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Science 331, 1052 (2011);
DOI: 10.1126/science.1199911

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The S<sub>3</sub><sup>-</sup> Ion Is Stable in Geological Fluids at Elevated Temperatures and Pressures

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The chemical speciation of sulfur in geological fluids is a controlling factor in a number of processes on Earth. The two major chemical forms of sulfur in crustal fluids over a wide range of temperature and pressure are believed to be sulfate and sulfide; however, we use in situ Raman spectroscopy to show that the dominant stable form of sulfur in aqueous solution above 250°C and 0.5 gigapascal is the trisulfur ion S<sub>3</sub><sup>-</sup>. The large stability range of S<sub>3</sub><sup>-</sup> enables efficient transport and concentration of sulfur and gold by geological fluids in deep metamorphic and subduction-zone settings. Furthermore, the formation of S<sub>3</sub><sup>-</sup> requires a revision of sulfur isotope–fractionation models between sulfides and sulfates in natural fluids.

Sulfur participates in numerous inorganic and biological reactions over a wide range of temperatures and pressures (1–3), during which it often exhibits large fractionations among its stable isotopes (4–7) and transforms between its multiple valence states and chemical forms. Earth’s surface and shallow crust are characterized by a large variation in sulfur forms, with sulfide (S<sub>2</sub><sup>-</sup>; H₂S/HS⁻/S<sup>-</sup>) and sulfate (S<sub>6</sub><sup>+</sup>; HSO₄⁻/SO₄<sup>2-</sup>) being the most common. Other sulfur species (for instance, elemental sulfur, sulfite, thiosulfate, polysulfides, polythionates, and organic sulfur) form as reaction intermediates in the sulfate-sulfide redox cycle but are relatively minor or thermodynamically metastable at near-surface conditions (1, 2). The sulfur speciation in deep and hot fluids from hydrothermal-magmatic systems and subduction zones hosting the major part of metallic resources on Earth (3) is far less constrained. This is due to intrinsic difficulties in studying such fluid systems, which are strongly sensitive to temperature, pressure, and redox conditions. For example, sulfur solubility in water increases by about five orders of magnitude over the hydrothermal temperature range (20° to 500°C) (8, 9). Intermediate-valence species, such as SO₃<sup>2-</sup>, that are potentially important in shallow magmatic-hydrothermal systems (3) tend to disproportionate rapidly into sulfate and hydrogen sulfide (with or without native sulfur) in the aqueous fluid or vapor phase on cooling (8–11). The rates of chemical and isotopic equilibration between sulfates and sulfides in an acidic aqueous solution at supercritical temperatures.

Fig. 1. (A) Raman spectra of an aqueous solution of 1 mol/kg potassium thiosulfate (K₂S₂O₃) at equilibrium at temperatures ranging from 150° to 350°C and pressures of 1.5 ± 0.3 GPa at the S-S and S-O bond-vibration region below 1250 cm⁻¹. Spectra at 150° to 250°C are magnified by a factor of 4 to emphasize small bands. (B) Raman spectra showing the S-H bands of H₂S and HS⁻. Spectra are normalized to 30-s acquisition time and offset vertically for clarity. Vertical dashed lines denote the vibration modes (ν = stretching, δ = bending) and positions of major Raman peaks for the indicated species; stars show the bands of diamond. The frequency shifts of the bands with temperature are typical for aqueous species. Spectra demonstrate the reversible formation of the S<sub>3</sub><sup>-</sup> ion at 300° and 350°C at the expense of sulfate and sulfide, which are stable at lower temperatures (13).

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atures are too fast (4) to allow a magmatic-hydrothermal fluid to preserve its original sulfur speciation on cooling. Thus, most aqueous fluids quenched or trapped as inclusions in minerals, both in nature and in the laboratory, contain almost exclusively sulfate and sulfide (2, 11, 12) and may not reflect the true speciation at high temperature and pressure.

Because of these limitations, the thermodynamic and kinetic properties and isotope fractionation could be investigated in detail only for the two aqueous end members, sulfate and sulfide, at temperature and pressure conditions of shallow magmatic-hydrothermal systems (2–4, 8). Together with chloride, these forms are regarded as the major transporting agents of metals in hydrothermal and metamorphic ore-bearing fluids (2, 3, 11, 12) and as responsible for sulfur isotope records in sulfide and sulfate minerals formed (2, 4). However, this simplicity of sulfur aqueous chemistry may only be apparent because of the lack of in situ approaches for analyzing geological fluids at elevated temperatures and pressures.

To fill this gap, we employed in situ Raman spectroscopy in a diamond anvil cell at 25° to 450°C and 0.5 to 3.5 GPa to measure the identity and stability of S-bearing species in model aqueous solutions of thiosulfate and sulfur across a range of acidity (1 < pH < 7) and of sulfur concentration (0.6 to 6 weight percent (wt %)) that are representative of crustal and subduction-zone fluids. These experimental conditions provide acidity and redox control on the system through the sulfate-sulfide equilibrium and can fix oxygen fugacity close to the magnetite-hematite mineral buffer. They also allow for the identification of the dominant S species directly from their characteristic Raman spectra (13).

Our data indicate the preponderance of sulfate and sulfide in solution at < 250°C and 0.5 to 3.5 GPa (Fig. 1), in agreement with thermodynamic predictions (fig. S1) (2, 4, 8, 12). In contrast, at higher temperatures, we systematically found that the dominant form of sulfur was the trisulfur S₃⁻ radical ion (Figs. 1 and 2). This ion is characterized by S-S symmetric bending (δ) and stretching (ν) modes at 240 ± 5 and 538 ± 5 cm⁻¹, respectively, and corresponding overtones δ₊ νₛ, 2νₛ, 3νₛ, and 4νₛ due to the Raman resonance phenomenon of S₃⁻ at the wavelength of the He-Ne laser (14–19). Kinetic and heating-cooling measurements (15) show that this species forms rapidly and reversibly at the expense of sulfate and sulfide (fths. S4 and S5). Owing to the radical nature of S₃⁻ that allows fast electron transfer, its presence accelerates the attainment of the sulfate-sulfide equilibrium at moderate temperatures (250° to 300°C) and near-neutral pH (6 to 8). Our findings demonstrate that S₃⁻ is a dominant, thermodynamically stable aqueous sulfur form at least in the range 250° to 450°C and 0.5 to 3.5 GPa.

The fast breakdown of S₃⁻ to sulfate and sulfide (with or without native sulfur) below 250°C explains the absence of reports of S₃⁻ in numerous studies employing fluid sampling or quenching techniques (4, 9, 11). The few in situ studies conducted so far using ultraviolet-visible (20) and Raman spectroscopy (21) in sulfur-bearing aqueous solutions at similar temperatures (150° to 500°C), but at much lower pressures (~0.1 GPa), may have erroneously attributed the S₃⁻ spectral pattern to other species. Compared with those
data, our high-pressure results indicate a larger $S_\text{3}^\text{−}$ stability domain in temperature, acidity, and sulfur concentration. This pressure-driven stabilization in dense aqueous solution may be due to solvation of $S_\text{3}^\text{−}$ by water molecules yielding energetically favorable coordination geometries, similar to those in zeolite cages of ultramarnine-type minerals in which tetrahedral coordination of $S_\text{3}^\text{−}$ by Na$^\text{+}$ strongly stabilizes this radical ion (18). The structure of $S_\text{3}^\text{−}$ with an S–S–S angle of 103° (17), close to that of H–O–H in H$_2$O, may favor its incorporation into the tetrahedral network of hydrogen bonds formed by water molecules in the dense liquid phase at elevated pressure.

We estimated the $S_\text{3}^\text{−}$ concentrations from comparison of the Raman intensities of the nonresonant sulfate and sulfide species and using the mass balance of the total dissolved sulfur and constraints of the relative fractions of $S_\text{6}^{\text{2−}}$ and $S_\text{2}^{\text{−}}$ forms imposed by the system composition (13). These data, combined with the robust thermodynamic properties for sulfate and sulfide species (8, 12), allow derivation of the formation constants (22) and prediction of the $S_\text{3}^\text{−}$ equilibrium amounts in natural fluids (Fig. 3). In an aqueous solution at 350°C and 0.5 to 1.5 GPa with 1 wt % S, which is a common S concentration for Au-Cu–bearing fluids in porphyry systems (3, 23) and metamorphic fluids generated by pyrite breakdown (24), $S_\text{3}^\text{−}$ accounts for a major part of dissolved sulfur (>50 to 95%) in a wide range of pH (2 to 7) at oxygen fugacity of the hematite-magnetite (HM) buffer. In more reducing environments, like those of the mantle-crust boundary or serpentinitization processes in subduction zones, approximated by the nickel-nickel oxide or quartz-fayalite-magnetite (QFM) equilibria, the $S_\text{3}^\text{−}$ ion attains amounts comparable with H$_2$S in fluids with ≥1000 parts per million (ppm) of dissolved S at near-neutral pH.

The $S_\text{3}^\text{−}$ ion has been studied extensively at low-pressure conditions in nonaqueous chemical systems such as sulfur-bearing organic solvents, alkali halide polysulfide solids and melts, S-doped aluminoborosilicate glasses, ultramarine pigments, and zeolite minerals such as lapis lazuli (14–19). However, the formation of $S_\text{3}^\text{−}$ also has consequences for sulfur geochemistry and metal transport in high-temperature and pressure crustal and mantle fluids. First, the formation of $S_\text{3}^\text{−}$ at the expense of H$_2$S/HS$^\text{−}$ and SO$_2^{\text{2−}}$ (Fig. 3) will enhance the sulfur mobility in the fluid phase by reducing the amount of sulfur retained in pyrite, pyrrhotite, anhydrite, and barite, the major sulfur-bearing minerals. In metamorphic settings of greenschist-amphibolite facies, $S_\text{3}^\text{−}$ will widen the temperature-pressure window of pyrite breakdown leading to generation of sulfur-rich fluids (24). In subduction processes, a part of oceanic crust sulfur is transported along the slab down to zones of dehydration and partial melting (3, 25, 26). If $S_\text{3}^\text{−}$ also forms at magmatic temperatures, as indicated by analyses of synthetic silicate glasses (19), it will enhance the release of sulfur from the FeS-bearing silicate melt into the degassing aqueous fluid phase, contributing to the sulfur flux in volcanic arcs. Evolved arc magmas and associated fluids are often characterized by high oxygen fugacity, up to QFM+2 (3, 26), favoring $S_\text{3}^\text{−}$ stability.

Second, because of its similarity to polysulfide ions like S$_\text{2}^\text{−}$ and S$_\text{4}^\text{2−}$ that form strong complexes with Au in aqueous solution (27), $S_\text{3}^\text{−}$ should exhibit a high affinity for Au and similar metals (Cu, Pt). This ligand will thus efficiently compete with HS$^\text{−}$ and H$_2$S, which are regarded so far as the main transporting agents of gold in hydrothermal fluids (2, 12, 23, 24). Experimental evidence (12, 28) in S-rich solutions at 300°C to 600°C and 0.05 to 0.5 GPa of Au solubilities up to three orders of magnitude higher than can be explained by the known Au(I)-sulfide complexes AuH$(\text{S}_\text{2})^\text{−}$ and Au(S$_\text{2}$)$_\text{2}^\text{−}$ (11, 12) may be due to Au binding with $S_\text{3}^\text{−}$. Optimal conditions for $S_\text{3}^\text{−}$ formation include temperatures around 350°C, redox conditions close to the HM buffer, and dissolved S concentrations above 1000 ppm (Fig. 3). Such conditions are met in porphyry and epithermal Au-Cu deposits associated with active convergent margins on Earth (3, 23). Aqueous $S_\text{3}^\text{−}$ released at high pressures from arc magmas into the fluid phase in such settings may extract Au and Cu from the slab and/or mantle wedge. Upon the fluid ascent and pressure decrease, $S_\text{3}^\text{−}$ will decompose into sulfate and sulfide. This may result in partial gold precipitation via sulfidation reactions with Fe-bearing rocks in porphyry and orogenic deposits (3, 25, 29), whereas some part of Au and associated metals (Cu, Se, Te) may be transported upward to shallower epithermal deposits by the low-density vapor phase as volatile species with H$_2$S and/or SO$_2$ (11, 23). Thus, $S_\text{3}^\text{−}$ may not only enhance the mobility of chalcophile metals at depth, but it may also provide the source of sulfur and metals for the important types of gold deposits in Achaean greenstone belts and porphyry-epithermal systems that imply high gold and sulfur fluxes over a wide temperature-pressure-acidity range (3, 23, 29) favored by the elevated solubility of $S_\text{3}^\text{−}$.

Third, if $S_\text{3}^\text{−}$ is abundant in hydrothermal fluids, it will influence thermodynamic properties of intermediate-valence aqueous sulfur species (8, 11, 12, 20, 24) and kinetic models of reactions between sulfates and sulfides (4, 30) in aqueous solution at high temperatures and pressures. This may, in turn, affect sulfur isotope–fractionation models (4–7, 25, 30), which ignore the formation of $S_\text{3}^\text{−}$ in geological fluids.