Solubility of Na₂SO₄ in silica-saturated solutions: Implications for REE mineralization

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Abstract

Sulfate is traditionally considered to have retrograde solubility in aqueous solutions. However, our recent hydrothermal diamond-anvil cell (HDAC) experiments have shown that the solubility of Na₂SO₄ changes from retrograde to prograde in the presence of silica, leading to the formation of sulfate-rich solutions at high temperatures, in line with observations on natural geofluids. In this study, we use synthetic inclusions of fused silica capillary capsules containing saturated Na₂SO₄ solutions and Na2SO4 crystals to quantitatively investigate the solubility of Na2SO4 at different temperatures in the Na₂SO₄-SiO₂-H₂O system. Sulfate concentrations were measured using Raman spectroscopy and calibrated using Cs₂SO₄ solutions with known concentrations. The solubility of crystalline Na₂SO₄ dropped slightly when heated from 50 to 225 °C and dramatically from 225 to 313 °C. At 313 °C, the Na₂SO₄ crystals began to melt, forming immiscible sulfate melt coexisting with the aqueous solution, with or without solid Na₂SO₄. With the formation of sulfate melt, the solubility of Na₂SO₄ was reversed to prograde (i.e., solubility increased considerably with increasing temperatures). The solubility of Na_2SO_4 in the measured solution was significantly higher than that predicted in the absence of SiO₂ over the entire temperature range (except for temperatures around 313 °C). This indicates that the presence of SiO_2 greatly changes the dissolution behavior of Na_2SO_4 , which may be caused by the formation of a sulfate-silicate intermediates such as Si(OH)₄SO₄²⁻. Considering that most crustal fluids are silicasaturated, the solubility curve of Na₂SO₄ obtained in this study can better reflect the characteristics of geofluids when compared to that of Na₂SO₄-H₂O binary system. At temperatures of 313–425 °C, the solubility of Na₂SO₄ increases with temperature following the function $C_{\text{sulfate}} = -3173.7/T + 5.9301$, where C_{sulfate} and T represent the solubility of Na₂SO₄ in mol/kg H₂O and temperature in Kelvin, respectively. As an application, this temperature-solubility relationship can be used to evaluate the sulfate contents in fluid inclusions that contain sulfate daughter minerals, based on the temperature of sulfate disappearance obtained from microthermometric analysis. The sulfate concentrations of the ore-forming fluids of the giant Maoniuping carbonatite-related rare earth element (REE) deposit (southwest China) were calculated to be 4.67-4.81 m (mol/kg H₂O). These sulfate concentrations were then used as internal standards to calibrate the previously reported semi-quantitative results of laser ablation-inductively coupled plasma-mass spectrometry (LA-ICP-MS) analysis of REE-forming stage fluid inclusions at this deposit. The calculated Ce concentrations in the REE-mineralizing fluid range from 0.42 to 0.49 wt%. The high fluid REE contents suggest that the sulfate-rich fluids are ideal solvents for REE transport. A mass-balance calculation was carried out to evaluate the minimal volume of carbonatite melt that was required for the formation of the giant Maoniuping REE deposit. The result indicates that the carbonatite dikes in the mining area are enough to provide the required fluids and metals, and thus a deep-seated magma chamber is not necessary for ore formation.

Keywords: Na₂SO₄ solubility, silica saturation, rare earth element, mineralizing fluid, FSCC

INTRODUCTION

Sulfate is the second most abundant solute in seawater and widely exists in the crustal mantle fluids and extraterrestrial aqueous environments, such as the surface of Mars and Europa (Chipera and Vaniman 2007; McCord et al. 1998). Sulfate is abundant in some hydrothermal systems related to ore formation, such as volcanogenic massive sulfide (VMS) deposits (Yang et al. 2018) and copper porphyry deposits (Sun et al.

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2013). Furthermore, syn-ore fluid inclusions containing sulfate daughter minerals have been reported in carbonatite-related rare earth element (REE) deposits, such as the world-class Bayan Obo in northern China (Xie et al. 2019) and the Maoniuping and Lizhuang in southwestern China (Xie et al. 2015). However, the presence of sulfate-rich geofluids contradicts the knowledge that sulfate salts have retrograde solubility, which would lead to low concentrations of dissolved sulfate in high-temperature solutions (Seward et al. 2014).

Our recent study shows that the presence of dissolved silica is a key to changing the temperature dependence of sulfate solubility. In the presence of quartz and water (SiO₂-saturated fluid), Na₂SO₄ crystal will melt at \sim 300 °C, forming immiscible sulfate melt in coexistence with the aqueous phase. Once the melt forms, the solubility of Na₂SO₄ changes from retrograde to prograde, leading to the formation of sulfate-rich fluids at high temperatures (Cui et al. 2020). However, the solubility of Na₂SO₄ in the Na₂SO₄-SiO₂-H₂O system has not yet been quantified.

Sulfate-rich fluids are in close association with REE mineralization, and sulfate minerals are commonly observed in carbonatite-related REE deposits (Fan et al. 2016; Xie et al. 2015; Xu et al. 2015; Yu et al. 2014). In the current paradigm of REE mineralization, sulfate ion is considered to be one of the potential ligands for REE migration (Migdisov et al. 2016). Moreover, previous laser ablation-inductively coupled plasmamass spectrometry (LA-ICP-MS) analyses on individual fluid inclusions at the Maoniuping and Lizhuang deposits showed strong REE signals in the sulfate-rich ore-fluids (Xie et al. 2015), manifesting the great ability of REE transport in sulfaterich fluids. However, the REE contents in these inclusions were not quantified due to the lack of available internal standards to calibrate the ICP-MS signals. The conventional approach to calibrate the trace element contents of inclusions is to use the concentration of dissolved NaCl as the internal standards, which can be obtained via the microthermometric analysis (Heinrich et al. 2003). However, this microthermometric method is not feasible for sulfate-predominated fluid systems because the equation of states to calculate the concentrations of dissolved sulfate is lacking. Instead, the Ce/K ratio of the mineralization stage fluid inclusions hosted in fluorite was reported by Xie et al. (2015). If the temperature dependence of sulfate solubility is known, the sulfate contents in inclusions can be obtained via microthermometric methods, according to the temperature of the disappearance of sulfate daughter minerals. The calculated sulfate concentrations can then be applied to calibrate the ICP-MS signals to obtain the REE concentrations in the fluids.

Experiments on mineral solubility are mostly conducted using hydrothermal autoclaves or piston-cylinder apparatus (Laudise et al. 1987; Tropper and Manning 2007), and the amounts of dissolved minerals can be analyzed using two major approaches. The first is to analyze the compositions of the separated (or quenched) solutions (e.g., Laudise et al. 1987) or weight losses of the reactants (e.g., Tropper and Manning 2007) after the devices are cooled down. If the solution is not quenchable, a design is required to separate solid reactants from the solutions at high temperatures, to prevent re-precipitation during cooling (e.g., double capsule method) (Manning and Boettcher 1994). The second approach takes advantage of in situ spectroscopic analyses (mostly with synchrotron X-rays) to directly analyze concentrations of dissolved elements in the aqueous phase at high temperatures (e.g., Migdisov et al. 2016). However, neither of these methods can be applied to determine the solubility of Na₂SO₄ in silica-saturated solutions, due to the unusual high-temperature phase behaviors of the system. As revealed by our previous study (Cui et al. 2020), Na₂SO₄ crystals will melt at ~300 °C and thus the liquid sulfate cannot be physically separated from the aqueous solution. In addition, dissolved Na₂SO₄ will precipitate rapidly during cooling, and the solution is thus unquenchable. Furthermore, due to the low X-ray energy of sulfur, its signal would be so weak that the quantitative measurement of the sulfate concentration is not feasible using synchrotron X-ray techniques. Raman spectra provide an ideal solution for the quantitative measurement of sulfate ions (Schmidt and Seward 2017; Qiu et al. 2020), and fused silica capillary capsules (FSCCs) (Chou et al. 2008) are suitable for high-temperature in situ Raman observations and can provide the silica-saturated condition.

In this study, fused silica capillary tubing was used to create synthetic fluid inclusions containing Na_2SO_4 -supersaturated fluid system. The solubility of Na_2SO_4 was determined by in situ Raman spectroscopy at varying temperatures in the Na_2SO_4 -SiO₂-H₂O system. Using the temperature-concentration relationship, we calculated the sulfate concentrations in the syn-ore fluoritehosted fluid inclusions at the Maoniuping deposit and then used the concentrations to evaluate the REE contents in the fluid.

MATERIALS AND METHODS

Experimental designs for measuring the Na₂SO₄ solubility in a silica-saturated system

Sample preparation. FSCCs containing standard Cs_2SO_4 solutions and the Na_2SO_4 -oversaturated system (solid Na_2SO_4 + saturated solution) were prepared using round cross-section fused silica capillary tubing with 100 µm of inner diameter and 365 µm of outer diameter (Polymicro Technologies, U.S.A.) (Wang et al. 2013). The polyimide coating of the tubing was burned off, and the tubing was cleaned with ethyl alcohol (analytical grade reagent) before sample loading. The procedures of sealing the capillary tubes were described in detail by Chou et al. (2008) and Wang et al. (2013).

Cesium sulfate solutions were prepared from distilled deionized water and Cs_2SO_4 crystals (analytical grade reagent, 99.9% in purity; Shanghai Aladdin Bio-Chem Technology Co., Ltd.), and loaded into fused silica capillary tubing as standard samples. The concentrations of standard samples were set as 0.10, 0.49, 0.87, 1.91, and 2.90 *m* (molality, mol/kg H₂O). For the Na₂SO₄-bearing experimental samples, Na₂SO₄-saturated solutions and excess analytical grade Na₂SO₄ crystals (>99.0% in purity) were loaded into the fused silica capillary tubing successively. First, Na₂SO₄-saturated solution was loaded into the tubing, as described in Wang et al. (2013). Then several Na₂SO₄ crystals were picked using a needle and loaded into the tubes. Finally, the sealed capsules were centrifuged to ensure that all the solids are at one end of the capsule (Fig. 1a). This allowed the solution to remain saturated with respect to Na₂SO₄ as temperatures increases from 50 to 425 °C so that the measured sulfate concentrations can represent the solubility of Na₂SO₄.

The degree of filling (i.e., the volumetric ratio between the liquid phase and the entire sample chamber) of standard samples were ~0.6. Two experimental samples were prepared with degrees of filling of 0.82 and 0.52. The former was heated from 50 to 425 °C, while the latter to only 313 °C to avoid explosion due to the high internal pressure. The prepared FSCCs were <1.5 cm in lengths so that they could be placed in the sample chamber of the Linkam THMSG 600 heating–cooling stage. The scaled FSCCs were washed in an ultrasonic cleaner before Raman spectroscopic analyses.

Acquisition and processing of Raman spectra. Fused silica capillary capsules were placed on a Linkam THMSG 600 heating-cooling stage for temperature control with an accuracy of ±0.1 °C, and the heating rate was set at 10 °C/min. To investigate the times that are required to achieve chemical equilibrium, three FSCCs containing solid Na2SO4 and saturated solution were maintained at 140, 330, and 400 °C. For each sample, Raman spectra of the solution were recorded repeatedly until the equilibrium of the system was acquired. Specifically, the times that are needed for getting constant peak area ratios between the stretching vibration band of dissolved sulfate $[v_1(SO_4^{2-})]$ and the OH stretching vibration band of water $[v_s(H_2O)]$ ($A_{sulfate}/A_{water}$) of the solutions were estimated (Fig. 2). It is revealed that Na2SO4 saturation can be reached within 400 s, considering the uncertainties of the measurements (Fig. 2). Therefore, all the samples were held for 10 min at each temperature before the spectral acquisition. For standard samples, Raman spectra were collected at temperature intervals of 50 °C for the temperature range of 50-450 °C. For experimental samples, the intervals were 50 and 25 °C for 50-200 and 200-450 °C, respectively. The Raman spectra for experimental samples



FIGURE 1. (a) Schematic diagram of the experimental method in FSCC experiments. (b) The microscopic photo of the experimental sample at 100 and 350 °C along with Raman spectra of the saturated solution at varying temperatures. Abbreviations: Aq = aqueous solution, S = solid Na₂SO₄, V = vapor phase, M = sulfate melt. (Color online.)

were shown in Figure 1b and those of standard samples in Supplemental Material¹

A Jobin Yvon/Horiba confocal LabRAM HR Evolution micro-Raman system at Peking University was used for Raman spectroscopic analyses. The ICS corrected Raman spectra were acquired with a 532 nm laser excitation (double frequency Nd: YAG), a 20× long-working distance objective with a 0.25 numerical aperture, and a 600 groove/mm grating. The laser power was 100 mW at the source. The confocal hole was set at 180 μ m, and the corresponding spectral pixel resolution was ~±2 cm⁻¹. Raman spectra in the ranges of 800–1200 and 2500–4000 cm⁻¹ were collected for the $v_1(SO_4^2)$ and $v_s(H_2O)$ bands, respectively, with an acquisition time of 20 s for each spectrum. The focus positions of the laser for all the samples, including experimental and standard samples, were set at the same depth by adjusting the focus distance till the boundaries of the cells are mostly clear (i.e., focus in the middle of the capillary). This can guarantee that the obtained Raman intensity is comparable among different samples (Schmidt and Seward 2017).

All Raman spectra measured in this study were processed using the PeakFit v.4.11 software package (SYSAT Software Inc., U.S.A.) (Yuan et al. 2016). A linear baseline was subtracted after the spectral region of interest was selected. Then the spectra of $v_1(SO_i^2)$ and $v_s(H_2O)$ bands were fitted using a Gaussian + Lorentzian function, and the peak areas of each band were obtained from the fitting results. For the $v_s(H_2O)$ band, the spectra were fitted with three sub-bands at ~3194, 3247, and 3444 cm⁻¹. The $v_1(SO_i^2)$ spectra were fitted as one component (Gaussian + Lorentzian function). The error of each fitted component was given by PeakFit.

The total error of the peak area ratio, A_{sulfac}/A_{water}, was calculated based on the errors of each fitted component and the rules of error propagation (Taylor 1996).

Calibration of sulfate concentrations in experimental samples. The integrated $v_s(H_2O)$ intensity was used as the internal standard to normalize the peak area of $v_1(SO_4^2)$ band. In theory, the H_2O -normalized Raman peak area of SO_4^2 (i.e., $A_{sulfate}/A_{sutec}$) should exhibit a linear relationship with the sulfate concentration (Qiu et al. 2020). If the Raman parameters (e.g., laser wavelength, grating notch groove density, cumulative time, laser power, confocal hole size, objective working distance, etc.) remain constant, the relationship between sulfate concentration and H_2O -normalized Raman intensity at a given temperature can be retrieved by measuring the intensities of the $v_1(SO_4^2)$ and $v_s(H_2O)$ of standard samples with known sulfate concentrations. Then the sulfate concentrations in experimental samples can be obtained by plotting their $A_{sulfate}/A_{water}$ ratios on the calibration curve (cf. Pokrovski and Dubessy 2015; Schmidt and Seward 2017).

Cesium sulfate solutions were chosen as the standard due to the high solubility of Cs₂SO₄ at room temperature, and thus standard samples with high sulfate concentrations can be prepared. Furthermore, Cs⁺ is not prone to associate with SO₄⁻ at elevated temperatures [i.e., negligible CsSO₄ and Cs₂SO₄(aq); inferred from the dissociation constant of KSO₄ in Table 4 of Helgeson (1969)], so the concentrations of free SO₄⁻ are well defined (nearly equal to that of total sulfur). After each run of the heating-cooling circle, the Raman spectra of the standard and experimental samples were collected again at room temperature and compared with



FIGURE 2. The investigation of equilibrium time of experimental samples. The peak area ratios, $A_{sulfate}/A_{water}$ for the experimental samples vary with experimental duration. The data suggest that equilibrium was attained after ~400 s at 140 and 330 °C and after ~300 s at 400 °C. Whiskers represent 95% confidence intervals. (Color online.)

those collected at the beginning of the experiment. It is shown that the intensities of the $\nu_1(SO_4^2)$ and $\nu_s(H_2O)$ bands remained constant before and after heating, confirming the functional stability of the Raman spectrometer throughout the experiments and the reliability of the calibration results.

Pressure determination in FSCCs. Pressures inside the experimental samples were calculated using the ISOC program of the FLUIDS v.1 software package (Bakker 2003). Because the EoS (equation of state) of the Na₂SO₄-H₂O system is lacking, that of the NaCl-H₂O system (Zhang and Frantz 1987) was used as an approximation for pressure evaluation. The model NaCl-H₂O solutions were set to have the same molarities and homogenization temperature as those of the Na₂SO₄-H₂O solutions in the experimental samples. Errors in pressure estimation may be introduced by (1) using the EoS of NaCl-H₂O instead of Na₂SO₄-H₂O, and (2) the thermal expansion of the cell at high temperatures (Chou et al. 2008). The latter might lead to the overestimation requires the system to be isochoric (after vapor disappearance). We estimated that the calculated pressures are accurate within a factor of 1–2.

Determination of experimental errors

The errors of the fitted peak areas for one component of $v_1(SO_4^{-1})$ band and three components of $v_s(H_2O)$ band were all directly obtained from PeakFit. According to the rules of error propagation (Taylor 1996), the total error of the peak area of $v_s(H_2O)$ band can be calculated by Equation 1:

$$E_{\rm w \ total} = E_{\rm w1} + E_{\rm w2} + E_{\rm w3} \tag{1}$$

where E_{w1} , E_{w2} , and E_{w3} denote the errors of the three components of v_s (H₂O) band, respectively, while $E_{w \text{ total}}$ is the total error of the peak area of v_s (H₂O). As for the area ratios between v_1 (SO₄²⁻) and v_s (H₂O) bands, the propagated errors of the area ratios can be calculated by Equation 2:

$$(E_{\text{Ratio}} = E_{\text{S}}/A_{\text{sulfate}} + E_{\text{w total}}/A_{\text{water}}) \times A_{\text{normalized}}$$
(2)

where $A_{\text{normalized}}$ is the peak area ratio between $v_1(\text{SO}_4^2)$ and $v_s(\text{H}_2\text{O})$ bands; E_s is the error of the peak area of $v_1(\text{SO}_4^2^-)$; A_{sulfate} and A_{water} are the areas of $v_1(\text{SO}_4^2^-)$ and $v_s(\text{H}_2\text{O})$, respectively; and E_{Ratio} is the propagated error of the area ratios.

Validation of the measured solubility using hydrothermal diamond-anvil cell

A hydrothermal diamond-anvil cell (HDAC) (Bassett et al. 1996) was used to independently measure the solubility of Na₂SO₄ in the presence of quartz or amorphous silica, as a validation of the solubility-temperature relationship obtained using FSCCs. Doubly polished and gold-lined Re gaskets with a 400 µm diameter hole at the center were used to avoid the reaction between rhenium and sulfate ion at high temperatures (Pokrovski and Dubrovinsky 2011). In experimental run 1 (labeled as HDAC_1), Na₂SO₄ solution (2.82 m; undersaturated at room temperature) was loaded in the sample chamber of the HDAC with a piece of silica glass. In experimental run 2 (labeled as HDAC_2), Na₂SO₄ solution (2.50 *m*) was loaded with a gem-grade quartz piece from the Donghai deposit in China. The use of quartz instead of silica glass was to investigate the influence of crystalized/amorphous silica in changing the Na₂SO₄ solubility. When loading the Na₂SO₄ solutions into the sample chambers, the relative humidity in the room was controlled to be >60% to minimize the water loss due to evaporation, which may lead to an elevation in the Na₂SO₄ concentration (e.g., Schmidt 2009).

RESULTS

Construction of the relationship between $A_{sulfate}/A_{water}$ and $C_{sulfate}$

The spectra of the Cs₂SO₄ standard samples were processed to retrieve the relationships between the $A_{\text{sulfate}}/A_{\text{water}}$ ratios and the concentrations of sulfate ($C_{sulfate}$). These relationships vary with temperature and thus were obtained at temperatures from 50 to 450 °C (Fig. 3). The relationships were retrieved as two steps. First, by heating the five standard samples with different concentrations (0.10, 0.49, 0.87, 1.91, and 2.90 m for standard 1-5, respectively), the relationships between the measured $A_{sulfate}/A_{water}$ ratio and experimental temperature were obtained, yielding the temperature dependence of $A_{\text{sulfate}}/A_{\text{water}}$ for solutions with given concentrations (Fig. 3; Table 1). Second, for each temperature of interest, the $A_{sulfate}/A_{water}$ - $C_{sulfate}$ relationships were determined by linear regression with five data points. Data points with low signal-to-noise ratios were excluded (e.g., 200 and 250 °C for standard samples and 200 and 275 °C for experimental samples), and the missing points were obtained from the $A_{\text{sulfate}}/A_{\text{water}}$ -T (temperature) relationship retrieved from the standard samples. In this way, the $A_{\text{sulfate}}/A_{\text{water}}$ - C_{sulfate} relationships were regressed for all temperature conditions with Pearson's correlation coefficients of $R^2 > 0.97$ (Fig. 4).



FIGURE 3. Changes in peak area ratios $A_{sulfate}/A_{water}$ of standard samples with increasing temperature (*T*). The standard samples (standard 1–5) have different concentrations (0.10, 0.49, 0.87, 1.91, and 2.90 *m*) of Cs₂SO₄. The relationships between the measured $A_{sulfate}/A_{water}$ and temperature were regressed. (Color online.)

Sample no.	Concentration (m)	Equation and R ²	
standard 1	0.10	$A_{\text{sulfate}}/A_{\text{water}} = 2 \times 10^{-5}T + 2.7 \times 10^{-3}, R^2 = 0.548$	
standard 2	0.49 A	$_{sulfate}/A_{water} = 4.42 \times 10^{-5}T + 1.72 \times 10^{-2}, R^2 = 0.938$	
standard 3	0.87	$A_{\text{sulfate}}/A_{\text{water}} = 2.62 \times 10^{-7}T^2 - 2.92 \times 10^{-5}T +$	
		3.72×10^{-2} , $R^2 = 0.981$	
standard 4	1.91	$A_{\text{sulfate}}/A_{\text{water}} = 6.76 \times 10^{-9} T^3 - 3.77 \times 10^{-6} T^2 +$	
		$7.01 \times 10^{-4}T + 5.26 \times 10^{-2}$, $R^2 = 0.987$	
standard 5	2.90	$A_{\text{sulfate}}/A_{\text{water}} = 8.50 \times 10^{-9} T^3 - 5.02 \times 10^{-6} T^2 +$	
		$1.01 \times 10^{-3}T + 7.81 \times 10^{-2}$, $R^2 = 0.978$	
<i>Note:</i> $A_{sulfate}/A_{water}$ is the peak area ratio between $v_1(SO_4^{2-})$ and $v_s(H_2O)$ bands and			
T is the temperature in Celsius.			

TABLE 1. The relationships between $A_{sulfate}/A_{water}$ and temperature (7)at different Cs_2SO_4 concentrations

Measurement of the solubility of sulfate in the Na_2SO_4 -SiO₂-H₂O system

Solubility of Na₂SO₄ in the SiO₂-saturated system. Based on the aforementioned $A_{sulfate}/A_{water}$ - $C_{sulfate}$ relationships, the sulfate concentrations in the Na₂SO₄-saturated solutions within two FSCCs were calculated using the obtained $A_{sulfate}/A_{water}$ ratio (Fig. 5; Table 2). Because SO₄²⁻ was the only aqueous sulfur species that could be detected by the Raman spectra and the solution was saturated in Na₂SO₄, the determined SO₄²⁻ concentration can represent the solubility of Na₂SO₄. This result is consistent with Cui et al. (2020), where the pairing between Na⁺ and SO₄²⁻



FIGURE 4. The relationships between the concentration of Cs_2SO_4 ($C_{sulfate}$) and peak area ratio $A_{sulfate}/A_{water}$ at each temperature. These relationships were used to calibrate the concentrations of sulfate in the experimental samples. (Color online.)

cannot be detected in the aqueous phase at high temperatures. The estimated concentrations with increasing temperature were plotted in Figure 5. For comparison, the solubility of Na_2SO_4 in pure water (silica-absent) was also calculated based on available thermodynamic properties using the HCh software package (Shvarov and Bastrakov 1999).

In both samples, Na₂SO₄ exhibited retrograde solubility at temperatures from 50–150 and 225–313 °C, and nearly constant solubility at 150–225 °C (Figs. 5 and 6a). The good reproducibility of the $C_{sulfate}$ -T relationships obtained from the two samples indicates that the measurements were reliable. The Na₂SO₄ solubility in SiO₂-saturated solutions between 50 and 313 °C can be described by a cubic polynomial function with $R^2 = 0.970$ (Eq. 3; Fig. 6a):

$$C_{\text{sulfate}} = -5.4873 \times 10^{-7} \times T^3 + 7.1399 \times 10^{-4} \times T^2 - 3.0661 \times 10^{-1} \times T + 46.251$$
(3)

where T is temperature in Kelvin.

As the temperature further increases, the solubility of Na₂SO₄



FIGURE 5. The solubility of Na_2SO_4 in silica-saturated solutions at temperatures of 50–425 °C. In addition, the solubility curve of Na_2SO_4 in SiO₂-free solutions is also shown for comparison [calculated using the HCh program, Shvarov and Bastrakov (1999)]. Whiskers represent 95% confidence intervals. (Color online.)

TABLE 2. The measured sulfate solubility in SiO₂-saturated Na₂SO₄ solutions (Sample FSCC_1−2) at temperatures from 50 to 425 °C

	725 C				
T (°C)	$A_{\rm sulfate}/A_{\rm water}$	Error for	(<i>m</i>)	Error for	Sample no.
		$A_{\text{sulfate}}/A_{\text{water}}$	C_{sulfate}	$C_{\text{sulfate}}(\pm m)$	
50	0.13	0.02	3.10	0.35	FSCC_1
100	0.13	0.01	2.92	0.26	FSCC_1
150	0.13	0.02	2.78	0.48	FSCC_1
225	0.15	0.01	2.91	0.42	FSCC_1
275	0.13	0.01	2.60	0.11	FSCC_1
300	0.11	0.01	1.65	0.18	FSCC_1
325	0.11	0.02	1.82	0.36	FSCC_1
350	0.14	0.00	2.34	0.05	FSCC_1
375	0.20	0.00	2.89	0.07	FSCC_1
400	0.26	0.01	3.41	0.07	FSCC_1
425	0.35	0.01	3.88	0.09	FSCC_1
50	0.14	0.02	3.22	0.37	FSCC_2
100	0.13	0.01	2.84	0.20	FSCC_2
150	0.13	0.01	2.70	0.20	FSCC_2
250	0.15	0.00	2.71	0.09	FSCC_2
275	0.14	0.01	2.42	0.18	FSCC_2
300	0.10	0.01	1.84	0.16	FSCC_2
313	0.07	0.01	1.25	0.11	FSCC_2



FIGURE 6. The temperature dependence of the Na₂SO₄ solubility in SiO₂-saturated solutions. (**a**) The Na₂SO₄ solubility within the temperature range of 323.15-586.15 K. (**b**) The Na₂SO₄ solubility under high temperatures (>586.15 K). (Color online.)

turned to be prograde (>313 °C; Fig. 5) due to the formation of sulfate melt (see below). The solubility of Na₂SO₄ reaches 3.88 *m* at 425 °C in the SiO₂-saturated system (Fig. 5; Table 2). For comparison, those in SiO₂-absent system remain retrograde throughout the whole temperature range and would decrease to ~0.6 *m* at 425 °C (Fig. 5). Even at temperatures below the melting temperature of Na₂SO₄ (~313 °C), the solubility of Na₂SO₄ in the SiO₂-saturated system is higher than that in the SiO₂-absent system (Fig. 5).

In the case of Na_2SO_4 dissolution at temperatures >313 °C, when the solubility is controlled by the dissolution of sulfate melt, the relationship between Na_2SO_4 solubility and temperature was fitted as Equation 4 with $R^2 = 0.993$ (Fig. 6b),

$$\ln C_{\rm sulfate} = -3173.7/T + 5.9301 \tag{4}$$

where *T* is the temperature in Kelvin and $586.15 \le T \le 698.15$.

Phase transitions of Na₂SO₄-SiO₂-H₂O system. During the initial stage of heating, Na₂SO₄ crystals began to precipitate around the pre-existing solid Na₂SO₄ in FSCCs with increasing temperature. In particular, precipitation of large amounts of solid was observed at temperatures >250 °C. This result is consistent with the measured solubility curve (Fig. 5) in which the solubility of Na₂SO₄ gradually decreased from 3.16 (50 °C) to 2.67 m (250 °C) and then sharply decreased to 1.25 m (313 °C). During heating, Na₂SO₄ crystals underwent two solid-state phase transitions, from type V to III at 200–250 °C and from III to I at 250–300 °C, which are consistent with previously reported experimental results (Cui et al. 2020). Cui et al. (2020) demonstrated that the incipient melting of Na₂SO₄ is coupled with the phase transition from type III to I, and this can be manifested by the solubility curve obtained in this study, which shows a change from retrograde to prograde solubility at \sim 313 °C due to the formation of sulfate melt (Fig. 5).

Validation experiments using HDAC

As temperature increased, the vapor in the HDAC gradually shrank and finally homogenized into a liquid phase at 336 and 300 °C for the two runs of HDAC experiments, respectively (Fig. 7). Upon heating, Na₂SO₄ crystals first precipitated from the solution and then melted at higher temperatures. Finally, the sulfate melts re-dissolved into the aqueous solution at 380 °C (sample HDAC_1) and 351 °C (sample HDAC_2). At these temperatures, the Na₂SO₄ solubility would be estimated as 2.91 and 2.33 *m* based on Equation 4 for samples HDAC_1 and _2, respectively. Comparing with the concentrations of the loaded solutions (2.82 and 2.50 *m* for sample HDAC_1 and _2), the differences are 3.37% and -7.36%, respectively (Table 3).

The pressures at which solid Na₂SO₄ disappeared were estimated to be 56 MPa at 380 °C for sample HDAC_1 and 66 MPa at 351 °C for sample HDAC_2, which are much lower than those in sample FSCC_1 (238 MPa at 351 °C and 298 MPa at 380 °C). The HDAC experiments indicated that the solubility-temperature relationship (Fig. 6b; Eq. 4) could describe the solubility of Na₂SO₄ in SiO₂-saturated solutions with errors of <~8% within the pressure range from 50 to at least 200 MPa (Table 4), regardless of the form of solid SiO₂ (quartz crystal or amorphous silica). Therefore, this relationship can be applied to investigate the Na₂SO₄ solubility in natural SiO₂-saturated crustal geofluids.

DISCUSSION

Influence of SiO₂ on the solubility of Na₂SO₄

Cui et al. (2020) reported that the presence of silica can significantly change the dissolution behaviors of sulfate through the formation of sulfate melt. This accounts for the rapid increase in the solubility of Na₂SO₄ at temperatures of >313 °C (Fig. 5). In this study, we further observed that at lower temperatures, the presence of silica also elevates the solubility of Na₂SO₄ (by ~35% at ~250 °C; Fig. 5) without the formation of sulfate melt (Fig. 5).

The changes in sulfate solubility may be a result of the formation of sulfate-silicic complexes, such as $Si(OH)_4SO_4^{2-}$, in the presence of dissolved silica (Marshall and Chen 1982). Previous studies have shown that the formation of such complexes

TABLE 3. The solubility of Na_2SO_4 in SiO_2 -saturated solutions from HDAC experiments

Sample	Silica	Actual	Calculated	T _{h,tot} ^a	Рь
110.	species	concentration	concentration		
		(<i>m</i>)	(<i>m</i>)	(°C)	(MPa)
HDAC_1	Glassy silica	2.82	2.91	380	56
HDAC_2	Quartz crystal	2.50	2.33	351	66
$\overline{T_{h,tot}}$ = totally homogenization temperature.					

^b P = pressure at $T_{\rm h,tot}$.

TABLE 4. Pressure estimation for FSCCs with different degrees of filling					
Sample no.	Degree of filling	T _{h,vapor} (°C) ^a	P ₄₂₅ (MPa) ^b		
5666 4		222	120		

FSCC_1	0.82	320	139		
FSCC_2	0.52	371	73		
${}^{a}T_{h,vapor}$ = aqueous-vapor homogenization temperature within the FSCC.					
^b P_{425} = the calculated inner pressure of FSCC at 425 °C.					



FIGURE 7. Phase behaviors of Na₂SO₄-SiO₂-H₂O system in an HDAC experiment (sample HDAC_2) at 180, 300, and 351 °C. Abbreviations: Aq = aqueous solution, S = solid Na₂SO₄, V = vapor phase, Qtz = quartz, V = vapor phase. (Color online.)

will increase the solubility of SiO_2 in sulfate-bearing solutions (Kotel'nikova and Kotel'nikov 2010; Wang et al. 2016). Here, we show that the elevation in solubility is mutual: the presence of silica could also enhance the dissolution of Na_2SO_4 and possibly other sulfate salts. Considering that quartz is ubiquitous in crustal rocks, the mobility of sulfate should be higher than previously estimated in crustal geofluids. Aside from the elevated solubility in the presence of silica, another factor that may lead to the formation of sulfate-rich fluids at high temperatures is the liquid-liquid immiscibility of sulfate solutions, which leads to the formation of a dense sulfate-rich liquid phase coexistent with the sulfate-poor solution (Wang et al. 2017; Wan et al. 2017).

Application to determine the fluid composition of oreforming fluids of the carbonatite-related REE deposit

For fluid inclusions containing halite and other daughter minerals, salinities are measured based on the temperatures at which the daughter minerals completely dissolve into the aqueous phase (Potter and Clynne 1978). Salinity calculated using this approach is conventionally expressed as equivalent NaCl concentration because the presence of other electrolytes is inevitable but difficult to be known quantitatively. Similarly, the sulfate concentrations in inclusions that contain sulfate daughter minerals can be estimated using Equation 4 and expressed as equivalent Na₂SO₄ contents, $C_{Na_2SO_4,eqv}$ (in m). This approach was applied to estimate the composition of the fluorite-hosted fluid inclusions from the Maoniuping deposit, the syn-ore mineral assemblage of which is characterized by the coexistence of quartz and REE minerals (mainly bastnaesite) (Xie et al. 2015). Previous studies indicated that the syn-ore fluid inclusions contain large volumes of sulfate daughter minerals and they will first melt and finally disappear at 450-455 °C upon heating (Xie et al. 2015). Based on Equation 4, these microthermometric results yielded sulfate concentrations ($C_{Na_2SO_4,eqv}$) ranging from 4.67 to 4.81 m.

Previous semi-quantitative LA-ICP-MS analyses of individual fluid inclusions (Xie et al. 2015) showed that the ore-fluids are mainly composed of dissolved Na-K-Sr-Ba-sulfates, with Na⁺ and K⁺ predominating over Sr²⁺ and Ba²⁺ [molar (K + Na)/(Sr + Ba) = 20], and Na⁺ over K⁺ (molar Na/K = 1.6). The ore-fluids were approximated as the Na-K-sulfate system in this calculation. The presence of K will lead to a deviation in the sulfate solubility from those calculated from simple Na_2SO_4 solution, but this deviation is not clear at present. We use the following equation to represent the influence of the presence of K:

$$C_{\text{Na2SO4,eqv}} = C_{\text{Na2SO4,real}} + \mathbf{A} \times C_{\text{K2SO4,real}}.$$
(5)

The form of Equation 5 follows that of the well-studied NaCl-KCl-H₂O system (Allan et al. 2005), and C_{Na2SO4,real} and $C_{K_2SO_4,real}$ are the real concentrations of Na₂SO₄ and K₂SO₄ in the fluids, respectively, and A is a factor describing the influence of the presence of K2SO4. For the NaCl-KCl-H2O system, A is usually set as 0.5, indicating that KCl is more soluble than NaCl at the same temperature. However, the exact A value for the Na₂SO₄-K₂SO₄-SiO₂-H₂O system is unknown. Three hypothetical end-member values were used to calculate $C_{\text{Na}_2\text{SO}_4,\text{real}}$ from $C_{\text{Na}_2\text{SO}_4,\text{eqv}}$: (1) A = 0.4 representing that K₂SO₄ is more soluble than Na_2SO_4 ; (2) A = 1 for equally soluble K_2SO_4 and Na_2SO_4 ; and (3) A = 1.2 representing that Na_2SO_4 is more soluble. The real sulfate concentration in the K2SO4-Na2SO4-H2O system should be within the three end-member hypotheses. Given that the Na/K molar ratio is 1.6, these end-member hypotheses yielded C_{Na2SO4,real} ranging from 3.25 to 4.20 m. Based on microscopic observations, Xie et al. (2015) estimated that the inclusions contain 70-75 vol% of daughter minerals (mostly hydrated Na-K-Ca-sulfates). If the daughter minerals are all assumed to be mirabilite (Na₂SO₄·10H₂O), this volume would yield sulfate contents of 3.9 to 4.16 m, close to our estimation. Taking the 1000 × Ce/Na ratio (40.7) obtained from previous LA-ICP-MS results (Xie et al. 2015), the Ce concentrations in the ore-forming fluids were calculated to be 0.42-0.49 wt%.

Volume of carbonatite magma required for the formation of giant REE deposit

Geological and geochemical observations on the worldclass Bayan Obo and the carbonatite-related REE deposits in southwestern China showed that sulfate-rich fluids were responsible for REE mineralization (Xie et al. 2009, 2015, 2019). Experimental studies also indicate that SO_4^2 forms strong bonds with REE³⁺ and is thus an ideal ligand for hydrothermal REE transport (Migdisov et al. 2016; Cui et al. 2020). In this study, it is shown that 0.36–0.42 wt% of dissolved Ce can be

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transported by the natural sulfate-rich fluids. Given that Ce typically contributes half of the total REE resource (ΣREE) of carbonatite-related deposits (e.g., Fan et al. 2016), the oreforming fluid at Maoniuping may have an ΣREE content as high as ~1 wt%, orders of magnitude higher than typical base metalforming fluids (typically 10 to 1000 ppm of dissolved Cu, Pb, or Zn) (Yardley 2005). The high concentration of REEs in the ore-forming fluid implies that large volumes of hydrothermal fluid and carbonate melt may not be necessary for the formation of a giant REE deposit. In the case of the giant Maoniuping REE deposit with a reserve of 3.66×10^4 tons of Ce (He et al. 2018), 9.83×10^6 tons of ore-forming fluids were required to transport the metals, as calculated based on the concentration of Ce in the fluid. Keppler (2003) showed that the water content in carbonatite magma relates well to the pressure of the magma. Given the pressure of the magma intruding at the Maoniuping (100-300 MPa) (Xie et al. 2009), the water content in the carbonatite melt is estimated to be ~15 wt%. Taking the estimated density of the carbonatite rock of 2.78 g/cm3 (cf. the densities of calcite and dolomite) into account, 1.78×10^7 to 2.11×10^7 m³ of carbonatite is required to provide the metals and fluids for the Maoniuping deposit. It should be noted that this volume is the lower limit required for the formation of the REE deposit. First, not all the REEs were partitioned into the aqueous phase during the melt-fluid separation. Second, not all REEs precipitated as ore minerals (e.g., bastnasite), with some existing as solid solutions in refractory minerals (e.g., apatite). Finally, not all REE minerals can be effectively concentrated to form economically significant REE orebodies, because many of them were dispersed and precipitated as sub-economic host rocks with REE anomalies.

Nevertheless, the estimation of the magma volume required for ore formation provides important constraints on the process of REE mineralization. Based on the geological map of the mining area and orebody cross sections (Xie et al. 2009), at least 1.44×10^8 m³ of carbonatite dikes were confirmed in the mine site of the Maoniuping deposit. This volume is nearly six times larger than the minimal magma volume required for REE mineralization. This indicates that the known carbonatite dikes are sufficient to account for the formation of the giant REE deposit and a deep-seated magma chamber is not necessary, consistent with the conclusion by Xie et al. (2015).

IMPLICATIONS

This study, together with our previous research (Cui et al. 2020), shows that the dissolution and melting process of sodium sulfate can be significantly influenced by the presence of trace amounts of silica, which is traditionally regarded as an inert component in the fluid system. Taken the ubiquitous presence of silica in the Earth's crust, this finding implies that our current understandings of the high-temperature hydrothermal behaviors of sulfates and possibly other types of minerals may need to be reevaluated, as dissolved silica can be active in reacting with other ligands. Furthermore, this study implies that the influences of dissolved silica cannot be neglected and should be envisaged for hydrothermal experiments using techniques such as fused silica capillary capsules and synthetic fluid inclusion in quartz (e.g., Kotel'nikova and Kotel'nikov 2010).

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