Discovery of Disulfane (H$_2$S$_2$) in Fluid Inclusions in Rubies from Yuanjiang, China, and Its Implications

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Abstract: Raman spectra of fluid inclusions in gem rubies from Yuanjiang deposit (China) within the Ailao Shan–Red River (ASRR) metamorphic belt showed the presence of compounds such as CO$_2$, COS, CH$_4$, H$_2$S, and elemental sulfur (S$_8$), accompanied by two bands at approximately 2499 and 2570 cm$^{-1}$. These two frequencies could be assigned to the vibrations of disulfane (H$_2$S$_2$). This is the second case of the sulfane-bearing fluid inclusions in geological samples reported, followed by the first in quartzite from Bastar Craton of India. The H$_2$S$_2$ was likely in situ enclosed by the host rubies rather than a reaction product that formed during the cooling of H$_2$S and S$_8$, suggesting sulfanes are stable at elevated temperatures (e.g., $>$600 °C). By comparing the lithologies and metamorphic conditions of these two sulfane-bearing cases (Bastar and Yuanjiang), it is suggested that amphibolite facies metamorphism of sedimentary sequence that deposited in a continental platform setting might favor the generation of sulfanes. Sulfanes may play an important role in the mobilization of Cr that is essential for ruby crystallization.

Keywords: rubies; fluid inclusion; Ailao Shan–Red River metamorphic belt

1. Introduction

Numerous volatile species (e.g., CO$_2$, CO, O$_2$, C$_3$H$_8$, and N$_2$) in fluid inclusions are well documented in the literature [1]. Sulfur, an element with redox states ranging from −2 to +6, can be found in fluid inclusions as sulfur compounds of different valence, e.g., S$_8$, COS, SO$_2$, and H$_2$S in gaseous, and SO$_4^{2−}$, HSO$_4^{−}$, and HS$^{−}$ solutes in aqueous fluids [1,2]. Polysulfanes (H$_2$S$_n$ with $n$ > 1; or hydrogen polysulfides) are generally found in natural gas and crude sulfate oils [3], but their presence in natural fluid inclusions is rare. Recently, disulfane (H$_2$S$_2$) was described for the first time in natural CH$_4$–H$_2$S inclusions in Archaean quartzite from Bastar Craton of India [4]; recognition and assignment of the H$_2$S$_2$-related Raman bands was accomplished according to quantum chemical calculations based on the density functional theory. Given that sulfanes have been observed in both sulfur-bearing fluid inclusions and sulfur-free types, the possibility that disulfane formed by the post-entrapment reaction between molten sulfur and H$_2$S was ruled out by Hurai et al. [4]. They accordingly inferred that the disulfane or any other unknown S-H-bearing species that could dissociate into disulfane along the cooling path should be present before the fluid was trapped by quartz. This is critical to understanding the fluid process and is thus of significance to investigate. On the other hand, many aspects regarding H$_2$S$_2$, e.g.,
the favorable formation condition, thermochemical stability, and its potential roles in the mobilization of specific elements, remain largely unclear and require further investigation.

For marble-hosted ruby deposits from Central and Southeast Asia, studies on fluid inclusions show that sulfur-bearing fluids with the composition of CO$_2$–H$_2$S (±COS ± S$_8$) occurred during corundum growth [2,5]. These complex fluids are metamorphic in origin and can be explained by decarbonation of limestone (CO$_2$) and reduction in sulfate (elemental sulfur, hydrogen sulfide, and COS) [2].

Here, we report the second occurrence of H$_2$S composition in ruby-hosted fluid inclusions from the Ailao Shan-Red River (ASRR) metamorphic belt. It corroborates the presence of sulfanes in high-grade metamorphic fluids. We further discussed its origin, thermochemical stability, and geochemical implications.

2. Geological Setting

The geology of the ASRR metamorphic belt has been documented in some detail by Tapponnier et al. [6] and Leloup et al. [7]. The ASRR metamorphic belt, a Tertiary left-lateral strike-slip fault zone, is traditionally interpreted to play an important role in the accommodation of the extrusion of the Indochina block during the India–Eurasia collision (Figure 1) [6,7]. The elongate Ailao Shan metamorphic massif is composed of mylonitic gneisses, migmatites, granitic pods, amphibole-bearing alkaline intrusions, and marble boudins [8,9]. U–Pb ages of zircon, xenotime, titanite from syntectonic leucogranites combined with Ar–Ar radiometric dating of biotite, muscovite, and K-feldspar within the Ailao Shan set the timing of left-lateral shearing activity from 35 Ma to 17 Ma [7–9]. In addition, 23.4 ± 0.3 Ma U–Pb ages of titanite syngenetic with rubies in marbles from Yuanjiang, which recorded the age of ruby mineralization, are also within this range [10]. Peak metamorphism occurred at 4.5 ± 1.5 kbar and 710 ± 70 °C, as revealed by a petrographic investigation on Ailao Shan massif [11]. Greenschist facies retrograde metamorphism occurred at pressures (P) of <3.8 kbar and temperatures (T) around 500 °C [11].

![Figure 1. Simplified tectonic sketch of SE Asia showing the ASRR metamorphic terranes (modified after Leloup et al. [7]) with marked study area. The ASRR strike-slip metamorphic zone is comprised of four metamorphic terranes: Ailao Shan, Xuelong Shan (XLS), Diancang Shan (DCS), and Day Nui Con Voi (DNCV).](image-url)
3. Materials and Methods

Corundum-bearing marbles in this investigation are from the Yuanjiang region in the central segment of the Ailao Shan mountain (Figure 2). They are exposed as boudins within meta-sedimentary hosts including gneiss, amphibolite, and leptynite. The marbles are medium- to coarse-grained in size and white and light gray in color. Most marble mainly comprises calcite with small amounts of micas, feldspars. Other minerals identified in marbles include ruby, rutile, spinel, sphene, graphite, pyrrhotite, and pyrite.

Figure 2. Structural map of Ailao Shan metamorphic range (modified after Leloup et al. [7]): MVF—mid-valley fault; RFF—range front fault.

We conducted fluid inclusion studies on both primary rubies in the marble matrix and secondary rubies (Figure 3) recovered from the sediments around the village of Shaku in Yuanjiang County. These ruby fragments were cut and doubly polished to a thickness of 150–200 µm. An optical microscope was used for petrographic observation of fluid inclusions. In addition, four pieces of ruby-bearing marbles were doubly polished for fluid inclusions investigations.

Raman analyses on fluid inclusions were completed at HORIBA Instrument (Shanghai) Co., Ltd., employing a confocal LabRAM HR Evolution Raman microspectrometer (Horiba, Ltd., Kyoto, Japan) coupled to an Olympus BX-51 microscope. Raman spectra were collected with a 100× objective lens. To suppress fluorescence from the host ruby, a 473 nm excitation wavelength of 25 mW, air-cooled solid-state Nd-YAG laser was selected, allowing detection of bands above 1500 cm⁻¹. The analytical settings included 600 grooves/mm.
gratings, 10–20 s acquisition time, and 2–3 times accumulations. Spectra processing was performed with baseline correction unless otherwise specified. Daily calibration of the spectrometer was completed with a monocrystalline silicon wafer (520.7 cm⁻¹).

4. Results

The fluid inclusions can be subdivided into three types according to their entrapment sequence relative to ruby growth, i.e., primary, pseudo-secondary, and secondary. A detailed description of fluid inclusion classification was given in Huang et al. [10] and is briefly outlined in Table 1. Mostly, these fluid inclusions are approximately 20–60 µm in diameter (Figure 4). Raman analyses showed complex fluid composition in the primary and pseudo-secondary fluid inclusions (Figures 4 and 5); they comprise CO₂, H₂S, COS, S₈, and CH₄ (Figure 5). Components such as CO₂, H₂S, COS, S₈, and CH₄ were also observed in the fluid inclusions of secondary origin. Moreover, solids in the fluid inclusions cavities, identified by Raman spectroscopy, include diaspore, arsenopyrite, dawsonite, gibbsite, pyrite, and rutile (Figures 4 and 5 and Table 1).

![Figure 4. Photomicrographs of H₂S₂–bearing fluid inclusions. These fluid inclusions sometimes contain rutile crystals (a,c) and diaspore crystals (a-d): L—liquid carbonic phase.](image)

| Inclusion Types | Occurrence Description | Fluid Components | Solids                        |
|----------------|------------------------|------------------|-------------------------------|
| Primary        | Oriented clusters throughout the host or small clusters in the core of the host, sometimes occurred as isolated or clusters within colored growth zonation | CO₂–H₂S–COS–S₈–H₂S₂–CH₄ | Diaspore, gibbsite, dawsonite, rutile, and arsenopyrite |
| Pseudo-secondary | In intragranular fractures that do not traverse into the rim of the host | CO₂–H₂S–COS–S₈–H₂S₂–CH₄ | Diaspore, arsenopyrite, and pyrite |
| Secondary      | In sealed fractures present in planar arrays that traversed the growth zone of crystals | CO₂–H₂S–COS–S₈–CH₄ | Diaspore |
Moreover, two Raman bands at 2499 and 2570 cm\(^{-1}\) were observed in fluid inclusions of both primary and pseudo-secondary origins (Figure 5).

![Raman spectra](image)

**Figure 5.** Raman spectra of the fluid inclusion as shown in Figure 4a: (a) the Raman spectrum corresponds to rutile; (b) Raman spectra showed that the transparent crystal was diaspore. Asterisk denoted the Raman signals from the hosting corundum; (c) the components of the included fluid included native sulfur (S\(_8\)), CO\(_2\), H\(_2\)S, and CH\(_4\). Two bands centered at 2499 and 2570 cm\(^{-1}\) can be clearly seen.

5. Discussion

5.1. Identification of Disulfane

The density functional theory method is a useful method for molecular structure prediction [3]. It is especially suitable for unstable reactive species (e.g., sulfur-rich compounds) whose structures and spectroscopic properties can hardly be obtained experimentally. Hurai et al. [4] employed the B3LYP hybrid functional and the basis set correlation-consistent polarized valence triple-zeta (cc-pVTZ) for the Raman frequency calculations of the polysulfanes. Quantum chemical calculations of both pure H\(_2\)S\(_n\) (n = 2–6) compounds and S\(_8\)-sulfane interactions had been conducted by these authors. They observed a similar pattern between the theoretical Raman bands of various systems comprising S\(_8\)-polysulfane and the examined sulfur-bearing fluid inclusions. Accordingly, they indicated that the
2488 cm$^{-1}$ Raman band corresponds to H-bonded S-H stretch vibrations of the H$_2$S$_2$ ⋯ S$_8$ complex, whereas the 2503 cm$^{-1}$ band corresponds to non-H-bonded S-H stretching vibration of the H$_2$S$_2$ ⋯ S$_8$ complex. Another band at 2574 cm$^{-1}$ was interpreted to reflect S-H stretch vibration in H$_2$S + H$_2$S$_2$ ⋯ S$_8$ [4].

Fluid inclusion in marble-hosted rubies from Central and Southeast Asia have been extensively studied [2,5]. However, the 2499 and 2570 cm$^{-1}$ Raman bands, which have not been detected in rubies from other deposits, are the most notable features of Raman bands of the investigated fluid inclusions within Yuanjiang rubies. The pioneering work of sulfane-bearing fluid inclusions carried out by Hurai et al. [4] enables us to interpret these two Raman bands. Deconvolution of the broad peak at 2499 cm$^{-1}$ using PeakFit software (version 4.12) revealed two bands centered at 2484 and 2499 cm$^{-1}$ (Figure 6c). Hurai et al. [4] have shown the broadband around 2497 cm$^{-1}$ actually comprised two peaks centered at 2482–2488 cm$^{-1}$ and 2497–2499 cm$^{-1}$. Therefore, the two bands of 2484 and 2499 cm$^{-1}$ observed in Yuanjiang rubies could be ascribed to H$_2$S$_2$-related vibrations and positively identified the presence of sulfanes.

![Figure 6](image_url)

**Figure 6.** Raman spectra of fluid inclusions showing the H$_2$S$_2$-related bands at approximately 2499 and 2570 cm$^{-1}$: (a) Raman spectrum without baseline correction of the inclusion as shown in Figure 4a; (b) Raman spectrum without baseline correction of the inclusion as shown in Figure 4b; (c) deconvolution of the 2499 cm$^{-1}$ band reveals a hidden band centered at 2484 cm$^{-1}$. The bands at 2484 and 2499 cm$^{-1}$ reflect the respective H-bonded and non–H–bonded SH-stretching vibrations of the H$_2$S$_2$ ⋯ S$_8$ ⋯ H$_2$S$_2$ complex.

### 5.2. Origin of Sulfane in Yuanjiang Rubies

The sulfur–sulfur bonds in S$_8$ have rather a high bond enthalpy ((264 kJ mol$^{-1}$), which results in sulfur atoms forming as chains, rings, or clusters in numerous compounds [3]. Sulfanes, H$_2$S$_n$, are among the most basic of such species. However, reports regarding the natural occurrences of sulfanes are still limited; they typically occur in crude sulfane oils [3]. Moreover, they could be found in hot underground deposits of natural gas where H$_2$S and elemental sulfur coexist [3] and sulfur-containing hydrothermal environments [12–14]. Both elemental sulfur and H$_2$S are common species coexisting in fluid inclusions [2,5], and sulfanes can be produced by the interaction of S$_8$ with H$_2$S via the reaction H$_2$S$_n$ + nS$_{liq} \rightarrow$ H$_2$S$_{n+1}$ [3]; however, their presence in fluid inclusions is still very rare. Thus far, the sulfane in fluid inclusions has been only reported in Archaean mica-rich quartzite from near the Saigon village within the Bastar Craton, India [4], to the best of our knowledge. These authors inferred that the H$_2$S$_2$ (or its precursor sulfur-bearing species) could have
been present during the quartz crystallization, as the H$_2$S$_2$-related Raman bands could be detected in sulfur-free CH$_4$—H$_2$S inclusions [4].

The coexistence of hydrogen sulfide and elemental sulfur is also a common characteristic of marble-hosted rubies from Central and Southeast Asia [5]. However, in addition to the Yuanjiang rubies, H$_2$S$_2$ has not been identified in any other deposits [2,5]. The Asian marble-hosted rubies formed under similar geological processes during the Cenozoic Himalayan orogenesis. Garnier et al. [15] suggested that they formed under amphibolite facies conditions. Since H$_2$S$_2$ has not been detected in other Asian marble-hosted rubies, it seems reasonable to infer that H$_2$S$_2$ in sulfur-bearing inclusions within Yuanjiang rubies does not result from a post-entrapment reaction between the elemental sulfur and the coexisting H$_2$S-bearing fluid. On the contrary, a most plausible explanation for its occurrence could be that H$_2$S$_2$ or its precursor mixed-valence sulfur-bearing species should have been present during the crystallization of Yuanjiang rubies.

Though the ASRR shear belt probably activated as a lithosphere-scale ductile strike-slip shear zone in the Tertiary [16], mantle-derived rocks have not been found within the mylonitic gneisses shear belt thus far [11]. Available evidence obtained from biotite-garnet and plagioclase-garnet thermobarometers suggests that the exhumed terrane along this belt corresponds to a paleo-depth of about 18 km [11]. The protoliths of the Ailao Shan high-grade metamorphic massif are of sedimentary origin [17]. Assuming a crustal thickness of 35–40 km for the study area [11,18], these observations suggest that rubies have a mid-crustal origin. As the sulfane composition in fluid inclusions is speculated to be enclosed during ruby growth (sulfane-bearing fluid inclusions are primary and pseudo-secondary), it must also have a mid-crustal origin.

### 5.3. Favorable Geological Condition for the Generation of Sulfanes

At both localities where H$_2$S$_2$ was detected (Bastar Craton in India and Ailao Shan metamorphic massif in China), rock types and metamorphic grades are comparable. The first documented sulfane-bearing fluid inclusions from India were found in metasedimentary sequence, which predominates in gneisses and contains minor schists, quartzites, marbles, and metabasalts. Peak metamorphism in this region occurred at amphibolite facies [4]. As for the Yuanjiang area, peak P-T conditions also correspond to amphibolite facies [11] and the exposed rocks are mainly composed of gneisses accompanied with some migmatites and granitic pods, and interlayers of metapelites, amphibolite, and marble. Comparison of the lithologies and P-T conditions between these two examples suggests amphibolite facies metamorphism of sedimentary sequence that deposited in a continental platform setting is favorable for the generation of H$_2$S$_2$. Even in similar-facies metamorphic terranes, the conditions can be very different in terms of fluid compositions, retrograde paths, and timeframes, etc., which may explain the H$_2$S$_2$ deficiency in fluid inclusions from other rubies in metasedimentary sequences. This explains the singularity of the localities where sulfanes have been found. Unfortunately, the specific mechanism for the generation of sulfanes in the above two occurrences remains unclear.

### 5.4. Implication for Thermochemical Stability of Sulfanes

If the explanation about H$_2$S$_2$ (or its precursor S-H-bearing species) formation was plausible, this would imply that the sulfanes or its precursor mixed-valence sulfur-bearing species could remain stable in metamorphic conditions of Yuanjiang corundum formation. The P-T estimates for the Yuanjiang ruby crystallization remain unconstrained at present. At another marble-hosted ruby deposit in the ASRR metamorphic belt, located in the Day Nui Con Voi range in Vietnam, the ruby mineralization temperature is calculated in the range of 600 to 625 °C by calcite-graphite isotopic thermometry [19]. Here, we assumed that the Yuanjiang rubies formed at a similar temperature range. This high formation temperature of Yuanjiang ruby implies that the thermochemical stability of sulfanes or its precursor S-H-bearing species could be as high as 600 °C. Thus far, there are few experiments and thermodynamic calculations to establish the stability limits of sulfanes in
natural fluid systems. Migdisov et al. [14] have experimentally shown that sulfanes remain stable at temperatures of 200–290 °C. However, the wider-range P-T stability of sulfanes is still unavailable. Nevertheless, based on their inference that H$_2$S$_2$ or its precursor sulfur-bearing compounds could have been present during the quartz crystallization, Hurai et al. [4] suggested high-temperature stability for mixed-valence sulfur-bearing species (>400 °C as estimated by graphite thermometer using Raman spectrometric method). If our inference is correct, it considerably expands the temperature stability of H$_2$S$_2$ or its precursor species. The presence of even traces of water tend to contribute to the decomposition of H$_2$S$_2$ [3]; though sulfanes and water were simultaneously present during entrapment in Yuanjiang rubies, water had been exhausted completely by reactions with the host corundum to produce diaspore along the cooling P-T path [10], making the preservation of sulfanes possible.

5.5. Geochemical Implications

The geological features at Yuanjiang share some similarities with those observed at the Bastar Craton region of India, where the sulfane-bearing inclusions were first documented. For example, they both occur in the high-grade gneissic terrane. Moreover, Cr-rich minerals were found in both of these locations, as represented by Cr-rich mica (up to 1.8 wt.% Cr$_2$O$_3$) and Cr-chlorite in Bastar [4], and ruby in Yuanjiang (up to 1.1 wt.% Cr$_2$O$_3$) [20], respectively.

Chromium, a siderophile element essential for the formation of rubies, is commonly considered as immobile in most geological fluids [21]. Short-distance mobilization of Cr has been proposed by molten salts during the ruby formation [5]. In other cases, it has been demonstrated that the polysulfide complexation in hydrothermal fluids plays an important role in transporting siderophile elements [22]. If our inference that sulfanes or its precursor S-H-bearing species were present in the course of Yuanjiang ruby precipitation is valid, then the role of sulfanes (or its precursor) in Cr mobilization could be expected.

6. Conclusions

The identification of H$_2$S$_2$ fluid composition in rubies from Yuanjiang marbles is the second example of sulfane-bearing fluid inclusions after quartz hosted H$_2$S$_2$-bearing fluids, confirming the presence of sulfane in geological fluids at elevated P-T conditions. Our observations suggest that H$_2$S$_2$ or its precursor S-H-bearing species should have been present during the Yuanjiang ruby precipitation. This further implies that sulfanes can survive high P-T and probably result from the amphibolite facies metamorphism of the sedimentary sequence. The presence of reduced volatiles such as H$_2$S also provides robust evidence for corundum crystallization in a reduced condition [13], aligning with the presence of H$_2$S and CH$_4$ in the fluid phase of Yuanjiang rubies.

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