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The mixing of multi-source fluids in the Wusihe Zn–Pb ore deposit in Sichuan Province, Southwestern China

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Abstract The Sichuan-Yunnan-Guizhou (SYG) metallogenic province of southwest China is one of the most important Zn–Pb ore zones in China, with ~ 200 Mt Zn– Pb ores at mean grades of 10 wt.% Zn and 5 wt.% Pb. The source and mechanism of the regional Zn-Pb mineralization remain controversial despite many investigations that have been conducted. The Wusihe Zn-Pb deposit is a representative large-scale Zn-Pb deposit in the northern SYG, which mainly occurs in the Dengying Formation and yields Zn–Pb resources of ~ 3.7 Mt. In this paper, Zn and S isotopes, and Fe and Cd contents of sphalerite from the Wusihe deposit were investigated in an attempt to constrain the controls on Zn and S isotopic variations, the potential sources of ore-forming components, and the possible mineralization mechanisms. Both the δ^{66} Zn and δ^{34} S values in sphalerite from the Wusihe deposit increase systematically from the bottom to the top of the strata-bound orebodies. Such spatial evolution in δ^{66} Zn and δ^{34} S values of sphalerite can be attributed to isotopic Rayleigh fractionation during sphalerite precipitation with temperature variations. The strong correlations between the Zn-S

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isotopic compositions and Fe-Cd concentrations in sphalerite suggest that their variations were dominated by a similar mechanism. However, the Rayleigh fractionation mechanism cannot explain the spatial variations of Fe and Cd concentrations of sphalerite in this deposit. It is noted that the bottom and top sphalerites from the strata-bound orebodies document contrasting Zn and S isotopic compositions which correspond to the Zn and S isotopic characteristics of basement rocks and host rocks, respectively. Therefore, the mixing of two-source fluids with distinct Zn-S isotopic signatures was responsible for the spatial variations of Zn-S isotopic compositions of sphalerite from the Wusihe deposit. The fluids from basement rocks are characterized by relatively lighter Zn ($\sim 0.2 \%$) and S (~ 5 ‰) isotopic compositions while the fluids from host rocks are marked by relatively heavier Zn (~ 0.6 ‰) and S ($\sim 15 \%$) isotopic compositions.

Keywords Sichuan–Yunnan–Guizhou \cdot Wusihe Zn–Pb deposit \cdot Zn–S isotopes \cdot Fe–Cd contents \cdot Two-source fluids

1 Introduction

The Sichuan–Yunnan–Guizhou (SYG) metallogenic province is an important component of the giant South China low-temperature metallogenic domain (Hu et al. 2017). More than 400 carbonate-hosted Zn–Pb deposits occur in the SYG, yielding total Zn–Pb ores of more than 200 million tonnes at mean grades of 10 wt.% Zn and 5 wt.% Pb (Liu and Lin 1999; Zhou et al. 2014a, b). The sources for the regional Zn–Pb mineralization in the SYG remain controversial despite many investigations that have been conducted, mostly in regard to the Emeishan basalts, folded basements, and host rocks (Huang et al. 2003; Han et al. 2007; Ye et al. 2011; Zhou et al. 2014b, 2018; Zhu et al. 2016; Xiong et al. 2018). Most deposits in the SYG have been suggested to be comparable to Mississippi Valleytype (MVT), including many large-scale Zn-Pb deposits (e.g., the Huize, Daliangzi, Tianbaoshan, and Maozu deposits) (Han et al. 2007; Zhou et al. 2013; He et al. 2016; Wang et al. 2016). Otherwise, many deposits (e.g., the Wusihe deposit) distributed in the northern SYG are characterized by well strata-bound orebodies within the Dengying Formation and produce total Zn-Pb resources of ~ 10 Mt (Xiong et al. 2018). The characteristics with well strata-bound orebodies of these deposits led them previously to be regarded as sedimentary exhalative (SEDEX) deposits (Lin 2005; Zhu et al. 2018). New evidence, however, shows that the Wusihe deposit could also be an MVT deposit based on detailed ore geology and geochemical data (Rb–Sr isochron age of 411 ± 10 Ma; Xiong et al. 2018).

Spatial variations of Zn isotopic in hydrothermal system have been well studied with three alternative interpretations, including temperature variation (Mason et al. 2005; Toutain et al. 2008; Pašava et al. 2014), Rayleigh fractionation (Wilkinson et al. 2005; Kelley et al. 2009; Gagnevin et al. 2012; Zhou et al. 2014a, b; Gao et al. 2017), and mixing of multi-source fluids (Wilkinson et al. 2005). Among the above three mechanisms, fluid mixing has not yet been fully understood because it is difficult to distinguish between Rayleigh fractionation and fluid mixing (Wilkinson et al. 2005). For example, although the fluid mixing has been alternatively proposed to explain the Zn isotopic variations of sphalerite from an Irish ore field, the dominant mechanism was preferentially attributed to Rayleigh fractionation (Wilkinson et al. 2005). Previous studies have demonstrated that Fe is preferentially precipitated during early stages under relatively high-temperature conditions, both experimentally (Seewald and Seyfried 1990) and in hydrothermal systems (Kelley et al. 2009; Liu et al. 2012). Conversely, Cd is inclined to substitute for Zn in sphalerite during late stages under relatively low-temperature conditions (Liu et al. 2012; He et al. 2016; Wen et al. 2016). Therefore, theoretically, the Fe and Cd concentrations of sphalerite should be decreasing and increasing respectively from the early to late stages during fluid evolution. Moreover, the trends toward higher δ^{66} Zn values in later precipitated sphalerite have been found to be mirrored by decreasing Fe concentration, which has been interpreted as Rayleigh fractionation (John et al. 2008; Kelley et al. 2009). Therefore, the variations of Fe and Cd concentrations in sphalerite can be used as additional proxies to identify the Rayleigh fractionation. However, during fluid mixing, the distributions of Fe and Cd contentions in sphalerite are dominated by multiple parameters (e.g., pH, temperature, species of Zn complex, and fluid components), not only the temperature-dependent Rayleigh fractionation (Fujii et al. 2011; Liu et al. 2012; Pašava et al. 2014; He et al. 2016; Wen et al. 2016). Therefore, during fluid mixing, the Fe and Cd concentrations in sphalerite could not change like the process of Rayleigh fractionation of a single fluid. For example, the decreasing Cd content in sphalerite from the early to late stages have also been reported from the Tianbaoshan and Daliangzi deposits in the SYG, which was interpreted as mixing of multi-source fluids based on geological and geochemical evidence (Liu et al. 2012; He et al. 2016). Therefore, the Fe and Cd concentrations in sphalerite could also be effective proxies to identify fluid mixing. If the Zn isotopic compositions of sphalerite are dominated by mixing of multi-source fluids, it would be feasible to trace the potential Zn sources of ore-forming fluids (Wilkinson et al. 2005).

In this study, the Zn and S isotopic compositions, and Fe and Cd concentrations of sphalerite from the Wusihe Zn– Pb deposit were investigated. The sphalerite samples were collected in a vertical direction from the well strata-bound orebodies at No. 12 adit in the Wusihe deposit. Sampling sphalerite ores from the bottom to the top of the orebodies allow us to distinguish the spatial evolution of Zn isotopic compositions of sphalerite. Combining with Fe and Cd concentrations, we would like to identify the mechanism that controls Zn isotopic variations of sphalerite in this deposit. Moreover, the new results allow us to trace the possible sources of ore-forming Zn and S in the Wusihe Zn–Pb deposit.

2 Geological setting

The SYG polymetallic mineralization province is tectonically located in the southwest margin of the Yangtze Craton (Fig. 1A). The SYG region is mainly composed of Mesoproterozoic folded basements, Ediacaran to Paleozoic marine sedimentary sequences, and Mesozoic to Cenozoic terrestrial sedimentary sequences (Liu and Lin 1999; Xiong et al. 2018). The folded basements include the Dongchuan, Kunyang, and Huili Groups that largely consist of phyllite, graywackes, dolostone, and volcanic rocks (Zhou et al. 2014b; Xiong et al. 2018). The Ediacaran to Paleozoic marine sedimentary sequences are composed of carbonates, black shales, clasolites, and widespread sulfate-bearing evaporate (Liu and Lin 1999; Zhou et al. 2014b). The Mesozoic to Cenozoic terrestrial sedimentary sequences are characterized by entirely continental origin (Liu and Lin 1999; Zhou et al. 2014b). The Ediacaran Dengying Formation (mainly dolostone) is the most important orehosting strata in the SYG area, which holds over 50% of



Fig. 1 Map of the tectonic framework of South China (A). Regional geological map of the Sichuan–Yunnan–Guizhou Zn–Pb metallogenic province, showing the distribution of major faults and Zn–Pb ore deposits (B) (modified from Xiong et al. 2016)

the regional Zn–Pb deposits (Luo et al. 2019). Tectonically, those deposits that occurred in the SYG are distributed in the triangular area enclosed by the Anninghe, Ziyun-Yadu, and Mile-Shizong regional fractures (Fig. 1B; Guan and Li 1999; Zhang et al. 2015). These regional fractures, which act as conduits for hydrothermal fluids, commonly control the migration of ore-forming fluids (Zhang et al. 2015; Xiong et al. 2018). And a series of thrust-folds are the primary ore-controlling and ore-hosting structures for the regional Zn–Pb mineralization (Zhang et al. 2015; Xiong et al. 2018). The primary magmatic activity in the SYG area is the Permian Emeishan flood basalts (\sim 250 Ma; Zhou et al. 2002), which cover an area of over 250,000 km² and spatially coexist with the Zn–Pb deposits occurred in the SYG (Zhou et al. 2014a, b).

The Wusihe Zn–Pb deposit is located in the northern SYG in Ya'an city, Sichuan province (Fig. 1B). Within the Wusihe ore field, exposed strata mainly include the Ediacaran Dengying Formation and Cambrian to Permian marine sedimentary sequences (Fig. 2). The Wusihe Zn–Pb deposit is structurally controlled by the Wangmaoshan thrust-fault and the Wanlicun syncline (Fig. 2). The Wangmaoshan thrust-fault, crosscutting the Dengying Formation and the Lower Cambrian strata and dipping 65°–70° (Zheng 2012; Xiong et al. 2018), is the major orecontrolling fault in the mining area. The NS-trending

Wanlicun syncline, including the Ediacaran and Paleozoic strata and plunging to the south, largely controls the distribution of the orebodies in the Wusihe deposit (Zheng 2012; Xiong et al. 2018). In the Wusihe Zn–Pb deposit, the main orebodies occurred in the Wanlicun syncline and were strata-bound within the Dengying Formation strata (Fig. 2). The Wusihe Zn-Pb deposit yields Zn-Pb resources of \sim 3.7 Mt at average grades of 12.50 wt.% Zn and 3.15 wt.% Pb (Zheng 2012). The ore minerals include sphalerite and galena, where the sphalerite is the primary economic mineral. The major gangue minerals are dolomite and quartz, also with minor pyrite and calcite. Sphalerite ores are characterized by disseminated, banded and veined structures (Fig. 3). The alteration type of the host rock principally includes silicification, pyritization, and dolomitization, which only occurred in local positions with a developed fissure (Zheng 2012).

The mineralization stages of the Wusihe deposit have been described in detail in the previous study (Xiong et al. 2018), where four stages were identified: Stage I is the pyrite stage (fine- to medium-grained pyrite disseminated within the dolomite), Stage II is the pyrite-pyrrhotitegalena-sphalerite-bitumen stage (most important economic mineralization stage), Stage III is the sphalerite-galena stage (subordinate economic stage), and Stage IV is the bitumen-calcite stage (final hydrothermal stage). Fig. 2 Geological map of the Wusihe deposit, showing the major exposed strata, orecontrolling structures and sampling locations (modified from Lin 2005)





3 Sampling and analytical methods

3.1 Samples

Nine representative sphalerite samples were collected from the strata-bound orebodies in a vertical direction in No. 12 adit (Fig. 2). Three host rock samples were collected from the outcrop in the Wusihe ore field (N29.26°, E102.90°). All of these sulfide ores have a similar mineral assemblage, including sphalerite, galena, pyrite, and organic matter (Fig. 3). Sphalerite and galena are ubiquitous among all sulfide ores. Pyrite occurs in the wsh-12 and wsh-15. The organic matter is also common among these sulfide ores. The wsh-4, wsh-7, wsh-8 and wsh-10 were obtained from the bottom of the strata-bound orebodies in No. 12 adit, all with densely disseminated structure (Fig. 3A). The wsh-12 (Fig. 3B), wsh-14 (Fig. 3C) and wsh-15 (Fig. 3D) were collected from the middle position of the strata-bound orebodies in No. 12 adit, with sparsely disseminated, banded-disseminated, and veined structures, respectively. The wsh-16 (Fig. 3E) and wsh-17 (Fig. 3F) were sampled from the top of the strata-bound orebodies in No. 12 adit, with banded and veined structures, respectively. The separated sphalerite grains in the wsh-4, wsh-7, wsh-8, and wsh-10 samples were hand-picked under a binocular microscope. Other sphalerite samples for analysis were obtained by the Micro-Drill Sampling System (MSS) at the State Key Laboratory of Ore Deposit Geochemistry, Institute of Geochemistry, Chinese Academy of Sciences. The tungsten steel used in the MSS has a diameter of 0.5 mm, which can greatly avoid the contamination from host rocks.

3.2 Methods

Sphalerite and carbonate samples were digested by HNO₃ (15 mol/L) and HCl (6 mol/L), respectively, at 120 °C for 24 h. Sample solutions were evaporated to dryness and then were digested in 2 mL 2N HCl. The Zn was purified from the other matrix elements using anion exchange chromatography, as reported by Tang et al. (2006). Zinc isotope ratios and concentrations were measured using Nu Plasma MC-ICP-MS in a low-resolution mode at the State Key Laboratory of Ore Deposit Geochemistry, Institute of Geochemistry, Chinese Academy of Sciences. Mass discrimination effects were corrected using a combined sample-standard bracketing (SSB) (Li et al. 2008; Zhou et al. 2014a, b). Accuracy and reproducibility were assessed by replicate analyses of the international standards BHVO-2 (basalt), which yielded an average δ^{66} Zn value of $0.30 \pm 0.04 \%$ (2 σ , n = 6), consistent with previously reported values (Herzog et al. 2009; Liu et al. 2016). Each result is the mean value over the N number of repeats and all results are reported relative to the JMC Lyon Zn standard (Maréchal et al. 1999) based on the difference between the IRMM3702 and JMC Lyon Zn solution (Δ^{66-} $Zn_{IRMM3702-JMC}$ Lyon = 0.30 %; Moeller et al. 2012; Samanta et al. 2016).

Sulfur isotope analyses were carried out using a Thermo Flash 2000 at the State Key Laboratory of Resource Utilization and Environmental Restoration, Institute of



Fig. 3 The specimen photo of the bottom wsh-7 sample, showing the densely disseminated structure (A); the specimen pictures of the middle part wsh-12 (B), wsh-14 (C) and wsh-15 (D) samples, showing the micro-drilled positions, and sparsely disseminated, banded-disseminated and veined structures, respectively; The specimen photos of the top wsh-16 (E) and wsh-17 (F) samples, showing the micro-drilled positions, and banded and veined structures, respectively; the specimen photo and photomicrographs of main mineralization (G, H and I), *Gn* galena, *Dol* dolomite, *Sp* sphalerite, *Py* pyrite, *OM* organic matter

Geographic Science and Natural Resources Research, Chinese Academy of Sciences. IAEA-S-1 (Ag₂S; δ^{34-} S_{CDT} = $-0.3 \pm 0.2 \%$, n = 6) were used as the external standards. The analysis errors (2σ) are better than 0.1 ‰ of the replication of standard materials. Sulfur isotopic compositions are reported relative to the Canyon Diablo Troilite (CDT).

The concentration of Fe and Cd of sphalerite were measured by the PinAAcle 900F atomic absorption spectrometer at the State Key Laboratory of Environmental Geochemistry, Institute of Geochemistry, Chinese Academy of Sciences. The Fe and Cd concentration of bulk-rock samples (host rocks) were measured by ICP-MS at ALS Chemex (Guangzhou) Co. Ltd. The relative analysis error of the Fe and Cd concentration is less than 5%. Moreover, we use the Fe_(ppm)/Zn_(%) and Cd_(ppm)/Zn_(%) to evaluate the Fe and Cd contents in sphalerite, which can diminish the

interference of different proportion incorporation of host rocks.

4 Results

Zinc and sulfur isotopic compositions, and Fe and Cd concentrations of sphalerite from the Wusihe Zn–Pb deposit are shown in Table 1. The δ^{66} Zn values in sphalerite vary from 0.14 to 0.62 ‰ and yield an increasing trend from the bottom to the top of the well strata-bound orebodies in No. 12 adit (Fig. 4). The δ^{34} S values of sphalerite vary from 5.1 to 15.4 ‰ and also yield an increasing trend from the bottom to the top of the well strata-bound orebodies in No. 12 adit, which becomes more obvious when the wsh-14 sample is excluded (Fig. 5). In addition, the δ^{66} Zn and δ^{34} S values of sphalerite are

Table 1	The δ^{66} Zn $-\delta^{34}$ S	values and Zn-Fe-C	Id contents in sphalerite	samples, host rock	samples, and basement	samples

No.	Positions	Objects	Zn (wt.%)	Fe (wt.%)	Cd (× 10^{-6})	δ ⁶⁶ Zn (‰)	δ ³⁴ S (‰)
WSH-4	Bottom orebody	Sphalerite	_	0.12	5864	0.18 ± 0.05	10.5
WSH-7	Bottom orebody	Sphalerite	_	0.07	7202	0.14 ± 0.06	10.5
WSH-8	Bottom orebody	Sphalerite	_	0.14	3058	0.21 ± 0.02	11.6
WSH-10	Bottom orebody	Sphalerite	_	0.08	9214	0.19 ± 0.06	8.4
WSH-12-1	Middle orebody	Sphalerite ores	22.54	0.68	776	0.35 ± 0.02	13.4
WSH-12-2	Middle orebody	Sphalerite	49.02	1.25	1633	0.38 ± 0.03	12.7
WSH-12-3	Middle orebody	Sphalerite	46.54	0.83	1983	0.31 ± 0.03	12.8
WSH-14-1	Middle orebody	Sphalerite	51.52	1.81	985	0.38 ± 0.03	7.1
WSH-14-2	Middle orebody	Sphalerite	52.02	1.76	716	0.51 ± 0.06	7.4
WSH-14-3	Middle orebody	Sphalerite	46.98	1.18	1154	0.44 ± 0.01	5.1
WSH-14-4	Middle orebody	Sphalerite	47.46	7.50	622	0.44 ± 0.07	11.7
WSH-14-5	Middle orebody	Sphalerite	54.39	1.85	884	0.41 ± 0.05	13.3
WSH-14-6	Middle orebody	Sphalerite	48.72	2.03	607	0.50 ± 0.08	11.5
WSH-14-7	Middle orebody	Sphalerite	49.99	2.16	629	0.41 ± 0.07	12.6
WSH-14-8	Middle orebody	Sphalerite	47.02	1.74	802	0.52 ± 0.11	12.5
WSH-14-9	Middle orebody	Sphalerite	42.41	1.81	1062	0.41 ± 0.09	13.0
WSH-15-4	Middle orebody	Sphalerite	40.96	2.24	737	0.46 ± 0.08	14.1
WSH-15-5	Middle orebody	Sphalerite	49.39	2.68	711	0.38 ± 0.05	14.1
WSH-15-6	Middle orebody	Sphalerite	52.70	2.82	777	0.44 ± 0.06	13.1
WSH-16-1	Top orebody	Sphalerite	45.99	4.32	1056	0.44 ± 0.06	14.9
WSH-16-2	Top orebody	Sphalerite	40.87	4.14	1095	0.39 ± 0.01	15.4
WSH-16-3	Top orebody	Sphalerite ores	37.43	4.08	734	0.39 ± 0.05	15.0
WSH-17-3	Top orebody	Sphalerite	47.82	6.38	468	0.60 ± 0.05	14.1
WSH-17-4	Top orebody	Sphalerite	48.28	7.31	426	0.62 ± 0.06	14.5
No.	Positions		Host rocks		Zn (ppm)		δ ⁶⁶ Zn (‰)
WSH ₁₈ -5	Shallow ground		Carbonates		204		0.23 ± 0.04
WSH18-8	Shallow ground		Carbonates		25		0.50 ± 0.01
WSH ₁₈ -16	Shallow ground		Carbonates		9		0.20 ± 0.01
No.	Positions		Basement rocks		Zn (ppm)		δ ⁶⁶ Zn (‰)
WC-1	Shallow ground		Carbonates		3		0.21 ± 0.04
HS-4	Shallow ground		Carbonates		24		0.10 ± 0.04
TBS ₁₆ -2	Shallow ground		Graywackes		61		0.30 ± 0.03
Td1900-8	Shallow ground		Phyllites		115		0.15 ± 0.05
Td1900-3	Shallow ground		Phyllites		199		0.34 ± 0.02

"-" mean not measured; the Zn concentration of the bottom sphalerites (hand-picked sphalerite separates) theoretically is $\sim 67\%$ considering their low Fe concentration; the data of the basement samples are collected from Zhang et al. (2019)

covariant in this deposit (Fig. 6). Three host rock samples contain Zn from 9 ppm to 204 ppm and yield δ^{66} Zn values from 0.20 to 0.50 ‰. Five basement samples contain Zn from 3 to 199 ppm and yield δ^{66} Zn values from 0.10 to 0.34 ‰.

5 Discussion

5.1 Possible mechanisms for Zn isotopic variations

The Zn isotopic compositions of sphalerite from the Wusihe Zn–Pb deposit range from 0.14 to 0.62 ‰ that increase from the bottom to the top of the well strata-bound orebodies (Fig. 4). The spatial evolution of Zn isotopic



Fig. 4 The correlation between δ^{66} Zn and Fe (ppm)/Zn (%) values (**A**); the correlation between δ^{66} Zn and Cd (ppm)/Zn (%) values (**B**). Using the Fe (ppm)/Zn (%) and Cd (ppm)/Zn (%) values to evaluate the Fe and Cd contents in sphalerite can diminish the interference from different proportion incorporation of host rocks. The gray polygon data were cited from Zhu et al. (2018)

composition in hydrothermal system has been attributed to three potential mechanisms, including temperature variation, Rayleigh fractionation, and mixing of multi-source fluids (Mason et al. 2005; Wilkinson et al. 2005; Toutain et al. 2008; Kelley et al. 2009; Gagnevin et al. 2012; Pašava et al. 2014; Zhou et al. 2014a, b; Gao et al. 2017).

The potential influence of temperature gradients on Zn isotopic fractionation has been examined in previous studies, but their results obtained at different temperature conditions are not consistent. For example, the systematically increasing δ^{66} Zn values (- 0.03 to 0.23 ‰) away from the hydrothermal vent (~ 300 °C) in the Alexandrinka VMS deposit were ascribed to temperature-controlled variations (Mason et al. 2005). Furthermore, in the Merapi volcano system (590-297 °C), the Zn isotopic composition of high-temperature fumarolic gases (0.05–0.85 ‰) is much lower than that of their low-temperature condensates (1.48–1.68 ‰), which was also temperature-dependent interpreted as Rayleigh



Fig. 5 The correlation between δ^{34} S and Fe (ppm)/Zn (%) values of all samples (A1), without the wsh-14 sample (A2); The correlation between δ^{34} S and Cd (ppm)/Zn (%) values of all samples (B1), without the wsh-14 sample (B2). The gray polygon data were cited from Zhu et al. (2018)



Fig. 6 The correlation between δ^{66} Zn and δ^{34} S values of all samples (A1), without the wsh-14 sample (A2). The gray polygon data were cited from Zhu et al. (2018)

fractionation (Toutain et al. 2008). However, several studies found that no correlation exists between the δ^{66} Zn values and temperatures when the temperatures were lower than 300 °C, either in an experiment (30 and 50 °C; Maréchal and Sheppard 2002) or in a hydrothermal system (60–250 °C; Wilkinson et al. 2005). These observations may indicate that the influence of temperature gradients on Zn isotope fractionation is negligible when the temperatures are lower than 300 °C. The homogenization temperatures of fluid inclusions in hydrothermal minerals from the Wusihe Zn–Pb deposit vary from 140 to 285 °C and mostly cluster between 150 and 250 °C (Lin 2005; Xiong et al. 2016). Therefore, it is suggested that temperature gradients could not dominate the Zn isotopic variations of sphalerite from the Wusihe Zn–Pb deposit.

Experiments demonstrated that early precipitated sulfides from the same solution are ⁶⁴Zn-enriched, whereas late precipitated sulfides are ⁶⁶Zn-enriched (Archer et al. 2004). Studies in hydrothermal systems have also confirmed that Rayleigh distillation could fractionate Zn isotopes between sphalerite and fluid, resulting in systematically increasing δ^{66} Zn values in sphalerite from the early to late stages (Wilkinson et al. 2005; Kelley et al. 2009; Gagnevin et al. 2012; Zhou et al. 2014a, b; Gao et al. 2017). Assuming that the systematically increasing δ^{66} Zn values of sphalerite from the Wusihe deposit were dominated by Rayleigh fractionation, it would indicate that, in a vertical direction, the bottom sphalerites precipitated earlier than the upper sphalerites. However, there is no evidence that the mineralization stages of this deposit are associated with the height positions of the well stratabound orebodies. And these sulfide ores are similarly composed of sphalerite, galena, and minor pyrite (Fig. 3), which could indicate these sulfide ores formed from the same stage (stage II; Xiong et al. 2018). Moreover, Rayleigh fractionation model cannot explain the correlations of δ^{66} Zn values and Fe–Cd concentrations in sphalerite from this deposit. Previous studies have demonstrated that Fe is preferentially precipitated during early stages under relatively high-temperature conditions, both experimentally (Seewald and Seyfried 1990) and in hydrothermal systems (Kelley et al. 2009; Liu et al. 2012). Conversely, Cd is inclined to substitute for Zn in sphalerite during late stages under relatively low-temperature conditions (Liu et al. 2012; He et al. 2016; Wen et al. 2016). Assuming that Zn isotopic variations of sphalerite from the deposit are dominated by Rayleigh fractionation, there would be negative correlations between δ^{66} Zn values and Fe content and positive correlations between δ^{66} Zn values and Cd content in sphalerite. For example, during Rayleigh fractionation processes, higher δ^{66} Zn values in later precipitated sphalerite have been found to possess lower Fe concentration in natural hydrothermal systems (John et al. 2008; Kelley

et al. 2009). In fact, the δ^{66} Zn values correlate with Fe positively and with Cd negatively in sphalerite from this deposit (Fig. 4). The clear correlations between δ^{66} Zn values and Fe and Cd contents in sphalerite indicate that their variations are dominated by a similar mechanism. Therefore, the Rayleigh fractionation could not be responsible for the spatial evolution in δ^{66} Zn values of sphalerite in the Wusihe deposit.

Similar to the Wusihe deposit, the decreasing Cd content in sphalerite from the early to late stages have also been reported from the Tianbaoshan and Daliangzi Zn-Pb deposits in the SYG, which was interpreted as mixing of multi-source fluids based on geological and geochemical evidence (Liu et al. 2012; He et al. 2016). The mixing of multi-source fluids could also result in a spatial evolution in δ^{66} Zn values of sphalerite on a deposit scale (Wilkinson et al. 2005). The Emeishan basalts, basements and host rocks are three potential sources for the regional Zn-Pb mineralization in the SYG (Huang et al. 2003; Han et al. 2007; Ye et al. 2011; Zhou et al. 2014b, 2018; Zhu et al. 2016; Xiong et al. 2018). Recently, the ore-forming age of the Wusihe deposit has been constrained to 411 ± 10 Ma via the Rb-Sr isochron (Xiong et al. 2018), and thus the much younger Emeishan basalts (~ 250 Ma; Zhou et al. 2002) are impossible to contribute to the mineralization of the Wusihe Zn-Pb deposit. Five basement samples, including carbonates, graywackes, and phyllites, yield low δ^{66} Zn values from 0.10 to 0.34 ‰, similar to the Zn isotopic signals documented in the bottom sphalerite samples of the well strata-bound orebodies in this deposit. Moreover, the basements of an Irish orefield, where the rock assemblage is similar to here (mainly graywackes and volcanic rocks), also have been reported to have low Zn isotopic signatures (0.00-0.18 %; Wilkinson et al. 2005; Gagnevin et al. 2012). On the other hand, three carbonate samples collected from the host rocks yield higher δ^{66} Zn values of 0.20-0.50 ‰, similar to the Zn isotopic signatures recorded in the upper sphalerites of the well stratabound orebodies in this deposit. Furthermore, biogenic carbonates have been studied with heavy Zn isotopic compositions (mean = 0.91 ‰; Pichat et al. 2003). Combining with the fact that the biogenic carbonates are widespread in the host rocks (Feng et al. 2009; He et al. 2016), the host rock-derived fluid could have higher Zn isotopic signature than that of basement-derived fluid. Therefore, the spatial variation in δ^{66} Zn values of sphalerite from the Wusihe Zn-Pb deposit could result from mixing basement-derived fluid of lighter Zn isotopic composition with host rock-derived fluid of heavier Zn isotopic composition (Figs. 4 and 7).

Fig. 7 Mixing model sketch of the Wusihe ore deposit. The mixing occurred under the black shales; the fluid from host rocks yields higher δ^{66} Zn and δ^{34} S values, whereas the fluid from folded basements yields lower δ^{66} Zn and δ^{34} S values



5.2 Possible mechanisms for sulfur isotopic variation

The sulfur isotopic compositions of sphalerite from the Wusihe Zn-Pb deposit vary from 5.1 to 15.4 ‰ (mean = 12.1 %; Fig. 6), consistent with the previous study where the δ^{34} S values of sulfide were reported to range from 7.1 to +15.5 % (mean = 11.4 %; Xiong et al. 2018). In the Wusihe Zn–Pb deposit, an increasing trend in δ^{34} S values from the bottom to the top of the well strata-bound orebodies also exists, which becomes more obvious when the wsh-14 sample is excluded (Fig. 5). The increasing δ^{34} S values in sulfide could be alternatively attributed to kinetic fractionation from single hydrothermal fluid due to fluid cooling (Rye and Ohmoto 1974; Böttcher et al. 1998; Gagnevin et al. 2012; Zhu et al. 2017). Similar to the δ^{66} Zn values in the deposit, the δ^{34} S values also correlate with Fe positively and with Cd negatively in sphalerite (Fig. 4), which cannot be attributed to kinetic fractionation (Bethke and Borton 1971; Liu et al. 2012; Belissont et al. 2014; He et al. 2016; Wen et al. 2016). As so, the kinetic fractionation could not be responsible for the sulfur isotopic variation of sphalerite in the Wusihe Zn-Pb deposit.

Previous studies also demonstrated that the temporally and spatially dependent $\delta^{34}S$ values could be interpreted as

mixing multi-source fluids (Rye and Ohmoto 1974; Bortnikov et al. 1995; Gagnevin et al. 2012; Thiessen et al. 2016; Zhu et al. 2017). The Emeishan basalts, basements, and host rocks are three potential sulfur sources for the regional Zn-Pb mineralization in the SYG (Huang et al. 2003; Han et al. 2007; Ye et al. 2011; Zhou et al. 2014b, 2018; Zhu et al. 2016; Xiong et al. 2018). The δ^{34} S values of sphalerite from the Wusihe Zn-Pb deposit range from 5.1 to 15.4 ‰ (mean = 12.1 ‰; Fig. 6), significantly different from mantle-derived magmatic sulfur (~ 0 %); Chaussidon et al. 1989), and thus that the magma-derived (Emeishan basalts) sulfur is ruled out. In the SYG, S-bearing evaporates are widespread in the host rocks, the sulfur isotopic compositions of these evaporates have been reported to range from 20.0 to 38.7 % (mean = 29.0 %); Zhang et al. 2004; Xiong et al. 2016). The homogenization temperatures of the ore-forming fluids in the deposit yield an average value of 197 °C (n = 47; Lin 2005), suggesting ~ 10 ‰ sulfur isotopic fractionation between sulfides and sulfates (200 °C, Δ^{34} S_{sulfate-sulfide} = 10 ‰; Machel et al. 1995). However, in the Wusihe deposit, the mean sulfur isotopic composition of sulfides is ~ 12.0 % (here; Xiong et al. 2018), which is ~ 17 % lower than that of evaporates (29.0 ‰). It is suggested, thus, that there could be another sulfur source providing lighter sulfur isotopes than

evaporates for the Wusihe deposit. The much lighter sulfur isotopic compositions of sulfide have been reported from the Tianbaoshan deposit in the SYG (0.2–5.0 ‰), which has been constrained to basement-derived sulfur after precluding the possibility from host rocks and mantle magma (Zhu et al. 2016). Therefore, the basements could be another potential sulfur source supplying lighter sulfur isotopes than evaporates for the Wusihe deposit. As so, similar to Zn isotopic system in the deposit, mixing of the ore-forming fluids from basement rocks and host rocks, respectively, could dominate the spatial variation in δ^{34} S values of sphalerite from the Wusihe Zn–Pb deposit (Figs. 5 and 7).

If it is the case, the mixing model may suggest an explanation to the sulfur isotopic outliers in the wsh-14 sample collected from the middle part of the well stratabound orebodies (Fig. 3C). For the other hand specimens, δ^{34} S values of micro-drilled sphalerite samples within the same specimen only slightly change, without exceeding 1.0 ‰, which suggests that these hand specimens could only record the homogeneous sulfur isotopic signature of a single fluid. However, the sulfur isotopic variation of the wsh-14 sample (5.16-13.05 %), with nine micro-drilled samples, almost covers the entire range of that of all the sphalerite samples from the deposit (5.16–15.43 ‰). Note that the wsh-14 sample was collected from the middle part of the well strata-bound orebodies, which may be a right position for fluid mixing. Here, we ascribe the largely varying δ^{34} S values in the wsh-14 sample to a localized heterogeneous mixing of two-source fluids and thus that the wsh-14 sample could document the sulfur isotopic characters of both the basements-derived fluid and host rockderived fluid.

5.3 Implications from Zn and S isotopes

Previous studies have different opinions on the origin and mechanism of the Wusihe deposit. Lin (2005) and Zhu et al. (2018) argued that ore-forming components of the Wusihe deposit had been sourced from SEDEX hydrothermal systems, namely from the Mesoproterozoic basements. However, Xiong et al. (2018) demonstrated that the Wusihe deposit belongs to MVT deposit, and further proposed that Pb and Sr have been sourced from both host and basement rocks.

In this contribution, both independent Zn and S isotopic systems indicate that there is a fluid mixing from both host and basement rocks (Fig. 7), which is consistent with the inferences by Xiong et al. (2018). The fluid from the host rocks yields higher δ^{66} Zn and δ^{34} S values, whereas the fluid from the basement rocks yields lower δ^{66} Zn and δ^{34} S values. Moreover, the previous study demonstrated that mixing of two-source fluids with contrasting zinc and

sulfur isotopic composition would induce covariant δ^{66} Zn and δ^{34} S values in sphalerite at a deposit scale (Wilkinson et al. 2005). The covariant δ^{66} Zn and δ^{34} S values in sphalerite are indeed observed in the Wusihe deposit (Fig. 6), which further validates the existence of fluid mixing. The mineralization deposition could happen when two source fluids mixed together below a black shale layer (Fig. 7), associating the changes in physical and chemical conditions of ore-forming fluids. The spatial variation in δ^{66} Zn and δ^{34} S values could be ascribed that the rock-derived brine migrated to the mineralization position earlier and inclined to fill stratigraphically higher spaces owing to fluid-pressure, while the subsequent basement-derived metamorphic fluids mainly migrated to stratigraphically lower position owing to limited spaces left.

6 Conclusion

In this contribution, Zn and S isotopes combined with Fe and Cd concentrations of sphalerite from the Wusihe Zn–Pb deposit in the SYG, Southwest China, have been examined. The results allow us to draw three conclusions as follows:

- 1. The spatial variation of δ^{66} Zn and δ^{34} S values of sphalerite from the Wusihe deposit resulted from the mixing of two-source fluids with contrasting Zn and S isotopic composition.
- 2. The fluid from the host rocks yields higher δ^{66} Zn (~ 0.6 ‰) and δ^{34} S values (~ 15 ‰), whereas the fluid from the basement rocks yields lower δ^{66} Zn (~ 0.2 ‰) and δ^{34} S values (~ 5 ‰).
- 3. The covariant isotopic compositions of Zn and S could be an effective proxy in reflecting fluids mixture, in which the Zn and S were transported together.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interest.

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