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Products of Wall-Rock Sulfidation by Deep Crustal **Fluids**

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Sediment-Hosted Gold Deposits in Guizhou, China: Products of Wall-Rock Sulfidation by Deep Crustal Fluids

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Abstract

Sediment-hosted gold deposits in Guizhou, China, are hosted in late Paleozoic and early Mesozoic sedimentary rocks along the southwest margin of the Precambrian Yangtze craton. They have characteristics similar to Carlin-type gold deposits in Nevada, USA and are notably enriched in As, Sb, Hg and Tl. The Shuiyindong and Yata deposits consist of disseminated, stratabound sulfides in Permian bioclastic limestone and fault-controlled mineralization in Middle Triassic calcareous clastic rocks, respectively. Mineralization in both deposits consists of barren milky quartz veins, disseminated gold-bearing arsenian pyrite and arsenopyrite, stibnite, realgar and orpiment. The barren milky quartz veins occur in the ore-controlling structures with an envelope of gold mineralization in the host rock consisting of disseminated gold-bearing arsenian pyrite and arsenopyrite and replacement-style quartz veinlets. Later drusy quartz, stibnite, realgar and orpiment fill fractures and vugs on the periphery of gold mineralization. Petrography, microthermometry, laser Raman spectroscopy and laser ablation inductively coupled plasma mass spectrometric (LA-ICP-MS) analyses of fluid inclusions are used to characterize the chemical evolution of ore fluids at Shuiyindong and Yata.

Early barren milky quartz veins trapped aqueous fluid with moderate salinity (6.0 wt.% NaCl equiv.) and minor CO_2 (<2.4 mole %) at a temperature of 230 \pm 30°C. This fluid contains measurable B, Na, K, Cs, Sr, As, Sb and Au $(3.8 \pm 0.5 \text{ to } 5.7 \pm 2.3 \mu g/g)$ determined by LA-ICP-MS and is interpreted to approximate the mineralizing fluid responsible for precipitation of early barren quartz veins. Low salinity (0.9 to 2.3 wt.% NaCl equiv.), $CO₂$ -bearing (6.3 to 8.4 mole %), aqueous fluids are recorded in replacement-style quartz veinlets. They formed gold-bearing arsenian pyrite and arsenopyrite and quartz veinlets at a temperature of $210 \pm 20^{\circ}$ C and pressures of 450 to 1150 bar, corresponding to depths between 1.7 and 4.3 km under lithostatic conditions. Fluid inclusions in late stibnite-orpimentrealgar-quartz veins contain high CO_2 (58 to 64 mole %) and N₂ (19.2 to 23.7 mole %) with trace CH_4 (up to 1.6 mole %). The increasing content of $CO₂$ and decrease in the concentrations of Au and As in aqueous-carbonic inclusions are interpreted to result from carbonate dissolution at the level of the deposits during gold mineralization. Covariance of Au, As, Sb, and Sr concentrations in the fluids is interpreted to reflect deposition of As- and Sb-sulfides and gold concurrent with carbonate crystallization. Iron is below detection limit $(~400 \text{ µg/g})$ in all fluid inclusion types, suggesting that the ore fluids were Fe-poor but possibly sulfur-rich, possibly explaining their exceptionally high gold contents. Iron in sulfide minerals was probably derived from dissolution of ferroan minerals in the host rocks, sulfidized by H₂S-rich fluids to precipitate arsenian pyrite, arsenopyrite, and gold. Pressure fluctuation induced by faulting resulted in local fluid immiscibility and led to late deposition of realgar, orpiment, stibnite, and calcite.

Hydrogen and oxygen isotope compositions of ore fluids (CDH_2 O = -35 to -68 ‰, N^{18} OH₂O = 4 to 16.5 ‰) indicate a metamorphic origin, possibly related to crustal thickening and prograde metamorphism during the late Yanshanian orogeny.

1. Introduction

Sediment-hosted gold deposits in Guizhou, China, are hosted in late Paleozoic and early Mesozoic sedimentary rocks along the southwest margin of the Precambrian Yangtze craton. They have characteristics similar to Carlin-type gold deposits in Nevada, USA, including notable enrichment in As, Sb, Hg and Tl (Hu et al., 2002; Peters et al., 2007). Most deposits are stratabound and/or faultcontrolled (Zhang et al., 2003; Peters et al., 2007). The stratabound mineralization is controlled by

anticlines and is hosted in Permian bioclastic limestone (e.g., Shuiyindong). The fault-controlled mineralization occurs along compressive shear zones in Middle or Lower Triassic siltstone and silty mudstone (e.g., Yata). Typical characteristics include impure carbonate or calcareous and carbonaceous host rock that contains disseminated pyrite and arsenopyrite. Gold occurs either as submicrometer-sized particles or invisibly as 'solid solution' in As-rich rims of pyrite and arsenopyrite (Mao, 1991; Zhang et al., 2005; Su et al., 2008). Late stibnite, realgar and orpiment fill fractures on the periphery of gold mineralization. Hydrothermal alteration caused decarbonation, silicification, argillization and sulfidation (Hu et al., 2002; Zhang et al., 2003; Su et al., 2008), similar to Carlin-type gold deposits in Nevada (Hofstra et al., 1991; Arehart, 1996; Hofstra and Cline, 2000; Emsbo et al., 2003; Kesler et al., 2003).

The physical and chemical characteristics of the ores are well constrained for Carlin-type deposits in both the USA and China. Sulfidation is recognized to be the most important mechanism of gold deposition, whereby gold and pyrite precipitated together from H_2S -rich fluids that reacted with Feminerals in the host rock package (Hofstra et al., 1991; Hofstra and Cline, 2000; Kesler et al., 2003). However, the source and evolution of ore fluids in Carlin-type systems remain a contentious issue (Cline and Hofstra, 2000), partly because the root of the ore-fluid system is difficult to identify. In the Shuiyindong and Yata deposits, quartz crystals in veins or stockworks associated with mineralization are somewhat coarser than those in most Carlin-type gold deposits. Fluid inclusions in the coarser quartz crystals are commonly 20 μm in diameter or larger and have clear paragenetic relationships of entrapment. These have been used to characterize the P-V-T-X evolution of fluids responsible for Carlin-type gold mineralization in the Shuiyindong and Yata deposits, by combining detailed petrography, microthermometry and laser Raman spectroscopy with laser ablation inductively coupled plasma mass spectrometric (LA-ICP-MS) analysis.

2. Regional Geological Setting

Carlin-type gold deposits in the southwestern Guizhou (Fig. 1) lie near the southwestern margin of the Yangtze craton, which is composed of Proterozoic low-grade metamorphic rocks overlain by thick carbonate and shale sequences of Cambrian to Triassic age (Wang et al., 1995). The outcropping strata in the area are mostly Permian and Triassic, with the Permian rocks generally exposed in the cores of anticlines. The rocks consist of limestone, dolomite, siltstone, sandstone and shale of shallowand deep-water facies. From the northwest to the southeast, the sedimentary environment gradually changes from continental and shallow-water platform to deep-water in the Youjiang rift basin. The basin margin formed in the Devonian during opening of the Paleo-Tethys (Liu et al., 2002). Most of the sedimentary rocks in the Youjiang basin have illite crystallinities indicative of low-grade burial metamorphism (anchi- to epizone: Suo et al., 1998).

Between the Middle Triassic and Early Jurassic, southwestern Guizhou was in a foreland fold- and -thrust setting on the northwest side of the Yanshanian orogen (Li and Li, 2007). The basin sequence was folded and uplifted by the orogeny as the Pacific Plate was subducted beneath the Eurasian Plate (Hu et al., 2002). The intensity of deformation caused by the orogenic event was highly variable (Wang et al., 1995). The Permian and Triassic carbonate and clastic platform facies were gently folded or domed and exhibit brittle deformation. Deep basin rocks affected by the orogenic event to the southeast (e.g., in the vicinity of Yata) have been tightly folded resulting in high-angle ductile-brittle reverse faults.

Granites and felsic volcanic rocks associated with the Yanshanian orogeny occur mainly in eastern China, far from the Carlin-type gold deposits. No felsic intrusive rocks were observed in the southwestern Guizhou or in the vicinity of the gold deposits. Small alkaline ultrabasic pipes and dikes cut Lower Permian to Middle Triassic strata, but well away from known gold deposits, along a N-Strending transitional sedimentary facies belt, an inferred Puding-Ceyang deep fault may have provided pathways for their emplacement (Fig. 1). The primary mineralogy of the dikes is obscured by alteration and weathering, except for relics of biotite and pyroxene. The least-altered biotite was dated at 85.0 \pm 0.2 Ma (40Ar/39Ar: Su, 2002). It may be interpreted as the onset of extension in response to the Yanshanian orogeny. The oldest volcanic rock, which only crops out in the northwestern part of the area, is alkaline flood basalt of late Permian age (Emeishan Basalt; Fig. 1). The flood basalts are highly variable in thickness and, generally, do not extend into the area of gold deposits.

Figure 1. Simplified geological map of southwestern Guizhou (modified after Zhang et al., 2003), showing the locations of the major Carlin-type gold (e.g. Shuiyindong and Yata) and antimony and mercury deposits.

The Carlin-type gold deposits in the southwestern Guizhou are hosted in both shallow- and deepwater facies sedimentary rocks of Permian and Triassic age (Fig. 1). Two styles of ore geometry have been recognized. Stratabound mineralization is controlled by anticlines or domes with ore hosted in Permian bioclastic limestone (e.g., Zimudang and Shuiyindong deposits). Fault-controlled mineralization occurs mainly along reverse faults on the flanks of anticlines or domes, with ore hosted in siltstone and silty mudstone of Middle or Lower Triassic age (e.g., Lannigou and Yata deposits). The age of gold mineralization in the southwestern Guizhou is poorly constrained because of the lack of minerals suitable for isotopic dating. However, fission-track ages on quartz from pyrite-quartz veins in the Lannigou and Baidi deposits are between 82 ± 8 Ma and 91 ± 6 Ma (Zhang and Yang, 1993). Recently, Su et al. (2009) reported Sm-Nd isochron ages of hydrothermal calcite veins coeval with gold mineralization at the Shuiyindong deposit, ranging from 134 ± 3 to 136 ± 3 Ma. Field observations suggest that bedding-parallel or reverse faults, which are important hosts for gold mineralization, were formed during the late stage of the Yanshanian orogeny (140 to 75 Ma: Hu et al., 2002), an important tectonic event in the southeastern China. Therefore, it is reasonable to conclude that gold mineralization in Guizhou probably took place during Late Jurassic to Cretaceous times.

3. Geology of the Shuiyindong and Yata Deposits

3.1. Shuiyindong

The Shuiyindong deposit is located about 20 km northwest of the town of Zhenfeng in the southwestern Guizhou (Fig. 1). It lies on the eastern limb of the Huijiabao anticline (Fig. 2A), which also hosts a cluster of deposits on its western limb, including the Zimudang deposit. Recent exploration and mining at Shuiyindong has proven gold reserves of 55 tons Au (1.8 Moz), with average gold grades from 7 to 18 g/t (Xia, 2005). All of the gold is refractory, occurring in arsenian pyrite and arsenopyrite (Su et al., 2008). The following geological description of the deposit is from Su et al., (2008).

Sedimentary rocks in the district consist of bioclastic limestone, siltstone and argillite of Middle and Upper Permian and Lower Triassic. The Middle Permian Maokou Formation, a massive bioclastic limestone, is overlain by the Upper Permian Longtan, Changxing and Dalong and Lower Triassic Yelang Formations. The Longtan Formation is about 300 meters thick in the Shuiyindong district and has been divided into three stratigraphic units (Liu, 2001). The lowest unit consists of calcareous siltstone, which grades into fine-grained sandstone at the top. The second unit consists of silty argillite intercalated with bioclastic limestone and coal seams. The third unit includes calcareous siltstone, sandstone, and muddy and bioclastic limestone. Gold mineralization is preferentially disseminated in bioclastic limestone and calcareous siltstone of the first and second units of the Longtan Formation at depths 100 to 300 meters below the surface (Fig. 2B). These rocks were deformed into an E-trending anticline with limbs that dip 10 to 20°. Stratabound gold orebodies are located mainly on the flanks of the anticline (Fig. 2B). The limbs of the anticline are cut by reverse faults F_{101} and F_{105} , respectively, which strike E-W and dip steeply to the north and south, respectively (Fig. 2A). The reverse faults host small orpiment and realgar bodies with low grade gold mineralization and are cut by a series of near N-S trending normal faults with steep dips (70-80°). The normal faults contain mercury-thallium deposits, such as Lanmuchang (Fig. 2A).

Figure 2. Simplified geological plan (**A**) and cross section (**B**) along the A-A' exploration line of Shuiyindong (modified from Liu, 2001). Note the stratiform nature of the numbered orebodies.

The three main orebodies at Shuiyindong (IIIa, IIIb, IIIc; Fig. 2B) contain approximately half of the gold reserves. They are 100 to 400 meters long, 50 to 350 meters wide, 1.7 to 1.9 meters thick, and have an average gold grade of 16 g/t (Xia, 2005). A lower grade orebody (Ia) is hosted in silicified, brecciated argillite and limestone at the unconformity between the Maokou Limestone and the first unit of the Longtan Formation. This unconformity controlled the distribution of other gold deposits regionally in the southwestern Guizhou, such as Getang and Nibao (Fig. 1), suggesting that it may have been a feeder conduit for gold mineralization. In these orebodies, milky quartz veins are frequently observed and occur nearly horizontal dips (Fig. 3A). Some of the milky quartz veins are cut by stibniterealgar-orpiment-quartz veinlets, suggesting that they formed later than the As-Sb mineralization but possibly predated the deposition of the bulk of disseminated gold-bearing arsenian pyrite and arsenopyrite in the host rock.

Figure 3. Examples of ores from Shuiyindong and Yata. **A.** Photograph of milky quartz veins from Shuiyindong. **B.** Slab of a quartz vein showing paragenetic relationships among calcite, stibnite and realgar from Yata. **C**. Microphotograph (crosspolarized light) of ore from Shuiyindong showing coprecipitated gold-bearing pyrite and arsenopyrite overgrowing Fe-poor dolomite. **D.** Microphotograph (cross-polarized light) of ore from Shuiyindong showing gold-bearing arsenian pyrite and arsenopyrite between quartz and dolomite on jasperoidal quartz. **E.** Backscattered electron (BSE) image showing arsenian pyrite disseminated within jasperoidal quartz from Shuiyindong. Note small relict inclusions of ferroan calcite in jasperoidal quartz. **F.** BSE image of arsenian pyrite in wall rock from Shuiyindong, with As-poor pyrite core overgrown by As-rich rim. **G.** and **H.** High-magnification BSE images of native gold grains in arsenian pyrite and calcite from Shuiyindong. **I.** Microphotograph (cross-polarized light) of typical ore from Yata showing intimate intergrowth of gold-bearing pyrite and quartz grains. **J.** Arsenopyrite and gold-bearing pyrite crystals with As-rich rims in a quartz vein from Yata (reflected light). **K.** Late-stage realgar which has overgrown euhedral quartz in a vein from Yata. Abbreviations: Asp = arsenopyrite, As-py = arsenian pyrite, Cal = calcite, Dol = dolomite, Fe-cal = ferroan calcite, Fe-dol = ferroan dolomite, Km = potassium mica, Py = pyrite, $Qz =$ quartz, Real = realgar, stb = stibnite.

Wall-rock alteration at Shuiyindong caused decarbonation, silicification, sulfidation and dolomitization. Decarbonation of limestone is evident from small relict inclusions of calcite and dolomite in jasperoidal quartz (cf. Hofstra and Cline, 2000). Small amounts of gold-bearing arsenian pyrite and arsenopyrite are enclosed within Fe-poor dolomite (Fig. 3C), whereas large amounts of gold-bearing arsenian pyrite and arsenopyrite have grown between quartz and dolomite (Fig. 3D) or enclosed in jasperoidal quartz (Fig. 3E). Sulfides in the deposit consist mainly of arsenian pyrite, arsenopyrite, marcasite, and small amounts of orpiment, realgar and stibnite. Gangue minerals are quartz, dolomite, calcite, and clay minerals (e.g., kaolinite). The dominant gold-bearing sulfides are arsenian pyrite and arsenopyrite. Arsenian pyrite formed earlier than arsenopyrite in the paragenetic sequence because arsenopyrite occurs as overgrowths on arsenian pyrite. Arsenian pyrite that contains significant gold typically occurs as tiny, irregular grains, generally less than 50 μm in size. Commonly, arsenian pyrite also forms rims on early pyrite crystals (e.g., Zhang et al., 2005, fig. 2e). The core of some early pyrite has a "porous" appearance, possibly owing to conversion of marcasite to pyrite (cf. Murowchick, 1992). Modal analyses showed that the dominant form of gold is "invisible" gold in iron sulfides (95%), with subordinate free native gold (5%) (Liu, 2003). Electron microprobe analyses (EMPA) showed that gold is predominantly in arsenian pyrite (400-3800 ppm Au, 3.37-14.06 wt.% As: Su et al., 2008), with lesser amounts in arsenopyrite (300-1500 ppm Au: Zhang et al., 2008). Lower-resolution LA-ICP-MS analyses of arsenian pyrite and arsenopyrite in ore-grade samples from Shuiyindong yielded an average of 200 to 500 ppm Au and demonstrated that Au is associated with As, Sb, Tl, Cu, Pb, Co and Ni, in decreasing order of correlation (Zhang et al., 2005). A few native gold grains were observed by Su et al. (2008). They occur in arsenian pyrite disseminated in the host rocks and in an arsenian pyrite veinlet (Fig. 3G, H). The size of these native gold grains ranges from 0.1 to 6 μm in diameter. Most are present at the edge of arsenian pyrite crystals in veinlets (Fig. 3G), and some occur in calcite (Fig. 3H), suggesting that ore fluids at Shuiyindong were locally saturated with gold (Su et al., 2008).

3.2. Yata

The Yata gold deposit is located about 15 km southwest of the town of Ceheng in the southwestern Guizhou (Fig. 1). The deposit had been mined for arsenic (realgar) at a small scale for many years. Gold mineralization was discovered as a result of reconnaissance sampling around the old realgar pits in the early 1980s. Intensive exploration was carried out by the Guizhou Bureau of Geology and Mineral Resources in the late 1980s. This resulted in the definition of more than 10 tons of gold reserves, with an average gold grade of 3 to 5 g/t (Tao et al., 1987). Geological descriptions of the deposit were given by Cunningham et al. (1988), Ashley et al. (1991) and Zhang et al. (2003).

Sedimentary rocks exposed in the district are mainly siltstone, sandstone, argillaceous limestone and shale of the Middle Triassic Xuman Formation, which is divided into four members based on sand grain size and bedding thickness (Tao et al., 1987). Gold mineralization occurs in Member 2, which is composed of interbedded sandstone, fine-grained sandstone, siltstone and mudstone. Unmineralized carbonaceous shale in this formation contains quartz, biotite, ferroan dolomite, illite, calcite and minor fine-grained diagenetic pyrite (Ashley et al., 1991).

Gold mineralization (Fig. 4) occurred along a narrow E-trending zone and comprises more than 40 orebodies (Han et al., 1999). Most of them occur along high-angle and strike-slip faults that cut the south limb of the E-trending Huangchang anticline and subsidiary folds. Limbs of the anticline dip at 35 to 70°, contain chevron folds, and mesoscopic compressive fracture zones. Faults F_1 , F_2 , F_3 and F_6 controlled mineralization within the zone. The F_2 and F_3 faults, which have easterly trends and dip south at 65 to 85°, controlled the largest mineralization zone (M_1) in the district. The M_1 zone is 1500 meters long, 40 to 60 meters wide, and 200 meters thick, with average gold grades varying from 1 to 3 g/t (Zhang et al., 2003). Smaller orebodies with average gold grades of 3 to 5 g/t occur as lenticular zones, veins, and vein-stockworks within the larger envelope of mineralization. Gold orebodies are preferentially hosted in altered calcareous siltstone and shale at intersections with high-angle faults that focused fluid flow during gold mineralization.

Figure 4. Simplified geological plan (**A**) and cross section (**B**) along the A-A' exploration line of Yata (after Zhang et al., 2003). Note the vertical (± fault-controlled) orientation of the orebodies.

Despite the stronger structural control than at Shuiyindong, wall-rock alteration at Yata is similar and extends well away from faults and fractures along more reactive or permeable clastic strata or calcareous shale and siltstone. Carbonate dissolution mostly occurred in carbonate strata. In proximal zones of carbonate dissolution, disseminated irregular or euhedral crystals of orpiment or realgar formed locally in porous decarbonated rocks. Silicification typically produced small bodies of jasperoidal quartz within the larger areas of altered rocks (Fig. 3I). Coarser quartz crystals occur in veins and veinlets (Fig. 3J, K). Quartz in the jasperoid has reticulate or xenomorphic textures, variable

grain sizes, and abundant small calcite or dolomite inclusions (Fig. 3I), suggesting that it may have formed at moderate temperatures and at greater depths (cf. Hofstra and Cline, 2000). More intense, pervasive silicification generally affected the coarser sandstone where it occurs close to feeder structures. Argillization has mainly produced illite or illite-quartz veinlets, many of which contain pyrite and arsenopyrite.

The dominant primary ore minerals at Yata are arsenian pyrite, arsenopyrite, marcasite, stibnite, orpiment and realgar (Zhang et al., 2003). Trace amounts of sphalerite, galena and chalcopyrite also occur. Gangue minerals include quartz, dolomite, calcite, and clay minerals (e.g., illite). Pyrite is the dominant sulfide in the ores (3 to 5 vol.%). It occurs disseminated in the host rocks as rounded pentagonal, dodecahedral, octahedral and cubic crystals, or as aggregates of all morphologies. Bulk analysis of pyrite has an average gold content of 83 ppm (Han et al., 1999) and arsenic contents of 1 to 7 wt. % (Ashley et al., 1991). EMPA showed that gold is predominantly in arsenian pyrite (540 ppm Au, 10.73 wt.% As: Ye et al., 1994), but a few gold grains (0.3-0.8 μm) have been observed in arsenian pyrite using scanning electron microscopy (Ye et al., 1994). Arsenopyrite is generally disseminated in the ore as tiny acicular to prismatic crystals. Some arsenopyrite occurs as rims on pyrite. Marcasite occurs as rare, tiny grains surrounding pyrite grains. Some stibnite formed early, although later than the gold-bearing pyrite, because stibnite crystals have overgrown on gold-bearing pyrite crystals at their margins. Most of the stibnite formed euhedral crystals filling fractures on the periphery of gold mineralization, local overgrowths of realgar and orpiment (Fig. 3B). Orpiment and realgar are also commonly observed in open cavities with euhedral quartz crystals (Fig. 3K) and fractures in the ore zones or their vicinities. Some calcite-orpiment-realgar veinlets cut pyrite-arsenopyrite-jasperoidal quartz veinlets.

In summary, the occurrence, textures, cross-cutting relationships and mineral assemblages of the ores at Shuiyindong and Yata indicate that ore-bearing minerals were deposited in three hydrothermal stages (Zhang et al., 2003; Hofstra et al., 2005): (1) early-stage milky quartz ± pyrite ± arsenopyrite veins; (2) main-stage vein and replacement-style quartz – dolomite – arsenian pyrite – arsenopyrite; (3) late-stage quartz – calcite \pm dolomite – stibnite – orpiment – realgar veinlets. The paragenetic sequence and associated fluid inclusion assemblages are summarized in Figure 5.

4. Samples and Fluid Inclusion Methods

Eight samples used in this study were collected from the main orebodies in the Shuiyindong and Yata deposits (Figs. 2B, 4B). Three are barren milky quartz veins that have early horizontal dips in the ore zones at Shuiyindong and fill fractures at Yata. Some of the milky quartz veins are colored orange or gray by inclusions of realgar or stibnite or are cut by stibnite-realgar-orpiment-quartz veinlets. A sample (YT-5) from Yata contains arsenian pyrite, arsenopyrite and quartz (Fig. 3J); other samples from Yata contain stibnite, realgar, orpiment, calcite and quartz (Fig. 3B, K), filling in fractures on the periphery of gold mineralization.

Doubly polished sample sections, about 200 μm thick, were examined. Fluid inclusion observations were made using a standard optical petrographic microscope and a scanning electron microscope cathodoluminescence (SEM-CL) imaging system to identify successive generations of fluid inclusionbearing vein quartz and to document its paragenetic relationship to ore minerals. These studies enabled successive assemblages of fluid inclusions to be defined. Representative samples of such assemblages were selected for detailed microthermometric, laser Raman spectroscopic, and LA-ICP-MS studies.

Mineral and fluid		Shuiyindong		Yata						
inclusion assemblages	Early-stage	Main-stage	Late-stage	Early-stage	Main-stage	Late-stage				
Vein quartz										
Arsenic-free pyrite										
Jasperoid quartz										
Arsenic-rich pyrite										
Arsenopyrite										
Marcasite										
Invisible gold										
Native gold										
Dolomite										
Illite and kaolinite										
Stibnite										
Orpiment										
Realgar										
Vein dolomite										
Vein calcite										
Galena										
Chalcopyrite										
Sphalerite										
Type Ia										
aqueous inclusions										
Type Ib aqueous										
- carbonic inclusions										
Type II carbonic- rich										
inclusions with variable										
phase proportions										
Type III mono- phase										
carbonic inclusions										

Figure 5. Paragenetic sequences of minerals and fluid inclusion assemblages at Shuiyindong and Yata, showing essentially similar mineralization history. Two-phase, liquid-rich aqueous inclusions (type Ia) occur in barren milky quartz veins in both deposits. Two- or three-phase aqueous-carbonic inclusions (type Ib) are mainly observed in quartz veinlets at Yata and in jasperoidal quartz with arsenian pyrite and arsenopyrite at Shuiyindong. Two- or monophase carbonic-rich inclusions (type II and III) occur in drusy quartz with realgar, stibnite and calcite at Yata.

Microthermometric measurements were made on a Linkam THMSG 600 programmable heatingfreezing stage mounted on a Leica microscope coupled to an image analysis system at the Institute of Geochemistry, Chinese Academy of Sciences (IGCAS) in Guiyang, China. The equipment was calibrated with melting-point standards (CCl₄, -22.99°C; KNO₃, 333°C) and the melting point of CO₂ (-56.6°C) in synthetic fluid inclusions (Sterner and Bodnar, 1984). Temperature measurements have an uncertainty of about ±0.2°C below 50°C and ±2°C for temperatures above 100°C. All fluid inclusions were analyzed by cycles of freezing down to -180°C and heating up at rate of 0.2 to 0.5°C/minute. Near the melting temperatures of carbonic phase (Tm $CO₂$) and clathrate (Tm cl), the heating rate was reduced to 0.1°C/minute.

For volatile-rich fluid inclusions, $CO₂$ was identified by melting of a solid phase in inclusions below –56.6°C. The volumetric fraction of phases in fluid inclusions was estimated at room temperature by reference to the volumetric chart of Roedder (1984), using regular-shaped, flat inclusions whenever possible according to the suggestion of Diamond (2001). Volatile species (e.g., CO_2 , CH_4 and N_2) in individual fluid inclusions that are representative of particular populations were analyzed by laser Raman spectroscopy, performed on a Renishaw RM 2000 Raman spectrometer at Nanjing University,

China. Molar fractions of CO_2 , CH₄ and N₂ were determined using the method of Burke (2001).

Salinities expressed as wt.% NaCl equivalent were calculated from the final melting temperature of ice in aqueous fluid inclusions according to the method of Bodnar (1993). The salinity of the $CO₂$ rich inclusions was calculated from the melting temperature of clathrate, using the computer programs of Bakker (1997) based on Diamond (1992). Bulk composition and molar volume were computed based on the P-T-V-X properties of individual inclusions in the C-H-O-N system (Thiéry et al., 1994; Bakker, 1997) using computer programs developed for fluid inclusion studies (Bakker and Brown, 2003). The P-T properties were modelled using the V-X data and the equation of state from Bowers and Helgeson (1983) as revised by Bakker (1999) for aqueous-carbonic fluid inclusions. For carbonic-rich inclusions, the equation of state from Duan et al. (1992, 1996) for C-H-O-N-S system was applied.

The compositions of fluid inclusions from selected assemblages were analyzed quantitatively by laser-ablation ICP-MS at ETH Zürich, Switzerland. Individual fluid inclusions were opened with a 193 nm Excimer laser (Compex 110 I, ArF, Lambda Physics), and the laser-ablation aerosol was transferred with helium carrier gas into the ICP-MS (ELAN 6100 DRC; Günther et al., 1998; Günther and Heinrich, 1999; Heinrich et al., 2003). ICP-MS conditions were similar to those reported in Pettke et al. (2004). Boron, Na, Mg, K, Mn, Fe, Cu, Pb, Zn, As, Sb, Au, Ag, Sr, Cs, and U concentrations and their detection limits were measured in each inclusion (App.), using SRM 610 from NIST as external standard and data reduction following Heinrich et al. (2003).

5. Fluid Inclusion Types and Petrographic Relations

Fluid inclusions observed in this study have negative crystal, elongate or irregular shapes (Fig. 6). Fluid inclusions types were classified based on their appearance at 25°C and by their Raman spectra, and occur in successive stages of the vein and alteration paragenesis, as summarized in Figure 5.

Type Ia inclusions are two-phase, liquid-rich aqueous inclusions with 10 to 20 vol.% of a lowdensity vapor bubble at room temperature. They occur in early barren milky quartz veins at Shuiyindong and Yata. Primary inclusions of this type occur along growth zones of quartz and have negative crystal shapes, generally less than 25 μm in diameter (Fig. 6B, E). Secondary inclusions are elongate or irregular and occur along trails crosscutting quartz grains or grain boundaries (Fig. 6C, F). Raman spectroscopy analysis has failed to accurately determine the composition of the vapor phase of this type of inclusion because the bubble moved as the laser beam was focused on it. Raman peaks of $CO₂$, N₂ and CH₄ have, however, been detected.

Type Ib inclusions are two- or three-phase aqueous-carbonic inclusions with a dominant aqueous liquid phase and a relatively constant carbonic (vapor $+$ CO₂ liquid) fraction of 15 vol.% (Fig. 6H). They are commonly observed in quartz veinlets with arsenian pyrite and arsenopyrite of the main stage of gold mineralization at Yata (Fig. 3J) and in jasperoidal quartz of the main stage at Shuiyindong. Primary inclusions are typically 20 μm in diameter, occur along growth zones of quartz, and have negative crystal shapes (Fig. 6G, H). Some inclusions along microfracture planes within quartz grains are pseudosecondary based on their spatial relationship to the growth zones and healed fractures. Both microthermometry and Raman spectroscopy analyses have revealed that the main component of the volatile phase of the inclusions is $CO₂$, with minor N₂ and trace CH₄.

Type II inclusions are rare, two-phase, aqueous-carbonic inclusions with variably high proportions of a carbonic phase ranging from 45 to 90 vol.% (Fig. 6K). In samples from Yata, they occur in late drusy quartz with realgar, stibnite and calcite (Fig. 3B, K).

Figure 6. Microphotographs of fluid inclusions in quartz. A. Early milky quartz (thin section) from Shuiyindong. B. Primary and C. secondary type Ia aqueous inclusions in the milky quartz from Shuiyindong. D. Photograph and sketch of early milky quartz from Yata. E. Primary and F. secondary type Ia aqueous inclusions in the milky quartz from Yata. G. SEM-CL image of a main ore-stage quartz crystal, and H. Photographs of Type Ib aqueous-carbonic inclusions from Yata. Circles denote the positions of inclusions analyzed by LA-ICP-MS. I. to L. Late drusy quartz crystal with type Ib, type II and type III aqueouscarbonic and carbonic-rich inclusions from Yata, respectively. Note that two end-member fluid inclusion assemblages are cogenetic in (L). For the various fluid inclusion types refer to Table 1. Scale bar is 20 μm unless defined otherwise.

Type III inclusions are monophase carbonic inclusions (Fig. 6L) and generally less than 15 μm in

diameter. They are restricted to late quartz-realgar veins or veinlets at Yata and occur along trails crosscutting quartz grains. Both microthermometry and Raman spectroscopy have revealed that the volatile phase is mainly composed of $CO₂$ and $N₂$, with trace CH₄.

Petrographic relationships indicate that $CO₂$ -poor aqueous inclusions of Type Ia can be interpreted to approximate the mineralizing fluid, which was responsible for precipitation of early ore minerals in veins but predates the deposition of the bulk of disseminated gold-bearing arsenian pyrite and arsenopyrite deposition in both deposits. CO₂-rich fluids of Type Ib are interpreted to correspond to the main gold-bearing fluid in both deposits. The same aqueous-carbonic fluids also occur together with Type II and III carbonic fluids in the late quartz-realgar veins at Yata. Type II and III fluids are interpreted to represent the waning or outflow stage of economic gold mineralization.

6. Quantitative Fluid Inclusion Results

6.1. Microthermometry and Raman data

Microthermometric measurements were made on primary, pseudosecondary and secondary inclusions in quartz crystals from milky quartz veins, replacement-style quartz veinlets and late drusy quartz. Three distinctive fluid compositions are interpreted to be representative of the different mineralization stages. Microthermometric data are summarized in Table 1 and Raman spectroscopy analyses of representative fluid inclusions are given in Table 2.

Type Ia primary and secondary aqueous inclusions (Fig. 6B-C, E-F) have initial ice-melting temperatures (Te) from –22.2 to –21.0°C, which is similar to the eutectic melting temperature in the NaCl-H2O system (Hall et al., 1988) but does not exclude more complex systems such as the NaCl-KCl-H2O system (Sterner et al., 1988). Final ice-melting temperatures (Tm) of primary aqueous inclusions range from –3.0 to –4.3°C, corresponding to salinities of 5.0 to 6.9 wt.% NaCl equivalent (Bodnar, 1993) with an average of 6.0 wt.% NaCl equivalent (Fig. 7B, D). Homogenization of these fluid inclusions was to the liquid phase at temperatures ranging from 190 to 258°C with a mode around 230°C (Fig. 7A, C). The Tm of secondary inclusions ranges from –1.2 to –4.5°C, corresponding to salinities of 2.1 to 7.2 wt.% NaCl equivalent (Fig. 7B, D). Secondary inclusions of type Ia also homogenized into the liquid phase at temperatures ranging from 151 to 261°C with a mode around 190 \degree C (Fig. 7C). No evidence of other phases, such as clathrate, liquid or solid CO₂ were observed, suggesting that this type of inclusion may contain as much as 2.4 mole % CO_2 dissolved in the aqueous phase without developing a separate $CO₂$ liquid phase at room temperature (Bodnar et al., 1985).

Type Ib primary and secondary aqueous-carbonic inclusions (Fig. 6H, J) always develop a vapor phase in the carbonic bubble, even if they are two-phase at room temperature. The melting temperature of CO_2 (Tm CO_2) ranges from -58.1 to -56.6°C with the majority of CO_2 melting occurring at -56.6° C (Fig. 7G). The carbonic phase always homogenized to the liquid (ThCO₂) at temperatures ranging from 10.2 to 26.1°C with a mode around 24.0°C (Fig. 7H). Clathrate observed in these inclusions exhibits a typical Q2 melting behavior (Bakker and Brown, 2003). The clathrate melting temperatures (Tm cl) of primary inclusions range from 8.3 to 9.2°C, corresponding to salinities of 1.6 to 3.3 wt.% NaCl equivalent (Diamond, 1992), with an average of 2.3 wt.% NaCl equivalent (Fig. 7F), which is two to three times lower than the salinity of the type Ia aqueous inclusions. The Tm cl of secondary inclusions ranges from 9.4 to 9.8°C, corresponding to salinities of 0.4 to 1.2 wt.% NaCl equivalent (Fig. 7F). These inclusions commonly decrepitated at temperatures below 200°C, before total homogenization was attained. The 35 inclusions that did not decrepitate homogenized into the liquid phase at temperatures from 190 to 245°C, with a mode around 220°C (Fig. 7E). Raman

Deposit	Stage	Fluid inclusion assemblage			Microthermometric data								
		Generation	Type	Number	TmCO ₂	Theo ₂	T _m	Tm cl	Th				
Shuiyindong	Early-stage	Primary in vein quartz crystals	Type Ia	14			-3.5 to -4.3		218 to 231				
	Vein quartz-pyrite	Secondary in vein quartz crystals	Type Ia	18			-2.3 to -3.3		194 to 229				
Yata	Early-stage	Primary in vein quartz crystals	Type Ia	23			-3.0 to -4.1		190 to 258				
	Vein quartz-pyrite												
	Main-stage	in veinlet quartz Secondary	Type Ia 22				-2.1 to -3.3		165 to 230				
	Quartz-arsenian	crystals							(190)				
	pyrite-arsenopyrite	Primary and secondary in veinlet Type Ib		67	-58.1 to -56.6	10.2 to 26.1		8.3 to 9.8	190 to 245				
		quartz crystals			(-56.6)	(24) L,		(9.5)	(220)				
	Late-stage	Secondary in veinlet quartz	Type Ia	23			-1.2 to -4.5		151 to 261				
	Quartz-stibnite-	crystals					(-3.5)		(190)				
	realgar -orpiment		Type II	6	-59.6 to -58.1	6.3 to 20.9 L,		9.5 to 10.7	205^{d} to 232^{d}				
			Type III	25	-60.5 to -59.6 (-60.1)	-24.3 to -22.5 (-24.0) L,							

TABLE. 1. Summary of fluid inclusion types associated mineral assemblages and microthermometric data

For microthermometric parameters, ranges (in parenthesis) and mode are given. Tm CO₂ = final melting temperature of CO₂, ThCO₂ = homogenization temperature of CO₂, Tm = melting temperature of ice, Tm cl = final melting temperature of clathrate, Th = homogenization temperature. L = homogenize to liquid. d = Decrepitation temperature. All values are in °C

Stage	Quartz type	Inclusion	Microthermometry				Raman data				
			$T_m co_2$	Thco ₂	Mode	Tm cl	Th	Mode	CO ₂	N_2	$\rm CH_{4}$
Main-stage	Veinlet quartz	41	-56.7	20.5	L	9.4	228	L	97.5	1.5	1.0
	crystals	42	-57.7	17.6	L	9.3	227	L	95.7	3.3	1.0
		55	-56.7	21.6	L	9.4	220	L	97.0	3.0	n.d.
		60	-56.8	22.2	L	8.8	215	L	96.8	2.0	1.2
		61	-56.6	22.9	L	9.4	210	L	99.5	0.5	n.d.
		65	-56.9	21.4	L		238	L	96.0	3.5	0.5
		66	-56.8	22.3	L		217	L	97.0	3.0	n.d.
		76	-56.7	23.4	L	9.6	223	L	96.8	2.5	0.7
		80	-56.7	23.3	L		206	L	98.2	1.8	n.d.
		93	-56.6	24.8	L	8.6	225	L	98.7	1.3	n.d.
		94	-56.7	24.5	L	9.4	210	L	97.5	1.8	0.7
		99	-56.7	17.9	L	9.0	247	L	96.5	2.7	0.8
		107	-56.7	20.8	L	9.8	216	L	97.2	1.9	0.9
Late-stage	Veinlet quartz	34	-58.7	6.3	L	9.5	232 ^d		86.5	13.5	n.d.
	crystals	35	-58.6	7.8	$\mathbf L$	9.3	228 ^d		89.2	10.0	0.8
		11	-60.5	-24.3	L				71.0	27.2	1.8
		12	-60.3	-24.0	L				73.0	27.0	n.d.
		14	-60.2	-23.3	L				75.0	25.0	n.d.
		15	-60.2	-23.2	L				77.0	23.0	n.d.

TABLE. 2. Chemical compositions of representive aqueous-carbonic and carbonic-rich inclusions obtained by Raman spectroscopic analysis and corresponding microthermometric data from Yata

n.d. = not detected, other abbreviations as in Table 1. Composition and temperature are given in mol. % and °C, respectively.

spectroscopy of the carbonic phase in individual fluid inclusions showed that $CO₂$ is the dominant volatile (> 96 mole %), N_2 is always detected (0.5 to 3.5 mole %), and CH₄ has been detected (up to 1.2 mole %) in a few inclusions (Table 2).

Figure 7. Histogram of homogenization temperature, final CO₂ melting temperature and salinity of fluid inclusions from Shuiyindong (**A-B**) and Yata (**C-H**).

Type II aqueous-carbonic inclusions (Fig. 6K) also always develop a vapor phase in the carbonic bubble during cooling runs. Their Tm $CO₂$ ranges from -59.6 to -58.1°C (Fig. 7G). Homogenization of the CO² was always to the liquid phase between 6.3 to 20.9°C. The Tm cl ranges from 9.5 to 10.7°C, corresponding to salinities of 0 to 8.9 wt.% NaCl equivalent (Bakker and Brown, 2003). Total homogenization temperatures were not obtained because these inclusions decrepitated when heating to above 200 $^{\circ}$ C. Raman spectroscopy revealed that their volatile phases contain major CO₂ (87-89 mole %), minor N_2 (10-14 mole %) and trace CH₄ (0.8 mole %; Table 2).

In the process of freezing (down to -180°C) and subsequent heating, Type III carbonic inclusions (Fig. 6L) underwent the following sequence of phase transitions: $S + V \rightarrow L + V \rightarrow L$. The Tm CO₂ ranges from –60.5 to –59.6°C with the majority at –60.1°C (Fig. 7G). The ThCO₂ of this type of inclusion ranges from –24.3 to –22.5°C (Fig. 7H) and was always into the liquid phase. Total homogenization temperatures could not be measured reliably owing to optical limitations (Diamond, 2003). Raman spectroscopy showed that the volatile phase of type III inclusions contains major CO_2 (71-77 mole %), minor N_2 (23-27 mole %) and trace CH₄ (up to 1.8 mole %; Table 2).

6.2. Bulk compositions of aqueous-carbonic and carbonic-rich fluids

Aqueous-carbonic inclusions in the main stage of pyrite-arsenopyrite-quartz veins and late stage of stibnite-realgar-orpiment-quartz veins at Yata have 10-15 vol.% volatile phase (nearly pure CO2), regardless of the sample localities. Calculated density of the volatile phase varies from 0.72 to 0.82 $g/cm³$, with an average of 0.75 g/cm³. Calculated bulk compositions showed that the fluids are dominated by H₂O (91 to 92 mole %) and CO₂ (6.3 to 8.4 mole %, with an average of 7.6 mole %). Their bulk densities range from 0.97 g/cm³ to 0.99 g/cm³. Their average salinities vary from 0.9 to 2.3 wt.% NaCl equivalent.

Calculated bulk compositions of carbonic-rich inclusions in the late stage of stibnite-realgarorpiment-quartz veins at Yata are dominantly CO_2 (58 to 64 mole %), N₂ (19.2 to 23.7 mole %) and H₂O (12.5 to 22.8 mole %), and trace CH₄ (up to 1.6 mole %), with volatile phase densities and bulk densities of 0.75 to 0.80 g/cm^3 and 0.77 to 0.82 g/cm^3 , respectively.

7. Pressure and Temperature Estimation

The compositions of type Ib inclusions plot close to the solvus of the H_2O-CO_2 system at 1.0 kb in Figure 8A. It is possible that the fluids of these inclusions may represent one end-member produced by unmixing of H_2O -CO₂ fluids. The compositions of a few type II inclusions plot in the two-phase field in Figure 8A. These inclusions may represent mechanical mixtures of two immiscible fluid phases. The local occurrence of type II inclusions at Yata indicates that the hydrothermal fluid reached a state of immiscibility at some time in its history (Diamond, 1990). Type III inclusions plot close to the 1.0 kb solvus in Figure 8A. If they coexisted with type Ib inclusions, then these inclusions may represent another end-member produced by unmixing of H_2O -CO₂ fluids.

The minimum P-T conditions of inclusion entrapment are constrained by intersecting points using the range of homogenization temperatures of type Ib inclusions (190 to 245°C) and the minimum of the bubble curves of 6 mole $\%$ CO₂ (Fig. 8B). The defined area in Figure 8B (shaded) ranges from 450 to 1150 bars, corresponding to a depth of 1.7 to 4.3 km under lithostatic load, using the average density of sedimentary rocks in southwestern Guizhou (2.67 g/cm3: Wang et al., 1995) and 4.5 to 11.5 km assuming hydrostatic pressure. As Yata was controlled by a fault, the estimated pressure may have been fluctuating between hydrostatic and lithostatic pressures. Decompression associated with episodes of faulting may have caused the immiscibility in the late stage of stibnite-realgar observed

in this deposit. Zhang et al. (2003) previously estimated pressures for the Lannigou deposit ranging from 600 to 1700 bars based on the $CO₂$ -bearing fluid inclusions, corresponding to a depth of 2.2 to 6.3 km under lithostatic conditions, or an unlikely 6 to 17 km under hydrostatic load. $CO₂$ contents of fluid inclusions at Yata (6 to 8 mole %) are somewhat lower than those of many orogenic lode gold deposits (10 to 25 mole % CO2: Ridley and Diamond, 2000) but higher than those of Carlin-type gold deposits in Nevada (2 to 4 mole % CO2: Hofstra and Cline, 2000). It is, therefore, reasonable to infer that the Yata deposit may have formed at depths intermediate between orogenic-type gold deposits and those of the Carlin trend in Nevada.

Figure 8. A. T-X diagram of the H₂O-CO₂ system at 1.0 kbar based on experimental data from Tödheide and Franck (1963) and Takenouchi and Kennedy (1964). **B.** P-T diagram showing the range of possible P-T conditions during mineralization at Yata. The miscibility boundaries for fluids with 6 and 7.6 mole % CO₂ are from experimental data of the H₂O-CO₂ system of Tödheide and Franck (1963) and Takenouchi and Kennedy (1964).

8. LA-ICP-MS Microanalyses of Fluid Inclusions

Representative fluid inclusion types from both deposits were analyzed by LA-ICP-MS. Boron, Na, Mg, K, Mn, Fe, Cu, Pb, Zn, As, Sb, Au, Ag, Sr, Cs and U concentrations were recorded in all analyses. Fluid inclusions of all types consistently contain detectable quantities of B, Na, K, Sr, Cs, As, and Sb, whereas Mg, Mn, Fe, Ag, Cu, Pb, Zn, U and Au are commonly near or below the limit of detection (LOD). Gold was above the $1\mathbb{Z}$ limit of detection in 5 aqueous inclusions and in 12 aqueous-carbonic inclusions. The element concentrations in each inclusion are listed in Appendix. The average element concentrations in the different fluid inclusion types are given in Table 3.

The results show systematic variations in metal concentrations of fluids between the different fluid inclusion types (Table 3 and Fig. 9). The B, Na, K, Cs, Sr, As, Sb and Au concentrations are higher in aqueous inclusions (Type Ia). Secondary aqueous inclusions have lower concentrations of these elements than primary aqueous inclusions. Concentrations of these elements are progressively lower in the aqueous-carbonic inclusions, but they do not decrease at the same rate. The Sr decrease is steeper than that of the other alkali metals (Fig.9), indicating that selective precipitation of Sr-bearing

minerals occurred during mineralization. Gold concentrations in the aqueous inclusions from Shuiyindong and Yata range from 3.8 ± 0.5 to 5.7 ± 2.3 μ g/g (Table 3), whereas the aqueous-carbonic inclusions from Yata have lower concentrations of Au, ranging from 0.1 to 0.8 μg/g (App., sample YT-5). Arsenic concentrations are approximately constant in the aqueous inclusions at Shuiyingdong, but decrease from 200 \pm 120 μg/g in the aqueous inclusions to 100 \pm 50 μg/g in the aqueous-carbonic inclusions at Yata. Antimony concentrations in aqueous-carbonic inclusions in the late stibniterealgar-quartz veins at Yata are higher (up to 90 μ g/g) than in the other inclusions. A decrease in Sb and As concentrations is probably due to precipitation of stibnite and realgar, which have been observed within late-stage quartz growth zones. However, other cations, such as Mg, are generally below the limit of detection in the fluid inclusions. Iron, which has a typical detection limit of 400 μg/g in LA-ICP-MS analysis, was not detected in any fluid inclusions analyzed during this study.

TABLE. 3. Summary of fluid inclusion composition (in µg/g) analyzed by laser ablation ICP-MS

b.d. = below detection. Abbreviations: $Qz =$ quartz, $py =$ pyrite, As-py = arsenian pyrite, Asp = arsenopyrite, cal = calcite, real = $realgar, stb = stibnite.$

^aNumbers in parenthesis are 1 standard deviation for an inclusion assemblage of which several inclusions have been analysed individually.

^b Numbers based on a single inclusion analysis with concentrations above the detection limit.

 \textdegree Numbers based on the average of multiple inclusion analyses with concentrations above the detection limit (38).

Figure 9. Chemical evolution of fluid inclusion assemblages across the paragenetic sequences based on LA-ICP-MS analysis. The relative timing of fluid inclusion assemblages is based on crosscutting relationships and quartz growth zones revealed by cathodoluminescence. Note the general decrease in element concentrations, except for As and to a lesser extent Sb, across the ore depositional event. The parallel and vertical grey lines represent the limit of detection of B, Na, K, Cs, As, Sb, Sr at 3σ and Au at 1σ respectively.

9. Discussion

9.1. Interpretation of fluid inclusion paragenesis and LA-ICP-MS data

LA-ICP-MS results showed that aqueous inclusions in barren milky quartz veins from both deposits have high concentrations of the characteristic ore elements (As, Sb and Au) compared to aqueouscarbonic inclusions. This fluid is interpreted to approximate the mineralizing fluid, which was responsible for precipitation of early milky quartz veins. Aqueous-carbonic fluid is interpreted to correspond to the main-stage gold-bearing fluid. The higher $CO₂$ content of this fluid is interpreted to result from carbonate dissolution at the level of the deposits. The decease of Au and As in aqueouscarbonic inclusions is interpreted to be due to deposition of disseminated gold-bearing arsenian pyrite and arsenopyrite in the main mineralization stage. Covariance of Au and As, as well as Sb and Sr, in ore fluids (Figs. 9, 10A), as observed in EMPA analyses of gold-bearing arsenian pyrite and bulk ore samples (Fig. 10B, C), is interpreted to reflect precipitation of As and Sb sulfides and gold concurrently with Sr-bearing carbonate. This is consistent with petrographic observations that goldbearing arsenian pyrite and arsenopyrite are accompanied in the main mineralization stage by the Fe-poor hydrothermal dolomite or calcite in both deposits (Fig. 3C, I). Dilution also may have occurred during gold mineralization because the salinities of the inclusions decrease two to three fold from aqueous inclusions to aqueous-carbonic inclusions.

Figure 10. Correlation of Au and As, Sb, Sr in ore fluid (A), gold-bearing arsenian pyrite (B), and ore (C). The EMPA data of arsenian pyrite and bulk ore samples are from Su et al. (2008) and Xia (2005), respectively.

9.2. Source of fluids

Oxygen and hydrogen isotopic compositions of early barren milky quartz veins and clay minerals (<2 μm) in the mineralized rock from Shuiyindong have been reported by Hofstra et al. (2005). The milky quartz veins have a narrow range of high δ^{18} O values (21-27 ‰). Calculated δ^{18} OH₂O values for

fluids in these samples vary from 10.5 to 16.5 ‰, using a homogenization temperature of 230°C from this study and the quartz-water isotopic equilibrium fractionation equation of Friedman and O'Neil (1977). Calculated $\delta^{18}OH_2O$ and δDH_2O values for fluids in equilibrium with clay minerals in the mineralized rocks at Shuiyindong range from 4 to 10 ‰ and -56 to -68 ‰, respectively (Hofstra et al., 2005). Similar fluid calculations for clay minerals in the mineralized rocks from Zimudang deposit, located 15 km west of Shuiyindong on the same large-scale dome structure, indicate $\delta^{18}OH_2O$ and δ DH₂O values from 13 to 16 ‰ and -35 to -40 ‰, respectively (Hofstra et al., 2005).

Li et al. (1989) and Zhu et al. (unpub. data) reported oxygen for the main-ore quartz and the lateore calcite from Yata. The main-ore quartz has a narrow range of δ^{18} O values (20.9-26.1 ‰), similar to milky quartz veins at Shuiyindong. The calculated $\delta^{18}OH_2O$ values for the main-ore fluids from Yata vary from 9.9 to 15.1 ‰ at 220°C. The $\delta^{18}OH_2O$ values of late-ore fluids (8.1 to 10.9 ‰) were calculated from measured δ^{18} O calcite values (18.2-20.8 ‰), the minimum homogenization temperature of 190 \degree C for CO₂-bearing fluid inclusions within calcite (Zhang et al., 2003), and the calcite-water isotopic equilibrium fractionation equation of Friedman and O'Neil (1977). The δDH2O values of water extracted from inclusion fluids in the main-ore quartz at Yata range from –51.1 to – 78.8 ‰ (Li et al., 1989), significantly higher than the average δDH2O value of local meteoric water during Jurassic and Cretaceous time (-85 ‰: Han et al., 1999).

The δ^{18} OH₂O and δ DH₂O values plot within or close to the metamorphic-water box (Hofstra et al., 2005, fig. 3), suggesting that the ore fluids are characterized by a predominantly metamorphic component. There is no evidence for meteoric water involvement, but a deep magmatic component cannot be excluded.

9.3. Processes of gold deposition

Recent studies of Carlin-type deposits in Nevada have concluded that gold-rich arsenian pyrite precipitated from H2S-rich fluids, which sulfidized iron-bearing minerals in the host rocks (Hofstra et al., 1991; Hofstra and Cline, 2000; Kesler et al., 2003). The chemical reaction of cooling H₂S-rich fluids in contact with iron-poor carbonate-bearing sedimentary rocks has been calculated from thermodynamic data by Heinrich (2005). An important prediction is that Au solubility may remain high to temperatures as low as 150°C, provided that an excess of H_2S over Fe is maintained in a moderately oxidized hydrothermal fluid, the pH of which is kept close to neutral by carbonate dissolution (Heinrich, 2005, fig. 6). This fluid-chemical evolution path can equally apply to fluids of magmatic or metamorphic origin, but clearly matches the low Fe contents of fluids measured in the inclusions from the Shuiyindong and Yata deposits.

At the stratabound Shuiyindong deposit, there is no evidence for significant phase separation during the main mineralization stage. At the fault-controlled Yata deposit, phase separation was minor during gold mineralization and common in the late stibnite-realgar stage. This fluid inclusion evidence suggests that phase separation was not the key process for deposition of gold and arsenian pyrite. The low Fe contents in the ore fluids (below 400 μg/g) measured by LA-ICP-MS of fluid inclusions and many relict inclusions of ferroan carbonate (with up to 7.0 wt.% Fe determined by EMPA: Su et al., 2008) enclosed in the jasperoidal quartz crystals suggest that iron in sulfide minerals was probably derived from dissolution of ferroan carbonate in the host rocks, as has been documented Carlin-type gold deposits in Nevada by lithogeochemistry of ores (Hofstra et al., 1991; Hofstra and Cline, 2002). Sulfidation of ferroan carbonate-rich host rocks by H₂S-rich ore fluids containing Au(HS)₂⁻ or Au(HS)⁰ (Seward, 1973, 1991), along with arsenic as $H_3AsO_{3(aq)}$ complex (Heinrich and Eadington, 1986; Pokrovski et al., 2002), would have effectively extracted gold from solution and transformed primary ferroan carbonate to secondary gold-bearing arsenian pyrite, possibly by a coupled reaction such as the following:

$$
2FeCO3 + 2H3AsO3 (aq) + H2S(aq) + 2Au(HS)2 (aq) + H2 (g)
$$

= Fe(S, As)₂ · Au₂S⁰ + FeS₂ + CO₃⁻² (aq) + CO₂ (g) + 7H₂O

Any loss of $CO₂$ by fluid phase separation would further promote the formation of gold-bearing arsenian pyrite by such a reaction. This is consistent with petrographic observations that gold-bearing arsenian pyrite and arsenopyrite occur together with Fe-poor hydrothermal dolomite (Fig. 3C). Moreover, it can explain the covariance of Au, As, Sb and Sr contents of the ore fluids (Fig. 9). The reaction involves reduction of As (+III) and Au (+I) by consumption of $H_{2(g)}$ which can be provided by reaction of the metal-transporting fluids with the local organic-rich sediments. This situation is likely to occur where fluids are channeled into a fluid conduit (e.g., along the unconformity between the Maokou Limestone and the Longtan Formation at Shuiyindong or the reverse faults at Yata). Here, they reacted with carbonaceous and Fe-carbonate-rich host rock, or were mixed with Fe⁺²- and CH₄bearing pore fluids previously equilibrated with the reduced host rock package, to deposit arsenian pyrite and arsenopyrite that host most of the gold in the deposits. The generation of secondary pervasive permeability and porosity by carbonate dissolution from the host rocks favors fluid focussing and the formation of stratabound orebodies (Heinrich, 2005), as is the case in the Shuiyindong. Therefore, fluid reduction and sulfidation of wall-rock iron by H_2S -rich ore fluids are proposed to be the most important mechanism of gold deposition as auriferous arsenian pyrite in the Chinese Carlin-type gold deposits.

Genetic models previously proposed for Carlin-type deposits in Nevada fall into three groups involving magmatic (Radtke et al., 1980; Ressel et al., 2000), metamorphic (Groves et al., 1998; Cline and Hofstra, 2000), and deeply or shallowly circulated meteoric waters (Ilchik and Barton, 1997; Emsbo et al., 2003). Our study of Carlin-type deposits in Guizhou indicates that the ore fluids are of low salinity but rich in high-density $CO₂$, and stable isotope data are consistent with a dominant metamorphic fluid source. The ore-forming fluids have extremely high contents of the characteristic ore elements (As, Sb and Au), and the mineralogical characteristics and high gold grade of the deposits can be explained by interaction of such fluids with chemically reactive carbonate-, iron- and carbonrich sedimentary host rocks. The deposits formed at similar temperatures as epithermal gold deposits, but at significantly higher pressure and greater depths (4-6 km), consistent with regionalmetamorphic temperature gradients. Their thermal regime and ore fluid characteristics are similar to those of the broad group of orogenic gold deposits, raising the possibility that the Carlin-type deposits in Guizhou might be the basin-hosted and relatively cool end-member of the 'crustal continuum' of orogenic gold deposit formed from fluids liberated by deep metamorphic dehydration or magmatism (Groves, 1993; Groves et al., 1998; Pettke et al., 2000).

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Appendix

Appendix 1 Element concentrations in aqueous inclusions in the early milky quartz veins at Shuiyindong and Yata

Sample	No. of	Salinity	B	Na	Mg	K	Mn	Fe	Cu	Zn	As	Sr	Ag	Sb	Cs	Au (3σ)	Au (1σ)	Pb	U
Number	inclusion	wt% NaCl eq	μ g/g	μ g/g	μ g/g	μ g/g													
SYD035-O2	mc02b03	6.9	1060	26800	$<$ 40	620	$<$ 20	$<$ 320	$<$ 20	$<$ 30	30	220	$<$ 2	$<$ 5	10	<1.7	< 0.6	$<$ 2	
	mc02b04	6.9	1040	26800	< 60	620	$<$ 30	$<$ 440	$<$ 20	50	$<$ 40	220	$<$ 3	$<$ 7	30	< 2.5	< 0.8	5	
	mc02b05	6.7	1230	26200	< 160	810	< 110	< 1330	50	< 140	110	280	20	$<$ 30	10	< 6.4	3.3	$<$ 10	$<$ 1
	mc02b07	6.6	1110	25700	< 130	680	< 100	< 1170	$<$ 40	< 110	90	180	$<$ 10	20	9	<6.6	4.4	$<$ 10	\leq 2
	mc02b09	6.9	820	26700	60	940	<70	< 720	$<$ 40	$<$ 50	< 80	250	$<$ 7	<10	20	<3.9	<1.3	$<$ 7	\leq 1
	mc02b10	6.7	1110	26300	< 80	690	<70	< 1200	$<$ 20	< 80	80	220	$<$ 5	<10	10	<3.1	<1.1	$<$ 9	$<$ 1
SYD035-Q1	mc02a08	5.4	790	21000	260	410	$<$ 40	< 540	30	$<$ 50	$<$ 40	160	5	$<$ 7	10	< 2.3	< 0.8	≤ 4	$<$ 1
	mc02a09	5.3	720	20300	150	930	< 100	<1610	150	< 180	< 120	180	20	$<$ 20	20	< 6.8	< 2.3	$<$ 10	\leq
	mc02a10	5.1	1060	20200	90	420	<70	< 1010	$<$ 50	< 100	<70	170	$<$ 6	$<$ 20	20	$<$ 4.8	< 1.6	$<$ 10	\leq 2
	mc02a11	5.3	390	20500	<70	290	$<$ 40	$<$ 370	$<$ 30	$<$ 40	90	140	\leq 3	$<$ 9	7	< 1.6	< 0.6	$<$ 5	$<$ 1
	mc02a12	5.3	870	20600	$<$ 40	150	$<$ 30	$<$ 240	$<$ 20	$<$ 30	110	170	$<$ 2	$<$ 6	10	< 1.2	< 0.4	$<$ 3	$<$ 1
	mc02a13	5.3	970	20600	90	270	< 60	< 820	$<$ 30	90	<70	120	< 8	<10	7	< 5.4	< 1.8	< 8	$<$ 1
	mc02a14	5.3	530	20700	≤ 40	< 140	$<$ 50	< 630	$<$ 40	<70	< 60	50	$<$ 6	<10	5	<4.2	< 1.4	$<$ 6	$<$ 3
	mc02a15	5.3	640	20600	< 100	210	70	< 840	$<$ 30	<70	<70	170	$<$ 6	<10	20	<3.8	< 1.3	< 6	\leq 1
	mc02a17	5.3	900	20600	< 60	300	$<$ 40	$<$ 430	50	< 60	70	140	$<$ 2	$<$ 9	8	< 1.8	< 0.6	$<$ 5	$<$ 1
O64	mc02c03	5.6	360	21500	$<$ 220	410	< 120	< 1360	70	200	200	30	30	$<$ 30	10	6.0	6.0	$<$ 20	$\overline{4}$
	mc02c05	5.7	500	21600	< 120	820	< 130	< 1480	< 110	< 150	< 150	30	$<$ 20	$<$ 40	5	< 8.7	< 2.9	$<$ 20	\leq
	mc02c06	5.7	1260	21800	$<$ 260	$<$ 490	$<$ 200	$<$ 2530	100	$<$ 280	340	50	$<$ 20	$<$ 50	10	<14.3	7.8	40	\leq 3
	mc02c07	5.6	540	21700	$<$ 40	340	$<$ 30	< 410	<10	$<$ 40	90	20	9	20	20	3.3	3.3	20	8

< values show the limit of detection (LOD) at 3 sigma. Detection limit of elements (38): $B \sim 80$, $Na \sim 10$, $Mg \sim 30$, $K \sim 80$, $Mn \sim 20$, $Fe \sim 400$, $Cu \sim 40$, $Zn \sim 30$, $As \sim 20$, $Sr \sim 1$, $Ag \sim 2$, $Sb \sim 5$, $Cs \sim 1$, A

 \sim 1, Pb \sim 4 and U \sim 1 µg/g, respectively.

Sample	No. of	Salinity	B	Na	Mg	K	Mn	Fe	Cu	Zn	As	Sr	Ag	Sb	Cs	Au (3σ)	Au (1σ)	Pb	U
Number	inclusion	wt.%NaCl eq.	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g									
$YT-5$	mc03a06	2.0	250	7700	$<$ 20	380	$<$ 20	$<$ 260	$<$ 10	$<$ 40	170		$<$ 2	$<$ 5	3	<1.7	< 0.6	$<$ 2	$<$ 1
	mc03a07	2.0	130	7800	$<$ 10	180	$<$ 10	<70	$<$ 10	$<$ 10	120	40	$<$ 1	$<$ 1	\overline{A}	< 0.5	< 0.2	$<$ 1	$<$ 1
	mc03a09	2.0	150	7800	10	140	$<$ 10	< 130	<10	$<$ 10	40	10	<1	\leq 2		0.6	0.6	<1	$<$ 1
	mc03a10	2.0	270	7800	<10	200	<10	$<$ 40	<10	<10	80	3	$<$ 1	5	5	< 0.2	0.2	1	$<$ 1
	mc03a11	2.0	570	7800	$<$ 10	310	$<$ 10	< 140	<10	$<$ 10	270	4	\leq 1	20	4	< 0.9	0.4	\leq 1	$<$ 1
	mc03a12	2.0	250	7900	$<$ 40	90	$<$ 30	$<$ 470	<10	< 60	200	1	$<$ 6	20	2	< 2.5	< 0.8	$<$ 5	$<$ 1
	mc03a13	2.0	350	7900	$<$ 10	50	$<$ 10	< 190	$<$ 10	$<$ 20	170	6	\leq 1	$<$ 3	5	< 0.9	< 0.3	$<$ 1	$<$ 1
	mc03a14	2.0	310	7800	$<$ 20	320	$<$ 10	< 150	$<$ 10	$<$ 20	150	3	\leq 1	$<$ 3	6	0.6	0.6	\leq 1	$<$ 1
	mc03a15	2.0	350	7700	20	350	$<$ 10	< 100	$<$ 10	$<$ 10	140	2		$<$ 3	5	< 0.6	0.3	\leq 1	$<$ 1
	mc03a16	2.0	290	6900	200	2460	<10	< 110	<10	$<$ 10	100	3	\leq 1	$<$ 2	10	< 0.6	0.2	$<$ 1	$<$ 1
	mc03a17	2.0	450	7800	$<$ 10	100	$<$ 10	< 100	$<$ 10	$<$ 10	140	2	$<$ 1	\leq 2	9	< 0.5	0.2	$<$ 1	$<$ 1
	mc03b03	2.0	330	7800	<70	140	$<$ 40	< 600	$<$ 30	$<$ 50	80	3	$<$ 6	<10	5	< 1.6	0.8	$<$ 5	$<$ 1
	mc03b04	2.0	210	7700	$<$ 30	340	$<$ 10	< 180	$<$ 10	$<$ 20	80	5	$<$ 1	$<$ 3	$\overline{4}$	< 0.6	0.3	\leq 2	$<$ 1
	mc03b06	2.0	330	7700	<10	490	<10	<70	<10	<10	80	3	<1	<1	7	< 0.3	< 0.1	<1	$<$ 1
	mc03b07	2.0	250	7700	20	$<$ 20	$<$ 10	< 120	$<$ 10	$<$ 10	70	2	$<$ 1	2	2	< 0.7	< 0.2	$<$ 1	$<$ 1
	mc03b08	2.0	260	7600	<10	610	$<$ 10	< 100	10	<10	90	2	\leq 1	2	3	< 0.2	0.1	$<$ 1	$<$ 1
	mc03b09	2.0	350	7800	$<$ 30	80	$<$ 20	$<$ 390	$<$ 40	$<$ 30	160	1	$<$ 2	≤ 4	4	< 1.3	< 0.5	≤ 4	$<$ 1
	mc03b11	2.0	270	7800	$<$ 20	160	<10	< 190	$<$ 10	$<$ 20	100	2	$<$ 1	$<$ 3	5	< 0.7	< 0.2	$<$ 2	$<$ 1
	mc03b12	2.0	290	7900	$<$ 10	30	$<$ 10	< 120	$<$ 10	$<$ 10	110	2	\leq 1	$<$ 2	\overline{A}	< 0.5	< 0.2	$<$ 1	$<$ l
	mc03b13	2.0	210	7700	20	550	$<$ 10	< 130	$<$ 10	$<$ 10	70	$\overline{2}$	$<$ 1	2	5	< 0.6	< 0.2	$<$ 1	$<$ 1
	mc03b14	2.0	260	7800	10	220	$<$ 10	$<$ 40	$<$ 10	10	70	2		$<$ 1	5	0.3	0.3	$<$ 1	$<$ 1
	mc03b15	2.0	400	7800	$<$ 10	50	$<$ 10	$<$ 40	2	<10	160	2	$<$ 1	2	7	< 0.2	< 0.1	$<$ 1	$<$ 1
	mc03b16	2.0	410	7700	<10	480	$<$ 10	< 60	<10	$<$ 10	170		$<$ 1	$<$ 1		< 0.2	< 0.1	\leq 1	$<$ 1

Element concentrations in aqueous-carbonic inclusions in the main-stage quartz at Yata

Appendix 1 (continued)

Sample	No. of	Salinity	B	Na	Mg	K	Mn	Fe	Cu	Zn	As	Sr	Ag	Sb	Cs	Au (3σ)	Au (1σ)	Pb	U
Number	inclusion	wt.% NaCl eq	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g	μ g/g									
YT1-012	mc02c14	1.2	190	4800	30	< 80	$<$ 30	$<$ 320	<10	$<$ 30	50	$\overline{4}$	≤ 4	<10	\leq 1	< 2.4	< 0.8	≤ 4	\leq 1
	mc02c11	1.0	420	3900	< 180	$<$ 450	< 190	$<$ 2380	200	< 220	480	10	16	140	$<$ 3	< 11.6	<3.9	27	$<$ 2
	mc02c12	0.8	$<$ 200	3200	< 120	290	< 130	< 1820	< 110	< 150	< 130	≤ 4	19	$<$ 30	2	< 5.6	<1.9	23	8
	mc02c13	0.8	$<$ 330	1900	< 190	3040	$<$ 200	< 1860	90	< 220	< 190	7	24	60	24	< 7.7	4.3	22	$<$ 5
	mc02c16	1.0	130	4000	<70	140	30	< 450	20	< 50	150	$\overline{\mathbf{c}}$	3	30		< 2.4	< 0.8	$<$ 3	
	mc02c17	1.2	220	4400	< 270	$<$ 300	< 110	< 1560	<70	< 130	220	$<$ 5	20	70	$<$ 3	< 5.6	<1.9	<15	$<$ 3
	mc02c15	0.8	200	3000	< 100	$<$ 270	< 110	< 1150	<70	< 150	420	≤ 4	< 6	160	1	< 5.3	< 1.8	<12	<1
	mc02c18	1.2	< 210	4100	< 240	990	90	< 1020	90	< 110	200	≤ 4	$<$ 7	20	3	<3.9	<1.3	< 9	$<$ 3
YT1-010	mc02e03	0.8	110	3200	$<$ 50	90	20	230	20	$<$ 30	$<$ 30	20	$<$ 5	<10	17	< 2.1	< 0.7	8	\leq 1
	mc02e04	0.8	80	3300	<10	40	<10	< 60	<10	<10	40	$\overline{4}$	\leq 1	$\overline{4}$		< 0.3	< 0.1	<1	
	mc02e05	0.8	80	3100	20	540	<10	< 130	<10	<10	460	8	$<$ 2	10	3	< 0.6	< 0.2	$<$ 2	
	mc02e07	0.8	160	3200	$<$ 40	280	30	$<$ 360	30	$<$ 30	80	4	\leq 3	10	3	< 1.0	< 0.3	$<$ 3	<1
YT1-013	mc02e10	0.6	130	2400	$<$ 30	140	20	< 250	<10	20	20	$<$ 1	$<$ $\!\!2$	5	<1	< 0.9	< 0.3	$<$ 2	\leq 1
	mc02el1	0.6	70	2500	< 80	< 60	$<$ 30	< 410	20	$<$ 30	50	$<$ 2	2	10		<1.7	< 0.6	$<$ 3	\leq 1
	mc02el2	0.6	180	2400	$<$ 50	< 80	$<$ 30	$<$ 360	$<$ 30	$<$ 40	120	1	$<$ 7	10		< 1.2	< 0.4	$<$ 3	<1
	mc02e13	0.6	100	2500	< 60	< 110	$<$ 40	$<$ 480	20	< 60	$<$ 40	$<$ 3	$<$ 7	10	3	<3.3	<1.1	$<$ 5	$<$ 1
$YT-4$	mc02a04	11.1	$<$ 4870	41300	< 1950	< 5680	< 1990	< 28130	< 1420	$<$ 2950	< 2250	<70	$<$ 220	$<$ 400	20	<65	<22	< 240	30
	mc02a05	11.1	< 2180	43600	< 1200	$<$ 3080	< 960	< 16000	< 570	< 1390	1880	$<$ 40	< 140	340	20	< 57	<19	< 140	20
	mc02a06	11.1	< 7810	43600	< 6170	$<$ 11450	$<$ 3980	$<$ 55880	< 2850	$<$ 5460	$<$ 3950	< 130	< 240	< 950	< 80	< 600	$<$ 200	$<$ 480	$<$ 50
	mc02c08	11.1	1060	43400	< 190	< 550	< 210	$<$ 3000	< 130	$<$ 290	< 240	50	$<$ 30	$<$ 40	\leq 3	$<$ 9	$<$ 3	20	$\overline{4}$

Element concentrations in aqueous-carbonic and vapor-rich inclusions in the late-stage quartz at Yata

* : vapor-rich inclusions.

Appendix 1 (continued)