly those with extensive hydrothermal alteration, should be good targets for 560 Pb-Zn-Ag resources.

561 The Pb-Zn-Ag veins are widespread in large areas of the southern NCC (Fig. 1). Although some 562 veins show close spatial and temporal relationships with porphyry Mo deposits (Fig. 1), such a 563 relationship, however, is lacking for other vein deposits. This contrast may reflect the variable degree 564 of erosion of each district and/or the difference in emplacement depth of individual Mo mineralized 565 porphyry systems. We predict that unexposed porphyry Mo deposits may occur beneath and/or to the 566 side of some Pb-Zn-Ag veins where magmatic intrusions are lacking at surface. The Pb-Zn-Ag veins at 567 Laodaizhanggou (Fig. 2) may be such an example. These veins predate the Donggou porphyry Mo 568 deposit by ca. 7 m.y. and thus are unlikely formed by magmatic-hydrothermal fluids from Donggou. 569 Laodaizhanggou is broadly coeval with the Taishanmiao intrusive complex and the Zhuyuangou quartz 570 - molybdenite veins, but the large distance between the former and the latter (>8 km; Fig. 2) suggests 571 that genetic link with the Taishanmiao magmatism is less likely. We therefore suggest that a concealed 572 magmatic intrusion, possibly with Mo mineralization, is located at depth beneath and/or to the side of 573 Laodaizhanggou. Given the distribution of Pb-Zn-Ag vein deposits in the area, we propose that a 574 porphyry Mo deposit may be located at depth between Laodaizhanggou, Xizaogou, and Liezishan (Fig. 575 2).

576 Silver-Pb-Zn veins are particularly developed in the Xiayu ore field, to the west of the 577 Xiong'ershan district (Fig. 1). These veins have been the largest silver and an important Pb + Zn 578 producer in central China (Chen et al., 2004; Mao et al., 2006; Li et al., 2013b). Previous studies favor 579 a magmatic origin for these Ag-Pb-Zn veins (Fig. 1; Mao et al., 2006; Ye, 2006; Gao et al., 2010a, 580 2011). While no porphyry Mo deposits have been recognized, as yet, with these large Ag-Pb-Zn 581 resources in the Xiayu ore field, we propose that Xiayu ore field may be a potential exploration target 582 for porphyry Mo deposits.

#### 583 Conclusions

The Fudian ore field in the southern NCC contains the Donggou porphyry Mo deposit and several Pb-Zn-Ag vein deposits (e.g., Sanyuangou, Wangpingxigou, Laodaizhanggou). New geochronological data show that the Donggou porphyry (LA-ICP-MS zircon U-Pb =  $117.8 \pm 0.9$  Ma), the Donggou Mo deposit (molybdenite Re-Os =  $117.5 \pm 0.8$  and  $116.4 \pm 0.6$  Ma), and the Sanyuangou Pb-Zn-Ag veins (sericite  ${}^{40}$ Ar/ ${}^{39}$ Ar =  $115.9 \pm 0.9$ ) formed contemporaneously in a short period during the Early 589 Cretaceous. Trace element compositions of pyrite from Donggou and Sanyuangou are consistent with 590 metal zonation from central Mo ores to peripheral Pb-Zn-sulfide veins, typical of porphyry Cu/Mo 591 deposits worldwide. In-situ lead isotopic analysis of pyrite suggests a common metal source for the 592 Donggou porphyry Mo deposit and Sanyuangou Pb-Zn-Ag vein deposit. The trace element and lead 593 isotope data of pyrite thus confirm a magmatic-hydrothermal origin for the Sanyuangou Pb-Zn-Ag 594 veins and its genetic link to the Donggou porphyry Mo deposit. Recognition of a porphyry-related 595 magmatic-hydrothermal system in the Fudian ore field has significant implications for future 596 exploration of Mo and Pb-Zn-Ag resources in the southern NCC. Where either base metal veins or 597 porphyry Mo ore bodies are known in a given area, they provide a potential exploration vector for the 598 other mineralization style.

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#### 945 Figure and table captions

951

Fig. 1 Geological map showing major mineral districts along the southern margin of the NCC
(modified from Mao et al., 2010; Li et al., 2013b). The insert map shows the tectonic location of the
southern NCC in Eastern China. Abbreviations: TNCO: Trans-North China Orogen, MCC:
metamorphic core complex.

950 Fig. 2 Geological map of the Fudian ore field showing the locations of the Donggou porphyry Mo

deposit and surrounding Pb-Zn-Ag vein deposits (simplified from Bureau of Geological Exploration

and Mineral Development of Henan Province, 1992). Abreviations: F1: Yangping-Wangping fault; F2:

953 Jincun-Fudian fault. For map location see Fig. 1.

Fig. 3 Geological map of the Donggou porphyry Mo deposit (after Ma et al., 2007). For map location
see Fig. 2.

Fig. 4 Representative cross sections of the Donggou porphyry Mo deposit, showing distribution and
morphology of ore bodies (after Ma et al., 2007). For map location see Fig. 3.

958 Fig. 5 Photographs showing occurrences and structures of ores at Donggou. (a) The contact zone 959 between the mineralized granite porphyry and the andesite of the Xiong'er Group. Note the 960 molybdenite disseminations both in the granite porphyry and the andesite. (b) Coarse-grained 961 molybdenite disseminated in the granite porphyry. Potassic alteration is well developed in the granite 962 porphyry. (c) Hydrothermal vein consisting of coarse-grained K-feldspar and molybdenite filling the 963 fracture of the andesite. (d) Both the K-feldspar - molybdenite vein and quartz - molybdenite vein 964 cutting the andesite. Note that the K-feldspar - molybdenite veins may be overprinted by a quartz -965 molybdenite vein. (e) Pyrite-sphalerite-molybdenite vein filling the fracture of the Mo-mineralized 966 andesite. (f) Polymetallic vein cutting the andesite. The sulfide minerals in the vein include pyrite, 967 molybdenite, and galena. Abbreviations: Mo: molybdenite, Kfs: K-feldspar, Qz: quartz, Py: pyrite, Sp: 968 sphalerite, Gn: galena.

969 Fig. 6 Transmitted-light (a) and reflected-light (b-f) photomicrographs illustrating the mineralogy and 970 textures of the granite porphyry and sulfide ores at Donggou, respectively. (a) Perthite and quartz 971 phenocrysts in the granite porphyry. (b) Stage D1 molybdenite aggregates intergrown with K-feldspar 972 and quartz. (c) Coarse-grained stage D2 molybdenite and quartz vein cutting the potassic zone. (d) 973 Stage D3 chalcopyrite and galena filling microfractures in coarse-grained, euhedral pyrite. (e) Stage D3 974 sphalerite intergrown with chalcopyrite and galena cutting massive quartz. (f) Stage D3 galena
975 replacing sphalerite. Abbreviations: Pth: Perthite, Ccp: chalcopyrite, other abbreviations as in Figure 5.

976 Fig. 7 Paragenetic sequence of the Donggou deposit.

977 Fig. 8 Photographs showing occurrences and structures of the ores at Sanyuangou. (a) Sulfide veinlets 978 penetrating quartz and ankerite in a steeply-dipping fault zone cutting the quartz diorite. (b) Quartz-979 galena-sphalerite vein as fracture infillings in the quartz diorite. (c) Coarse-grained sphalerite and 980 galena aggregates in quartz diorite. (d) Massive sphalerite and galena ores replacing quartz diorite 981 breccias. (e) Massive galena-dominated ores. (f) Textural relationship of a vein consisting of the outer 982 siderite and ankerite, the intermediate galena and quartz, and the inner coarse-grained calcite. 983 Abbreviations: Sd: siderite, Ank: ankerite, Cal: calcite; other abbreviations as in Figure 5.

984 Fig. 9 Reflected-light (a-d) and transmitted-light (e-f) photomicrographs illustrating the mineralogy 985 and textures of sulfide ores and associated alteration at Sanyuangou. (a) Stage S1 pyrite was fractured 986 and filled by paragenetically late chalcopyrite, sphalerite, and galena. (b) Stage S2 bornite and 987 sphalerite replacing pyrite; all of which were altered by galena. Note that tetrahedrite is intergrown 988 with bornite. (c) Stage S2 covellite forming selvages around foam-like pyrite; chalcopyrite forming 989 inclusions in sphalerite. (d) Stage S2 sphalerite with chalcopyrite inclusions filled and replaced by 990 galena. (e) Stage S3 calcite intergrown with coarse-grained quartz. (f) Sericite aggregates intergrown 991 associated with Pb-Zn sulfide mineralization. Abbreviations: Ccp: chalcopyrite, Bn: bornite, Ttr: 992 tetrahedrite, Cv: covellite, Cal: calcite, Ser: sericite; other abbreviations as in Figure 5.

993 Fig. 10 Paragenetic sequence of the Sanyuangou deposit.

994 Fig. 11 U-Pb concordia diagrams of zircons from the Donggou granite porphyry (a), the Sanyuangou

995 quartz diorite (b), and the Wangpingxigou quartz monzonite (c).

**Fig. 12**<sup>40</sup>Ar/<sup>39</sup>Ar age spectra (a) and inverse isochron (b) of stage S2 sericite from Sanyuangou.

997 Fig. 13 Diagrams showing the concentrations of selected trace elements for pyrite from Donggou and

- 998 Sanyuangou. Red dotted line indicates the detection limits of the trace elements.
- Fig. 14 Correlation between Co and Ni for pyrite from Donggou and Sanyuangou. The shadow areasshow the average detection limit of the trace elements.
- 1001 Fig. 15 Thorogenic (a) and uranogenic (b) lead isotopic diagrams for pyrites from Donggou and
- 1002 Sanyuangou. The dotted lines are trend lines for the spots. NHRL shows the North hemisphere
- 1003 reference line (Hart, 1984).

- 1005 **Table 1** Re-Os isotope data of molybdenite from the Donggou deposit.
- **Table 2**<sup>40</sup>Ar/<sup>39</sup>Ar analytical results of stage S2 sericite from Sanyuangou.
- 1007 **Table 3** Lead isotopic compositions of pyrite from Donggou and Sanyuangou
- 1008 Appendix 1 Laser ablation ICP-MS zircon U-Pb dating results for the Donggou granite porphyry,
- 1009 Sanyuangou quartz diorite, and Wangpingxigou quartz monzonite.
- 1010 Appendix 2 Laser ablation ICP-MS spot analyses of pyrite from Donggou and Sanyuangou
- 1011













Stage Mineral	D1	D2	D3	D4
Molybdenite				
Scheelite				
Magnetite				
Pyrite				
Chalcopyrite				
Sphalerite				
Galena				
K-feldspar				
Quartz				
Biotite				
Sericite				
Chlorite				
Fluorite				
Anhydrite				
Calcite				

abundant — common ----- minor





Stage	S1	S2	S3
Pyrite			
Pyrrhotite			
Sphalerite			
Galena			
Chalcopyrite			
Bornite			
Tetrahedrite			
Covellite			
Quartz			
Sericite			
Chlorite			
Siderite			
Ankerite			
Calcite			
Fluorite			
	I	I	I

abundant ----

----- minor











Table 1 Re-Os isotopic data of molybdenite from the Donggou deposit

	1 5		00	1							
Sample No.	Location	Weight	Re	±	<sup>187</sup> Re	±	<sup>187</sup> Os	±	Age (Ma)	±a	±b
		(ing)	(ppiii)		(ppm)		(ppu)		(Ivia)		
DG03	Quartz-molybdenite vein	27	2.6	0.01	1.7	0.01	3.2	0.02	117.5	0.8	0.9
FDG08	Quartz-polymetallic sulfide vein	20	7.1	0.03	4.4	0.02	8.6	0.03	116.4	0.6	0.7

Note: <sup>a</sup> analytical uncertainty, <sup>b</sup> analytical and decay constant uncertainty.

 Table 2 <sup>40</sup>Ar/<sup>39</sup>Ar analytical results of the ore-related sericite from the Sanyuangou deposit

Table 2	Al/Al allaly	fucal results of t	the ore-related s	sencite nom u	le Saliyualigou	deposit				
T (°C)	<sup>40</sup> Ar/ <sup>39</sup> Ar	<sup>36</sup> Ar/ <sup>39</sup> Ar	$^{36}\text{Ar}/^{39}\text{Ar}$ $^{37}\text{Ar}/^{39}\text{Ar}$ $^{38}\text{Ar}/^{39}\text{Ar}$ $^{40}\text{Ar}^{*}(\%)$ $^{40}\text{Ar}^{*}/^{39}\text{Ar}_{K}$		<sup>39</sup> Ar <sub>K</sub> (×10 <sup>-14</sup> mol)	<sup>39</sup> Ar <sub>Cumulative</sub> (%)	Age (Ma)	± 1σ (Ma)		
Sample S	SY02, J value	= 0.004606, Tot	al age = 115.1 ]							
600	137.919	0.443	0.000	0.103	5.03	6.931	0.05	0.21	57	10
700	31.308	0.064	0.129	0.027	39.51	12.371	0.23	1.28	100	1.8
800	30.214	0.053	0.000	0.022	48.44	14.636	1.57	8.42	117.7	1.2
850	15.851	0.005	0.000	0.013	90.53	14.351	0.58	11.67	115.5	1.1
900	14.868	0.001	0.007	0.013	97.12	14.441	1.29	16.94	116.1	1.1
950	14.569	0.001	0.007	0.013	98.76	14.388	1.91	25.63	115.6	1.1
1000	14.866	0.001	0.032	0.013	97.22	14.453	11.72	79.05	116.3	1.1
1050	14.760	0.001	0.067	0.013	97.52	14.395	2.27	89.41	115.8	1.1
1150	15.173	0.003	0.383	0.014	93.62	14.210	0.99	93.93	114.4	1.2
1250	22.021	0.028	0.051	0.018	62.07	13.669	1.02	98.58	110.1	1.2
1400	147.832	0.047	0.000	0.099	6.88	10.164	0.31	100	82.5	2.3
Note: the	e terms <sup>40</sup> Ar* a	nd <sup>39</sup> Ar <sub>K</sub> denote	radiogenic <sup>40</sup> A	r and necleoge	nic <sup>39</sup> Ar, respec	ctively.				

Table 3 In-situ lead isotopic data for pyrite from Donggou and Sanyuangou

Deposit	Sample no.	Pb <sup>208</sup> /Pb <sup>206</sup>	2 σ	Pb <sup>207</sup> /Pb <sup>206</sup>	2 σ	Pb <sup>208</sup> /Pb <sup>204</sup>	2 σ	Pb <sup>207</sup> /Pb <sup>204</sup>	2 σ	Pb <sup>206</sup> /Pb <sup>204</sup>	2 σ	<sup>205</sup> Tl/ <sup>203</sup> Tl	2 σ	Total Pb	Total Tl
*	FDG09-1	2.198	0.0002	0.890	0.0001	38.147	0.009	15.445	0.003	17.357	0.002	2.425	0.00009	30.08	2.58
	FDG09-2	2.198	0.0001	0.890	0.0001	38.164	0.006	15.447	0.003	17.361	0.002	2.425	0.00009	37.13	2.59
	FDG09-3	2.198	0.0014	0.889	0.0004	38.155	0.098	15.419	0.027	17.328	0.034	2.425	0.00008	15.30	2.60
	FDG10-1	2.198	0.0006	0.890	0.0004	38.140	0.087	15.427	0.036	17.332	0.039	2.426	0.00009	1.79	2.76
Donggou	FDG10-2	2.199	0.0005	0.889	0.0002	38.243	0.067	15.452	0.022	17.385	0.022	2.426	0.00011	0.95	2.77
	FDG10-3	2.198	0.0004	0.890	0.0003	38.426	0.089	15.551	0.038	17.471	0.038	2.426	0.00010	0.54	2.75
	FDG10-4	2.199	0.0002	0.889	0.0002	38.118	0.039	15.408	0.014	17.335	0.016	2.426	0.00010	8.16	2.72
	FDG11-1	2.189	0.0071	0.890	0.0008	38.396	0.196	15.550	0.052	17.489	0.046	2.426	0.00010	0.62	2.60
	FDG11-2	2.196	0.0011	0.891	0.0006	38.080	0.168	15.439	0.069	17.336	0.085	2.426	0.00009	0.37	2.77
	FDG11-3	2.199	0.0008	0.888	0.0007	38.436	0.191	15.527	0.090	17.517	0.088	2.426	0.00010	7.12	2.73
	mean	2.197	0.0013	0.890	0.0004	38.230	0.095	15.466	0.035	17.391	0.037	2.426	0.00009	10.21	2.69
	SY04-1	2.198	0.0002	0.892	0.0001	38.012	0.011	15.432	0.005	17.296	0.005	2.426	0.00010	11.49	2.71
	SY04-2	2.199	0.0001	0.893	0.0000	38.021	0.003	15.443	0.001	17.295	0.001	2.426	0.00010	37.55	2.73
	SY04-3	2.195	0.0012	0.890	0.0006	38.008	0.106	15.454	0.040	17.323	0.046	2.425	0.00009	0.48	2.73
	SY04-4	2.191	0.0012	0.888	0.0005	38.218	0.085	15.489	0.037	17.440	0.040	2.425	0.00007	0.48	2.76
	SY04-5	2.198	0.0009	0.889	0.0006	38.337	0.175	15.566	0.076	17.495	0.084	2.425	0.00009	0.26	2.66
Sanyuangou	SY05-1	2.196	0.0005	0.892	0.0003	38.048	0.036	15.451	0.017	17.318	0.017	2.425	0.00009	6.23	2.69
	SY05-2	2.203	0.0003	0.894	0.0002	38.047	0.007	15.445	0.003	17.278	0.003	2.425	0.00008	27.78	2.66
	SY06-1	2.198	0.0002	0.893	0.0002	38.006	0.014	15.432	0.007	17.286	0.005	2.425	0.00009	17.19	2.71
	SY06-2	2.199	0.0002	0.893	0.0001	37.991	0.024	15.431	0.010	17.273	0.011	2.425	0.00012	1.45	2.73
	SY06-3	2.199	0.0001	0.893	0.0001	38.012	0.005	15.435	0.002	17.286	0.002	2.425	0.00010	55.60	2.75
	mean	2.198	0.0005	0.892	0.0003	38.070	0.047	15.458	0.020	17.329	0.021	2.425	0.00009	15.85	2.71

Appendix 1 Results of LA-ICP-MS zircon U-Pb dating for the Donggou granite porphyry, Sanyuangou quartz diorite, and Wangpingxigou quartz monzonite

<b>C</b> .	ть	TI	Th/II			Isotope	ratio					Ag (Ma	)		
Spot	Th	U	Th/U	<sup>207</sup> Pb/ <sup>206</sup> Pb	$\pm 1\sigma$	<sup>207</sup> Pb/ <sup>235</sup> U	± 1σ	<sup>206</sup> Pb/ <sup>238</sup> U	$\pm 1\sigma$	<sup>207</sup> Pb/ <sup>206</sup> Pb	±1σ	<sup>207</sup> Pb/ <sup>235</sup> U	$\pm 1\sigma$	<sup>206</sup> Pb/ <sup>238</sup> U	±1σ
Donggou gra	nite porpl	nyry, san	nple DO	G01											
DG01-1	389	381	1.02	0.0490	0.0029	0.1236	0.0071	0.0184	0.0003	150	58	118	6	118	2
DG01-2	247	385	0.64	0.0496	0.0030	0.1238	0.0069	0.0184	0.0002	176	147	119	6	118	2
DG01-3	1911	3722	0.51	0.0550	0.0012	0.1411	0.0033	0.0184	0.0002	413	50	134	3	118	1
DG01-4	279	302	0.92	0.0495	0.0032	0.1229	0.0074	0.0185	0.0003	172	150	118	7	118	2
DG01-5	177	417	0.43	0.0488	0.0023	0.1239	0.0058	0.0185	0.0002	200	118	119	5	118	1
DG01-6	296	335	0.88	0.0501	0.0039	0.1253	0.0097	0.0184	0.0003	211	181	120	9	118	2
DG01-7	389	379	1.03	0.0491	0.0025	0.1232	0.0060	0.0184	0.0002	154	150	118	5	118	2
DG01-8	411	437	0.94	0.0513	0.0033	0.1289	0.0087	0.0185	0.0003	254	148	123	8	118	2
DG01-9	801	693	1.16	0.0485	0.0017	0.1240	0.0045	0.0185	0.0002	124	83	119	4	118	1
DG01-10	1026	600	1.71	0.0500	0.0033	0.1281	0.0086	0.0184	0.0002	195	156	122	8	118	1
DG01-11	320	279	1.15	0.0507	0.0044	0.1254	0.0107	0.0185	0.0004	228	204	120	10	118	3
DG01-12	311	360	0.87	0.0498	0.0031	0.1234	0.0074	0.0185	0.0003	187	146	118	7	118	2
DG01-13	457	539	0.85	0.0589	0.0035	0.1440	0.0076	0.0184	0.0003	561	127	137	7	117	2
Sanyuangou	quartz di	orite, sai	mple S	Y01											
SY01-1	2403	1350	1.78	0.1085	0.0016	4.7579	0.0842	0.3168	0.0041	1776	27	1778	15	1774	20
SY01-2	1058	739	1.43	0.1096	0.0013	4.8453	0.0646	0.3187	0.0021	1794	22	1793	11	1783	10
SY01-3	997	739	1.35	0.1078	0.0013	4.6202	0.0592	0.3093	0.0019	1762	22	1753	11	1737	10
SY01-4	531	393	1.35	0.1075	0.0011	4.6456	0.0577	0.312	0.0024	1767	19	1758	10	1750	12
SY01-5	701	544	1.29	0.1079	0.0008	4.6809	0.0437	0.3129	0.0017	1765	13	1764	8	1755	8
SY01-6	1362	761	1.79	0.1071	0.0007	4.6555	0.0382	0.3131	0.0012	1751	13	1759	7	1756	6
SY01-7	1224	815	1.50	0.1069	0.0007	4.6203	0.0378	0.3113	0.0019	1748	11	1753	7	1747	9
SY01-8	358	518	0.69	0.1062	0.001	4.6717	0.0699	0.3165	0.0038	1735	17	1762	13	1772	18
SY01-9	987	835	1.18	0.1062	0.0006	4.6689	0.0355	0.3163	0.0017	1735	10	1762	6	1772	9
SY01-10	1275	942	1.35	0.1068	0.0007	4.7115	0.0436	0.3173	0.0017	1746	21	1769	8	1776	9
SY01-11	450	419	1.07	0.1086	0.0011	4.7737	0.0613	0.3167	0.0025	1776	14	1780	11	1774	12
SY01-12	491	481	1.02	0.1077	0.0012	4.7042	0.0645	0.3149	0.0024	1761	21	1768	11	1765	12
SY01-13	1149	889	1.29	0.1084	0.0014	4.8193	0.0706	0.3211	0.0025	1773	24	1788	12	1795	12
SY01-14	650	612	1.06	0.1074	0.0017	4.6636	0.0745	0.3141	0.0022	1767	28	1761	13	1761	11
Wanpingxigo	ou quartz i	monzoni	te, sam	ple WP01						1000					
WP01-1	993	885	1.12	0.1118	0.0017	5.0921	0.0813	0.3282	0.0023	1829	23	1835	14	1830	11
WP01-2	343	324	1.06	0.1119	0.0019	5.0678	0.0838	0.3272	0.0027	1831	25	1831	14	1825	13
WP01-3	251	214	1.17	0.1166	0.0019	5.3102	0.0992	0.3282	0.0027	1905	34	1871	16	1830	13
WP01-4	433	397	1.09	0.1118	0.0014	5.05	0.0627	0.3265	0.0018	1829	23	1828	11	1821	9
WP01-5	/38	592	1.25	0.1134	0.0008	5.1546	0.0536	0.3283	0.0023	1855	45	1845	9	1830	11
WP01-6	195	221	0.88	0.1141	0.0012	5.1774	0.0668	0.3276	0.0021	1865	19	1849	11	1827	10
WP01-/	/12	618	1.15	0.1137	0.001	5.1542	0.04/8	0.3272	0.0015	1859	17	1845	8 10	1825	/
WP01-8	46/	400	1.1/	0.1116	0.0017	5.0022	0.0736	0.3231	0.0023	1826	28	1820	12	1805	11
WP01-9	455	419	1.09	0.1127	0.0009	5.1369	0.04/5	0.3278	0.0017	1844	15	1842	8 10	1828	8 10
WP01-10	16/	258	0.65	0.1103	0.0016	4.9928	0.0706	0.3257	0.0026	1806	2/	1818	12	1817	13
WP01-11	426	420	1.01	0.1102	0.0012	4.9/1/	0.058/	0.324	0.0010	1802	19	1815	10	1809	8
WP01-12	2/9	308	0.91	0.1114	0.0013	5.0624	0.1120	0.3264	0.0018	1022	10	1030	11	1021	9 17
WPU1-13	203	205	U./I 1 40	0.1119	0.0023	5.0896 E 1200	0.1138	0.3269	0.0035	1031	69 27	1034	19	1025	17
WP01-14	000 201	44/ 595	1.40 1.27	0.1122	0.0017	5.1300	0.0002	0.3293	0.0025	1820	∠/ 30	1043	15	1822	12
WP01-15	001	202	1.37	0.1112	0.0020	5.0050	0.0300	0.3207	0.0019	1020	32	1034	15	1032	9

Doporit	-ICP-WI3 Spot alla	Analusia na	C-	Ma	Fail	Ca	NI:	C.u.	7	۸	6. M		۸	C.I	6-	ch.	Te	P.	C.I	116	Ta	147	D.	A	71	DI-	D:
Deposit	Sample no.	Analysis no.	CT	20.22	Fe"	01.67	INI 70.12	106.75	2007.60 Ga	AS	5e M	0 26	Ag	22.20	50	31.32	10	ва	<0.02	HI	1a	W 0.20	11	AU 0.22	0.21	209.04	20.90
	FDG09-2-1 FDC09-2-2	AU01A005	<4.89	38.23	460000	6 20	/0.12	106.75	3007.09 0.	251.35	2.94 2	2.30	55.55 10.27	22.26	0.00	31.22	4.91	0.21	<0.03	0.02	<0.01	0.28	~0.02	0.32	0.21	398.04	29.80
	FDG09-2-2	AU01A008	<2.49	100.50	460000	7.03	6.40	67.29	454.05 0.	9 44.00	1.54 <0.04	0.00	0.07	2.52	0.20	0.72	0.27	<0.06	~0.01	<0.01	<0.01	<0.01	<0.01	<0.01	0.52	22 72	2.54
	FDG09-2-3	AU01A007	<3.10	2.60	460000	7.03	4.69	12.10	24.78 <0.02	0.34	2.95	0.00	1.57	0.15	0.20	1.20	1.56	<0.00	<0.02	<0.02 0.02	<0.01	<0.01 0.10	<0.01	~0.01 0.0C	0.02	25.72	4.76
	FDG09-3-4	AU01A008	<2.50	<0.11	460000	6.72	4.00	2.09	0.21 <0.02	4.30	1.16 < 0.02	4.23	0.09	0.43 2.43	0.10	0.94	0.92	<0.07	<0.01	<0.02	<0.01	<0.10	<0.01	0.00	<0.05	52.79	4.70
	FDG09-3-5	AU01A009	< 3.05	0.11	460000	25.15	10.44	15 77	27.44 <0.01	1021.40	1.16 <0.02		0.09	~0.07 0.20	0.12	1.02	0.02	~0.05	~0.04	<0.01	<0.01	<0.01	<0.01	<0.01	0.01	7 20	2.70
	FDC09.4.7	AU01A011	<2.75	10.40	460000	1.17	0.09	1.47	0.49 <0.02	4.55 5.49	2.54 <0.01		0.41	<0.09	0.11	0.00	0.00	<0.05	<0.02	<0.01	<0.01	<0.01	<0.02	-0.01	0.01	9.37	0.00
	FDC09.4.9	AU01A012	<2.54	<0.15	460000	1.17	0.00	1.47	0.45 <0.02	2.74	2.34 <0.01		0.17	~0.00	0.14	0.05	0.00	<0.03	~0.01	<0.01	<0.01	<0.01	<0.01	0.01	<0.01	2.92	0.03
	FDG09-4-9	AU01A012	<2.57	0.15	460000	2 34	1.40	6.41	0.03 <0.02	17.32	2.03 <0.04		0.52	<0.10	0.14	0.10	1.27	-0.04	<0.01	<0.02	<0.01	<0.02	<0.02	0.03	0.02	10.38	0.03
	FDG09-4-10	AU01A014	<4 54	78.99	460000	4.56	5.11	271 54	7512.98 <0.03	113.81	1.80	0.13	3.12	49 72	1 31	0.58	9.73	<0.04	<0.01	<0.04	<0.01	<0.01	<0.02	<0.04	0.01	23 34	0.41
Donggou	FDG11-1-1	AU01A015	<3.62	<0.10	459000	0.94	0.19	6.63	78.28 0	1362.45	2.49	0.05	3 50	0.35	0.10	0.19	<0.07	<0.06	<0.01	<0.04	<0.01	<0.01	<0.01	0.15	0.00	39.12	0.01
	FDG11-1-2	AU01A016	<2.82	<0.10	459000	0.14	0.10	0.05	0.35 <0.02	390.22	2.45	<0.05	02 4	<0.05	0.10	<0.01	<0.07	<0.00	<0.01	<0.01	<0.01	<0.01	<0.01	<0.01	<0.00	0.07 <	:0.01
	FDG11-3-3	AU01A017	<3.73	<0.11	459000	3.12	1.52	0.84	0.32 <0.02	1.76	2.05 0.05	0.03	0.05	<0.05	0.12	0.07	0.46	0.01	<0.03	0.01	<0.01	<0.01	<0.02	<0.01	<0.01	2.28	0.01
	FDC11.2.4	AU01A019	<3.10	<0.13	450000	24.99	11.02	4 59	0.32 <0.02	04.21	2.03	0.03	0.05	~0.05	0.12	0.67	1.49	<0.07	<0.04	<0.04	<0.01	<0.01	<0.02	<0.01	0.01	7.27	1.00
	FDG11-3-5	AU01A019	<3.80	-0.12	459000	491.57	115.34	1.73	<0.32 <0.02	37.90	2.51 0	0.05	0.03	<0.07	0.11	0.02	3.13	<0.01	<0.01	<0.01	<0.01	<0.01	<0.01	<0.01	0.01	11.65	1.00
	FDG11-3-6	AU01A020	<2.54	<0.11	459000	55.37	12.99	0.40	0.39 <0.02	1.84	2.63 <0.02	<0.00	01 4	<0.06	0.10	0.02	<0.17	<0.00	<0.04	<0.01	<0.01	<0.01	<0.01	0.01	<0.01	0.89	0.43
	FDG11-4-7	AU01A021	27.20	25.43	459000	180.72	23.06	3.76	3.49 2	5 74.88	6.05 (	0.44	1 30 4	<0.11	0.68	0.62	2.65	2.83	18 39	<0.01	<0.01	0.51	<0.02	<0.01	0.39	133 21	4 10
	FDG11-4-8	AU01A022	7 74	118.06	459000	112.44	22.00	8.78	15.96 5	79 53.67	1.95	1.76	2.38	<0.13	1 72	2 71	3.62	12.00	0.48	0.02	<0.01	0.35	<0.02	<0.01	1.55	156.19	2 77
	FDG11-4-9	AU01A023	<2.73	3.84	459000	61.63	36.62	2.25	104 0	38 136.08	1.59 < 0.02	1.70	0.31	<0.06	0.23	2.71	0.48	1.53	<0.40	<0.01	<0.01	0.07	<0.01	<0.01	0.49	25.63	1.63
	FDG11-4-10	AU01A024	<2.97	9.71	459000	1224.09	116.10	2.93	<0.45 0.	27 49.95	8.86 < 0.05		1.63	<0.00	1.77	0.20	0.40	0.85	1.26	0.05	0.01	0.05	<0.02	0.16	<0.45	10.86	5.77
	SV04-1-1	AU01A025	<2.06	<0.08	454000	3.05	1 13	7.36	0.63 <0.02	17213 55	1.47 <0.01		0.05	<0.03	0.09	<0.01	<0.12	<0.03	<0.01	<0.03	<0.03	<0.01	<0.01	11.96	<0.01	0.26 <	0.01
	SY04-1-2	AU01A026	<2.82	<0.10	454000	5.84	0.29	4.25	0.29 < 0.01	4916.60	2.17 < 0.03		0.10	<0.05	0.11	<0.02	<0.08	<0.03	< 0.01	< 0.01	0.00	<0.01	< 0.01	0.58	< 0.01	0.05 <	:0.01
	SY04-1-3	AU01A027	<2.55	<0.08	454000	382.93	14.25	2.90	0.25 < 0.02	7337.75	2.36 < 0.03	<0.	02 .	<0.07	0.12	<0.02	<0.10	<0.03	< 0.03	< 0.01	< 0.01	< 0.01	< 0.01	4.63	<0.01	0.12	0.01
	SY04-2-4	AU01A028	<2.70	<0.13	454000	1289.78	34.08	475.78	0.54 < 0.02	2928.28	2.25 < 0.01		5.19	<0.08	0.13	0.14	<0.14	<0.07	0.03	< 0.01	< 0.01	< 0.01	< 0.01	0.44	<0.01	8.64	0.01
	SY04-2-5	AU01A029	<2.54	<0.08	454000	121.18	1.48	2.73	0.20 < 0.02	9937.71	2.11 < 0.01		0.02	<0.08	0.14	<0.01	<0.10	<0.04	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	21.96	<0.01	0.02 <	:0.01
	SY04-2-6	AU01A030	6.25	< 0.09	454000	33.12	2.86	9.37	10.67 < 0.02	13113.92	2.14 < 0.02		0.27	0.07	0.10	1.79	<0.13	0.04	< 0.02	< 0.01	< 0.01	< 0.01	< 0.02	8.46	<0.01	1.04 <	:0.01
	SY04-2-7	AU01A031	<2.51	<0.10	454000	1.64	0.26	1355.14	3.18 < 0.01	4469.19	2.26 < 0.02		8.62	<0.04	0.11	<0.01	<0.09	<0.03	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	1.09	<0.01	2.50 <	:0.01
	SY04-2-8	AU01A032	<2.67	3.97	454000	3.37	2.53	185.19	840.75 0.	05 7810.34	<1.31 <0.02		4.38	3.39	1.12	2.31	< 0.14	< 0.01	< 0.03	< 0.01	< 0.01	< 0.01	< 0.02	12.18	< 0.01	18.18 <	:0.01
	SY04-4-9	AU01A033	<2.78	0.51	454000	601.95	255.99	7939.22	1.39 < 0.01	< 0.30	<1.36 <0.03		1091.60	0.28	0.61	0.73	0.10	< 0.01	< 0.01	< 0.02	< 0.01	0.25	< 0.01	< 0.01	2.18	167.53	0.01
	SY04-4-10	AU01A034	<2.12	0.16	454000	636,78	300.24	6161.43	59.80 0.	0.83	2.15 < 0.01		205.95	1.22	5.50	7.49	0.11	< 0.01	< 0.02	< 0.01	0.00	0.11	< 0.01	< 0.01	1.51	637.60 <	<0.01
	SY03-1-1	NO06A003	3.16	84.97	456000	30.66	37.47	2650.71	11.47 0.	16 1102.18	1.56 < 0.01		16.00	0.14	0.16	2.06	< 0.12	< 0.02	< 0.02	0.01	0.00	0.14	< 0.01	< 0.01	0.04	34.31 <	:0.01
	SY03-1-2	NO06A004	< 0.44	2.46	456000	62.20	74.73	11087.95	26893.67 0.	04 229.96	1.94 < 0.01		85.69	203.54	0.12	0.77	< 0.13	< 0.06	< 0.04	< 0.01	< 0.01	< 0.02	< 0.01	< 0.01	0.01	101.06 <	:0.01
	SY03-1-3	NO06A005	1.03	17.25	456000	55.29	55.72	12666.89	2051.15 0.	04 136.12	0.90	0.02	68.54	17.99	0.12	0.56	< 0.17	< 0.07	< 0.02	0.03	0.02	0.06	< 0.01	< 0.01	0.01	43.08 <	:0.01
	SY03-1-4	NO06A006	< 0.33	10.59	456000	64.37	4.41	11506.79	8944.66 < 0.02	2955.85	1.72 < 0.02		146.72	69.70	0.16	16.77	< 0.17	< 0.03	< 0.02	0.02	< 0.01	0.01	< 0.01	0.05	0.07	234.89	0.03
	SY03-1-5	NO06A007	< 0.47	< 0.11	456000	38.62	60.01	9428.25	15250.56 < 0.03	1071.85	1.51 < 0.03		119.18	101.27	0.37	9.79	0.21	0.04	< 0.02	< 0.01	< 0.01	< 0.01	< 0.01	0.02	0.03	176.81	0.01
Sanyuangou	SY03-1-6	NO06A008	< 0.33	2.04	456000	19.07	2.15	2588.46	4670.34 0.	3 599.00	2.57 < 0.02		47.50	38.23	0.17	7.37	< 0.19	< 0.03	< 0.03	< 0.01	0.00	< 0.01	< 0.01	< 0.01	0.02	96.38	0.01
	SY03-1-7	NO06A009	< 0.32	0.37	456000	75.41	61.23	2075.28	13.07 < 0.02	106.77	0.77 < 0.02		18.33	0.27	0.16	0.43	< 0.14	< 0.02	< 0.01	0.00	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	13.65	0.02
	SY03-1-8	NO06A010	3.13	302.27	456000	210.43	33.24	23420.20	9034.94 0.	79 300.07	1.58	0.27	170.03	69.31	0.99	1.13	< 0.16	0.39	< 0.01	0.16	< 0.01	0.05	< 0.01	< 0.01	0.03	179.43	0.02
	SY03-1-9	NO06A011	< 0.37	7.26	456000	217.89	190.66	1752.82	35.07 < 0.02	343.79	1.27 < 0.02		15.57	0.17	0.11	1.10	< 0.17	< 0.05	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	0.01	53.49 <	:0.01
	SY03-1-10	NO06A012	< 0.45	10.63	456000	26.74	7.01	4622.61	545.19 < 0.03	1628.50	0.89 < 0.01		50.72	6.88	0.13	0.24	< 0.16	< 0.02	< 0.03	< 0.01	< 0.01	0.03	< 0.01	< 0.01	0.01	43.71 <	:0.01
	SY03-2-1	NO06A013	< 0.38	1.67	456000	646.47	56.47	782.26	388.43 0.	19 2658.79	1.63 < 0.01		22.07	1.58	0.81	14.12	< 0.15	0.37	< 0.03	< 0.01	0.00	< 0.01	< 0.01	1.81	0.01	154.17 <	:0.01
	SY03-2-2	NO06A014	3.09	6.27	456000	159.52	73.23	608.32	57.03 < 0.04	1088.15	0.93 < 0.02		4.96	0.23	0.64	1.68	0.14	0.63	0.03	0.93	< 0.01	< 0.01	< 0.01	0.28	0.02	22.19 <	:0.01
	SY03-2-3	NO06A015	29.00	18.99	456000	2.19	2.68	4.57	3.10 0.	41 3286.03	1.07 < 0.02		1.34	< 0.04	0.10	1.51	< 0.13	0.87	0.11	0.60	0.01	0.23	< 0.01	0.07	0.05	34.25 <	0.01
	SY03-2-4	NO06A016	8.13	5.25	456000	5.47	2.48	3.78	0.69 0.	66 12418.22	1.84 0	0.06	0.70	0.09	0.34	0.95	0.18	4.73	0.07	0.25	0.27	0.19	< 0.01	2.45	0.06	20.02 <	0.01
	SY03-2-5	NO06A017	0.50	32.21	456000	0.47	1.15	1046.99	4513.82 0.	69 4196.21	<0.61 <0.02		29.15	17.18	2.17	14.53	< 0.18	< 0.03	0.01	< 0.01	< 0.01	< 0.01	< 0.01	2.40	0.02	165.76	0.02
	SY03-2-6	NO06A018	< 0.47	5.11	456000	29.48	8.56	295.84	455.71 0.	06 11234.28	1.26 < 0.01		5.42	1.56	0.27	2.74	< 0.17	< 0.04	0.01	< 0.01	< 0.01	< 0.01	< 0.01	3.38	< 0.01	24.68 <	:0.01
	SY03-2-7	NO06A019	< 0.32	< 0.10	456000	10.54	0.71	29.50	0.31 < 0.03	14624.10	<0.85 <0.01		0.19	< 0.06	0.09	0.04	< 0.12	< 0.02	< 0.02	< 0.01	< 0.01	< 0.01	< 0.01	15.39	< 0.01	1.00 <	0.01
	SY03-2-8	NO06A020	< 0.46	0.43	456000	9.28	0.89	100.66	47.82 <0.04	13559.60	1.28 < 0.03		2.17	0.20	0.17	0.82	< 0.19	< 0.05	< 0.03	< 0.01	< 0.01	0.01	< 0.01	12.20	< 0.01	7.57	0.01
	SY03-2-9	NO06A021	1.85	2.96	456000	137.63	49.86	175.70	132.51 0.	74 304.43	1.34	0.02	2.77	0.57	0.33	0.90	<0.24	13.18	0.02	< 0.01	< 0.01	0.03	< 0.01	0.06	0.02	31.93	0.02
	SY03-2-10	NO06A022	0.74	<0.11	456000	496.69	52.89	3.00	2.71 < 0.03	17913.81	1.75 < 0.03		0.35	<0.06	0.11	0.69	<0.20	< 0.04	< 0.04	< 0.01	< 0.01	< 0.01	< 0.01	3.53	0.02	23.33 <	:0.01
Note: " Iron was	used as the interna	il standard, and the	Fe contents	s were determi	ned by EMP	analyses																					

#### **Highlights:**

1) New geochronological data indicate that the Donggou Mo deposit and the spatially associated Sanyuangou Pb-Zn-Ag veins formed contemporaneously in the early Cretaceous (117.5-115.9 Ma).

2) Trace element compositions and lead isotopes of pyrite by in-situ LA-ICP-MS analyses confirm a magmatic-hydrothermal origin for the Sanyuangou Pb-Zn-Ag deposit.

3) The Donggou and Sanyuangou deposits form a typical porphyry-related magmatic-hydrothermal system. This model has broad implications for further exploration of Mo and Pb-Zn-Ag resources in the area.