Geochemical Characteristics of Rare Earth Elements in the Chaluo Hot Springs in Western Sichuan Province, China

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HIGH-TEMPERATURE HYDROTHERMAL ACTIVITY AREAS IN WESTERN SICHUAN PROVINCE, CHINA ARE IDEAL OBJECTS FOR STUDYING DEEP EARTH SCIENCE, Extreme ecological environments, and comprehensive geothermal utilization. TO UNDERSTAND THE GEOCHEMICAL CHARACTERISTICS OF RARE EARTH ELEMENTS (REEs) IN THE CHALUO HOT SPRINGS IN WESTERN SICHUAN PROVINCE, THE AUTHORS ANALYZED THE COMPOSITION AND FRACTIONATION OF REEs IN THE HOT SPRINGS THROUGH HYDROCHEMICAL ANALYSIS, REE TESTS, AND NORTH AMERICAN SHALE COMPOSITE-NORMALIZED REE PATTERNS. Moreover, the composition and complex species of REEs in the geothermal water in the Chaluo area were determined through calculation and simulation analysis using the Visual MINTEQ 3.0 software. The results are as follows. In terms of hydrochemical type, all geothermal water in the Chaluo area is of the Na-HCO3 type. The cations in the geothermal water are mainly controlled by water-rock interactions and evaporation, the anions are determined by water-rock interactions, and the hydrochemical processes are primarily controlled by the dissolution of silicate minerals. The total REE content of the geothermal water in the Chaluo hot springs is 0.306 ± 0.103 ug/L. It is low compared to the Kangding area and is primarily affected by the reductive dissolution of Fe oxides/hydroxides, followed by pH. The geothermal water in the Chaluo area is rich in light rare Earth elements (LREEs) because of the presence of Fe oxides. It shows positive Eu and Ce anomalies due to the combined effects of the dissolution of Eh and Mn oxides and surface water. Furthermore, the positive Eu anomalies are also caused by the water-rock interactions between the Qugasi Formation and deep geothermal water. Similar to alkaline water bodies, the complex species of REEs in the geothermal water mainly include Ln(CO3)2−, LnCO3+, and LnOH2+, which is caused by the stability constants of complexation reactions.

Keywords: Chaluo hot springs, rare earth elements, North American shale standardized REE distribution patterns, Eu anomaly, complexes of the species
INTRODUCTION

Western Sichuan Province is located at the easternmost end of the Mediterranean-Himalayan geothermal zone and at the boundary of the collision between the Eurasian Plate and the African and Indian plates (Royden et al., 2008; Xu et al., 2011; Figure 1A). This region is mainly composed of three strike-slip faults—the Xianshuihe, Ganzi-Litang, and Jinsihaijiang faults (Dewey et al., 1988; Xu et al., 2005; Yan and Lin, 2015), the Songpan-Ganzi fold belt, and the Yidun arc (SBGMR, 1991; Xu et al., 1992; Zhang et al., 2013; Tang et al., 2017; Figure 1B). It is characterized by strong neotectonic movements, frequent earthquakes, active hot springs, and geothermal anomalies (Tang et al., 2017; Zhang Jian. et al., 2017). Statistics reveal that 248 hot springs are distributed in this region (Luo, 1994). Similar to the Yangbajing geothermal field in Tibet and the Rehai geothermal field in Tengchong City, Yunnan Province (Tong et al., 1981; Tong and Zhang, 1989), the high-temperature geothermal resources in western Sichuan Province have various geothermal manifestations including geysers, boiling springs, and boiling spouters, hydrothermal explosions, and hydrothermal alterations (Fan et al., 2019; Figure 1B). The hot springs in western Sichuan Province are mainly controlled by the Jinsihaijiang, Dege-Xiangcheng, Ganzi-Litang, and Xianshuihe faults in the giant structure in the shape of Chinese character ‘‘回’’, which reach as deep as the Moho (Zhang Jian. et al., 2017; Figure 1B). Medium- and high-temperature hot springs are intensively exposed in the Kangding, Batang, and Litang areas. They exhibit strong hydrothermal activities, bear rich high-temperature thermal energy, and therefore have considerable potential for geothermal resource development (Fu and Yin, 2009). These high-temperature hydrothermal activity areas, which are ideal for understanding deep Earth science, extreme ecological environments, and the comprehensive utilization of geothermal energy, have attracted the attention of domestic and foreign experts and researchers in geothermal, geological, geochemical, and ecological fields.

Previous hydrogeochemical studies on the hot spring system in western Sichuan Province have mainly focused on the material origin and the recharge, runoff, and discharge conditions of hot springs, the characteristics and genetic mechanisms of geothermal reservoirs, and the structure, formation, and evolution of crust-mantle heat flow (Zhang Jian. et al., 2017; Yuan et al., 2017; Zhang Y. et al., 2018; Li et al., 2018; Zhang et al., 2019; Cao et al., 2021). Zhao J. Y. et al. (2019) analyzed the evolution of geothermal water in the Batang area by calculating the thermal storage temperature, cold water mixing ratio, and thermal circulation depth. Cao et al. (2021) studied the recharge sources, recharge elevation, geothermal water circulation depth, and geothermal storage temperature of the Chaluo geysers. Yuan et al. (2017) concluded that the differences in hydrochemical characteristics in western Sichuan Province are primarily caused by the dissolution and precipitation of minerals and cation exchanges. Previous research methods are mostly limited to traditional water chemistry and hydrogen and oxygen isotopes, and lack of combined analyses of macronutrients, trace elements and rare Earth elements, which make it difficult to accurately reflect the complex hydrogeochemical evolution of water-rock processes in deep and large fractured geothermal systems. Therefore, the combined hydrogeochemical and rare-earth element analyses are used in this study to provide a more detailed picture of the water-rock interaction in hot springs.

REEs consist of a group of elements with similar physicochemical properties (Hederson, 1984). They generally do not break in geochemical processes and thus have been widely used to trace the processes such as water-rock interactions in surface water (Goldstein and Jacobsen, 1988; Ménager et al., 1992; Johannesson et al., 1997; Dia et al., 2000; Göb et al., 2013). However, there are relatively few studies on the application of REEs in geothermal systems (Möller, 2000; Wood and Shannon, 2003; Sanada et al., 2006). Since REEs serve as a valuable tool for retrospectively studying the origin of geothermal fluids, the study of REEs is fundamental to understanding fluid-rock systems. Therefore, it is necessary to study the geochemical characteristics of REEs in geothermal fluids to improve the understanding of the behavior of geothermal fluids. This study analyzed the composition, fractionation, and key affecting factors of REEs in the geothermal system in the Chaluo hot springs. It will improve the understanding of the genesis of the geothermal system and provide guidance for increasing the development and utilization efficiency of local geothermal resources.

REGIONAL GEOTHERMAL GEOLOGICAL CHARACTERISTICS

The study area is located in the Jinsihaijiang fault and has cut downward to the upper mantle (Zhang Jian. et al., 2017; Zhang, 2020). In this area, the Late Yanshanian-Himalayan faults are superimposed on the earlier tectonic traces and cut through earlier structures. As a result, a checkerboard structural pattern with obvious fracture zones and fracture surfaces has been formed, providing favorable channels for the recharge and runoff of geothermal water. Meanwhile, the checkerboard structural pattern noticeably restricts hydrothermal activities, and the hot springs mostly spread along the fracture zones. By contrast, Indonesian structures in the study area are long and deep. They control the magmatic activities in the area and further control deep hydrothermal convection in the study area. The intersections of these structures of different eras show intensive high-temperature hydrothermal activities. The Rekeng and Reshuitang hot springs in Chaluo Town investigated in this study are also located at the intersections.

The study area lies in a hilly plateau subject to tectonic erosion. The mountains show high-grade slopes, and valleys are wide and gentle, with middle and lower parts downcutting deeply. The height differences between mountain peaks and valley bottom are generally less than 500 m. The rainfall in the Batang area is low and shows an uneven spatial-temporal distribution. It is mainly recharged by precipitation and a small amount of snowmelt.
The study area features stratigraphic discontinuity, where Mesozoic strata dominated by Triassic strata (especially Upper Triassic strata) are widely distributed and outcrops also include Yanshanian biotite granites and monzonitic granites (Figure 1C). Hot springs in the study area are exposed in the Triassic Tumugou and Qugasi formations. The Qugasi Formation
(T₃q) is exposed in the central part of the study area and spreads along Chaluo and Leiwo villages in the NS direction. With the Chalong-Ranbu fracture as a boundary, the Qugasi Formation can be roughly divided into the upper and lower members (Figure 1C). The upper member (T₃q²) has a thickness of 2,295 m and is mainly composed of miscellaneous quartz conglomerates, sandy slates, slates, and dolomitic crystalline tuffs. Meanwhile, lamellibranchia and ammonite fossils occur in this member. The lower member (T₃q¹) has a thickness of 1,558 m. It is composed of light gray—dark gray crystalline tuffs, metamorphic sandstones, and slates and is interbedded with quartz conglomerates. The fossils occurring in this member include lamellibranchia, brachiopoda, and coral. The Tumugou Formation (T₃t), which is widely distributed in the study area, is a set of intermediate-acidic volcanic-sedimentary rock assemblages consisting of neritic-facies volcanic conglomerates, metamorphic sandstones, argillaceous argillites, and crystalline tuffs containing various types of volcanic conglomerates. This formation is in conformable contact with the overlying Lanashan Formation. It can be divided into the upper, middle, and lower members. The upper member (T₃t³) of the formation has a thickness of 972–4,832 m and is composed of sandstones and slates interbedded with intermediate and basal volcanic rocks and crystalline tuffs. The middle member (T₃t²) has a thickness of 579–4,852 m. It is composed of interlayers with different thicknesses consisting of grayish-black lithic feldspar-quartz sandstones and dark slates, interbedded with rhyolites and limestones. The lower member (T₃t¹) has a thickness of 1,830 m and is composed of metamorphic sandstones and argillites. It bears conglomerates in most phase zones and is interbedded with acidic volcanic rocks locally. Moreover, it is in unconformable contact with the underlying strata in local areas.

The Chaluo hot springs refer to a group of hot springs distributed along both sides of the Baqu River in the NE direction from Chaluo Town to the Cuopu Lake in Batang County. They are mainly affected by the Zamagang-Maoyaba, Biezonglongba-Quwengcuo, Chaluo-Saduo, and Chalong-Ranbu faults (Zhang, 2020) and are mostly distributed in the shape of bands along the fault zones in the low-lying areas of the river valley, especially in the area between the Rekeng and Reshuitang areas (Figure 1C).

The hot springs in the Rekeng area (also referred to as the Rekeng hot springs) occur as a spring group. Their water mostly violently boils since its temperature is higher than the local boiling point (89°C; Cao et al., 2021). The hot and boiling springs have temperatures of 80–89°C and their fumaroles have a temperature of up to 99°C. The Rekeng hot springs are composed of more than a thousand springs in an area with a length of about 1,000 m, a width of 100–200 m, and an area of 0.15 km². Their outlets are mainly composed of strongly weathered zones consisting of elluvium-proluvium, pebble gravel, and bedrock. A large amount of calcareous tufa occurs below the outlets (Figure 2A), and its surrounding rocks are oxidized and brown. Meanwhile, a large area of tufa waterfalls is also present below the spring outlets (Figure 2B), with a height of about 50 m and a maximum thickness of 3 m. Local geothermal manifestations include boiling spouters, many boiling springs,
fumaroles, steaming ground, hot springs, and geysers (Figure 2C).

The hot springs in the Reshuitang area (also referred to as the Reshuitang hot springs) are also present as a spring group. Their temperatures are 27°C–89°C dominated by 80°C–87°C, mostly exceeding the local boiling point. These hot springs are primarily located on the high platform of a floodplain. The platform is made of calcareous tufa and spreads in the EW direction in the shape of an oval with a long axis of about 250 m and a short axis of about 150 m in length. The top of the platform is about 10 m above the water level. Hot and boiling springs are concentrated in an area of about 100 × 60 m² in the upper part of the high platform. They are composed of more than one hundred springs, each of which shows small water bubbles at dozens of positions. In addition, a few springs are distributed along the edge of the river valley. The surface geothermal manifestations include boiling spouters, many boiling springs, fumaroles, bubbling ground, and hot springs (Figure 2D). They are accompanied by a large amount of gas overflow and a small amount of H₂S odors, with milky white calcified sediments occurring in surrounding areas. In addition, geothermal wells (Figure 2E) and the Baqu River (Figure 2F) are also distributed near the Reshuitang hot springs.

**SAMPLING AND TESTS**

Samples were collected in September 2016. They include 10 samples collected from the Rekeng and Reshuitang hot springs in the area from Chaluo Village to the Cuopu Lake, one geothermal water sample and some core samples taken from the high platform of a geothermal well next to the Reshuitang hot springs (the geothermal well was subsumed into the Reshuitang hot spring system), and two surface water samples collected from the Cuopu Lake and the Baqu River. The pH, Eh, total dissolved solids (TDS), and temperature of water were measured on site using portable water quality analyzers. 500 ml polyethylene bottles were used to collect the water samples after being cleaned and rinsed. The hydrochemical analysis of the water samples was conducted at the Key Laboratory of Groundwater Science and Engineering of the Ministry of Natural Resources, the Institute of Hydrogeology and Environmental Geology, Chinese Academy of Geological Sciences. The REE content of the water and rock samples was determined at the Central South Mineral Resources Supervision and Testing Center, Ministry of Land and Resources. The water samples used for tests of cations, trace elements, and isotopes were acidized until their pH was <2 by adding some drops of concentrated hydrochloric acid, while water samples used for anion tests were not acidified. Before tests, hydrochloric acid was added into the water samples using the titration method to eliminate the impacts of microorganisms and other impurities. Cations, anions, and trace elements were determined primarily using ICP-MS as per GB/TB 538-2008 Methods for examination of drinking natural mineral water. The REE contents of the water samples were tested using a plasma mass spectrometer, with analytical accuracy of better than 5%.

**RESULTS AND DISCUSSION**

**Hydrochemical Characteristics**

The hydrochemical compositions of the water samples are shown in Table 1. The Schöeller diagram (Figure 3) can be used to reflect the physicochemical properties of geothermal water from hot springs and geothermal wells and surface water from lakes and rivers. The geothermal water is neutral to alkaline and surface water is alkaline, and they have similar chemical compositions. However, the geothermal water contains higher concentrations of major ions, especially Na ions, than the surface water (Figure 3). The TDS content of the surface water and the geothermal water is 75.61–98.98 mg/L and 305.5–1,179 mg/L, respectively. For most water samples, the anions are dominated by HCO₃⁻, and their contents are in the order of HCO₃⁻ > SO₄²⁻ > Cl⁻. Meanwhile, cations in the sample waters mainly included Na⁺, and their contents are in the order of Na⁺ > K⁺ > Ca²⁺ > Mg²⁺. The total dissolved cations (TZ⁺ = Na⁺ + K⁺ + 2Mg²⁺ + 2Ca²⁺) and total dissolved anions (TZ⁻ = Cl⁻ + 2SO₄²⁻ + HCO₃⁻ + 2CO₃²⁻) are well balanced: all samples have normalized inorganic charge balance, NICB = (TZ⁺ + −TZ⁻)/TZ⁺, and all samples have NICB ≤10% (Table 1) (Moon et al., 2007). The Piper diagram of the water samples (Piper, 1994; Figure 4) illustrates the distribution of these cations and anions and the hydrochemical classification of the water samples. According to this diagram, three hydrochemical types were identified. All the geothermal water is of Na-HCO₃ type, the river water is of the mixed Ca-Na-HCO₃ type, and lake water is of Ca-HCO₃ type. Moreover, the greater concentrations of major ions and the higher TDS content in the geothermal water indicate a longer retention time and stronger water-rock interactions of the geothermal water (Zhang Y. et al., 2018).

**Hydrochemical Processes**

Soluble ions in water originate from various natural processes, such as precipitation, water-rock interactions, and evaporation, and these different processes can be distinguished using Gibbs diagrams (Gibbs, 1970). In the Gibbs diagram of TDS versus Na⁺/ (Na⁺ + Ca²⁺) and Cl⁻/(Cl⁻ + HCO₃⁻) of the geothermal water (Figure 5), the Na⁺/(Na⁺ + Ca²⁺) ratio is 0.87–0.99, falling in the zones of water-rock interactions and evaporation. Meanwhile, the Cl⁻/(Cl⁻ + HCO₃⁻) ratio of the geothermal water is 0.05–0.17, falling into the water-rock interaction zone. These results indicate that cations in the geothermal water are mainly controlled by water-rock interactions and evaporation, while the anions in the geothermal water are dominated by water-rock interactions. The surface water samples fall in the water-rock interaction zone in the Gibbs diagram (Figure 5), indicating that the hydrochemical composition of the surface water is controlled by water-rock interactions.

The hydrochemical processes of water-rock interactions can be reflected from the correlation between the anion and cation...
According to the lithology of the strata in the study area, the rock formations contain minerals such as silicates, carbonates, sulphates and halides, and the main ions in the water samples originate from the dissolution of these minerals. Among these ions, Cl\(^-\) is rarely altered by water-rock interactions and mineral adsorption, even under high temperature and pressure conditions. Therefore, the correlation between Cl\(^-\) and other ions can account for the hydrochemical process in the geothermal water cycle (Li et al., 2020).

If Na\(^+\) is derived from halite dissolution, the molar ratio of Na\(^+\) to Cl\(^-\) should be 1:1. Figure 6A shows that the surface water samples are distributed near the origin of coordinates in which the Na\(^+\)/Cl\(^-\) ratios for CPW and RW were 0.56, 12.74 respectively. While the geothermal water shows a Na\(^+\)/Cl\(^-\) molar ratio of much higher than 1, implying that the excess Na\(^+\) in the geothermal water may originate from silicate weathering (Zhang Y. et al., 2018). Ca\(^{2+}\) and SO\(_4\)\(^{2-}\) in groundwater are affected by the dissolution and precipitation of gypsum. If merely gypsum dissolves and precipitates in groundwater, as shown in Eq. 1, the molar ratio of Ca\(^{2+}\) to SO\(_4\)\(^{2-}\) should be 1:1. In the plot of Ca\(^{2+}\)

| Sample | Location | T(°C) | Eh(mv) | pH | K\(^+\) | Na\(^+\) | Ca\(^{2+}\) | Mg\(^{2+}\) |
|--------|----------|-------|--------|----|--------|---------|------------|------------|
| BT13   | Rekeng   | 89    | -514.5 | 9.18 | 34.53  | 358.9   | 1.34       | 0.02       |
| BT14   | Rekeng   | 89    | -574   | 9.27 | 29.81  | 317     | 0.81       | 0.013      |
| BT15   | Rekeng   | 90    | -413.4 | 8.43 | 27.62  | 290.8   | 4.02       | 0.21       |
| BT18   | Rekeng   | 89    | -517   | 8.97 | 26.86  | 343.8   | 0.65       | 0.013      |
| BT20   | Rekeng   | 53    | 45.1   | 8.26 | 2.41   | 90.24   | 11.61      | 0.81       |
| BT22   | Reshuitang| 86    | -550   | 8.19 | 26     | 353.2   | 6.41       | 2.92       |
| BT24   | Reshuitang| 90    | -442.1 | 9.1  | 26.23  | 351.6   | 0.77       | 0.1        |
| BT25   | Reshuitang| 90    | 55.6   | 7.76 | 26.8   | 358.2   | 1.67       | 0.2        |
| BT26   | Reshuitang| 90    | 64.8   | 8.03 | 25.24  | 345.2   | 0.8        | 0.1        |
| BT27   | Reshuitang| 70    | 45.3   | 7.64 | 30.51  | 380.4   | 7.77       | 2.51       |
| CPH    | Cuopu Lake| 15    | 75     | 8.96 | 0.75   | 0.65    | 19.55      | 1.81       |
| RW     | Baqu River| 13    | 90     | 8.2  | 2.3    | 14.71   | 10.95      | 2.51       |

**TABLE 1** | Hydrochemical composition of water samples in the study area (mg/L).

| Sample | Cl\(^-\) | SO\(_4\)\(^{2-}\) | HCO\(_3\)\(^-\) | CO\(_3\)\(^{2-}\) | TDS | NICB\(^a\) (%) | Fe | Mn |
|--------|---------|-----------------|----------------|----------------|-----|----------------|----|----|
| BT13   | 19.19   | 32.84           | 379.1          | 222.5          | 1,079 | 10            | 0.242 | 0.032 |
| BT14   | 39.1    | 30.17           | 365            | 194.2          | 991.2   | 2.79          | 0.012 | 0.032 |
| BT15   | 42.65   | 42.06           | 617.5          | 42.09          | 909    | -0.21         | 0.021 | 0.032 |
| BT18   | 51.9    | 25.45           | 428            | 180.4          | 1,041   | 4.48          | 0.012 | 0.001 |
| BT20   | 8.89    | 20.74           | 217            | -              | 305.5   | 8.50          | 0.036 | 0.006 |
| BT23   | 33.77   | 19.47           | 904.9          | 18.04          | 1,071   | -1.21         | 0.17  | 0.011 |
| BT24   | 23.1    | 20.3            | 679.3          | 114.3          | 1,039   | -0.03         | 0.425 | 0.009 |
| BT25   | 33.77   | 42.62           | 885.9          | -              | 1,071   | 0.04          | 0.381 | 0.004 |
| BT26   | 33.06   | 42.09           | 862.1          | -              | 1,033   | -1.46         | 0.512 | 0.009 |
| BT27   | 32.7    | 25.9            | 1,015          | -              | 1,179   | -0.98         | 0.016 | 0.01  |
| CPH    | 1.78    | 22.89           | 42.8           | -              | 75.61   | -4.80         | 0.01  | 0.004 |
| RW     | 1.78    | 17.95           | 61.75          | -              | 98.98   | 1.08          | 0.162 | 0.003 |

\(^{a}\)NICB, normalized inorganic charge balance = (TZ\(^+\) - TZ\(^-\))/TZ\(^+\) × 100%.
versus $\text{SO}_4^{2-}$ (Figure 6B), only a few geothermal water and river water samples are distributed along the line with a slope of 1:1, and most of the geothermal water samples fall below the line with a slope of 1:1. These results indicate that the excess $\text{Ca}^{2+}$ may originate from the dissolution of carbonate and silicate minerals. In addition, the relatively high $\text{SO}_4^{2-}$ of some samples from the geothermal water and the Cuopu Lake may be influenced by human activities (Zhang Y. et al., 2018).

If the molar ratio of $\text{Ca}^{2+}$ to $\text{HCO}_3^-$ is 1:2, $\text{Ca}^{2+}$ in groundwater originates from the dissolution of carbonate minerals (Eq. 2; Guo et al., 2020). In the plot of $\text{Ca}^{2+}$ versus $\text{HCO}_3^-$ (Figure 6C), the samples of river water and Cuopu Lake are distributed along the line with a slope of 1:2, and the molar ratio of $\text{Ca}^{2+}$ to $\text{HCO}_3^-$ of samples from the geothermal water are much less than 1:2. These results indicate that the $\text{Ca}^{2+}$ in the geothermal water are derived from the dissolution of silicate minerals, while the $\text{Ca}^{2+}$ in the
river water and Cuopu Lake are derived from the dissolution of carbonate minerals.

The origin of Ca$^{2+}$, Mg$^{2+}$, HCO$_3^-$, and SO$_4^{2-}$ in groundwater can be reflected from the molar ratio of (Ca$^{2+}$ + Mg$^{2+}$) to (HCO$_3^-$ + SO$_4^{2-}$) in water. Specifically, a molar ratio of 1:1 indicates that these ions are obtained from the dissolution of carbonate rocks and sulfate minerals, a molar ratio of greater than 1:1 indicates that ion exchange process dominates (Eq. 3), and a molar ratio of less than 1:1 indicates the presence of reverse ion exchange (Eq. 4). As shown in plot of (Ca$^{2+}$ + Mg$^{2+}$) versus (HCO$_3^-$ + SO$_4^{2-}$) of the water samples (Figure 6D), the samples of river water and Cuopu Lake are distributed along the line with a slope of 1:1, while the water samples of the geothermal water are distributed below the line. Meanwhile, the concentration of Ca$^{2+}$ is far higher than that of Mg$^{2+}$ in the geothermal water. Therefore, it can be inferred that the relatively depleted Ca$^{2+}$ in the geothermal water may be affected by ion exchange (Eq. 3), which is caused by silicate weathering.

In the diagram of (Na$^+$ + K$^+$)-Cl$^-$ versus (Ca$^{2+}$ + Mg$^{2+}$)-(SO$_4^{2-}$ + HCO$_3^-$ + CO$_3^{2-}$) (Figure 6E), all samples are essentially distributed along a 1:1 line. These results indicate that the hydrochemical processes of the geothermal water and the surface water are mainly controlled by cation exchanges and adsorption in the dissolution of silicate minerals (Li et al., 2020).
Cation exchanges and adsorption in the geothermal water result in a decrease in Ca$^{2+}$ and Mg$^{2+}$ concentrations and an increase in Na$^+$ concentration (Figure 3, Wang et al., 2021). According to the molar ratio diagrams of the major ions in water, the hydrochemical processes in the study area are mainly controlled by the dissolution of silicate minerals.

$$\text{CaSO}_4 \cdot 2\text{H}_2\text{O} = \text{Ca}^{2+} + \text{SO}_4^{2-} + 2\text{H}_2\text{O} \quad (1)$$

$$\text{CaCO}_3 + \text{H}_2\text{CO}_3 \rightarrow \text{Ca}^{2+} + 2\text{HCO}_3^- \quad (2)$$

Ion exchange: $2\text{Na}^+ - \text{Clay} + \text{Ca}^{2+} \rightarrow 2\text{Na}^+ + \text{Ca}^{2+} - \text{Clay}_2$ \quad (3)

Reverse ion exchange: $2\text{Na}^+ + \text{Ca}^{2+} - \text{Clay}_2 2\text{Na}^+ - \text{Clay}$ + $\text{Ca}^{2+} \rightarrow 2\text{Na}^+ - \text{Clay} + \text{Ca}^{2+} \quad (4)$

**Characteristics and Controlling Factors of Rare Earth Element Contents in Geothermal Water**

The composition of REEs in geothermal water in the study area is shown in Table 2. The contents of REEs (except for Yttrium) in the Chaluо hot springs are lower than those of the hot springs around the Kangding area in western Sichuan Province. They are $0.306 \pm 0.103$ μg/L and $0.669 \pm 0.367$ μg/L, respectively (Zhang, 2018b). In addition, the water in the Cuopu Lake has low REE contents, while the river water has high REE contents. The difference in the REE contents between the surface water and the geothermal water may be related to the material exchanges between water and rocks (Fan et al., 2021).

Studies have shown that REEs in water bodies are derived from the weathering or dissolution of minerals, the reductive dissolution of Fe and Mn oxides in sediments, the leaching and dissolution of secondary minerals, and human activities (Markert and Zhang, 1991; Dia et al., 2000; Tweed et al., 2006; Kynicky et al., 2012). The dominant factors of these REE sources include pH, oxidation-reduction potential (OPR), and mineral adsorption/dissolution (Leybourne et al., 2000; Noack et al., 2014; Gruau et al., 2004; Koenpenkastrop and De Carlo, 1992; Coppin et al., 2002). pH can control the REE contents by directly influencing the chemical weathering or adsorption/precipitation of REE minerals and by indirectly affecting the complexation (Namely, inorganic ions in the water column such as CO$_3^{2-}$, PO$_4^{3-}$, F$^-$, SO$_4^{2-}$, Cl$^-$, NO$_3^-$ as ligands combine with rare Earth elements to form complexes) or adsorption of REEs (Miller, 1992; Noack et al., 2014). Generally, there is a close negative correlation between the pH and the REE contents of water bodies (Noack et al., 2014). Figure 7A shows almost no correlation between the pH and REE ($R^2 = 0.02$) of geothermal water in the study area. However, the REE tends to increase with a decrease in pH when pH < 9, and there is a significant positive correlation between the REE and pH ($R^2 = 0.84$) when pH > 9. These results are consistent with the study of Noack et al. (2014). It is speculated that the adsorption-absorption regulation of REEs decreases as alkalinity increases in the water bodies. Meanwhile, the increase in the dissolution of CO$_2$ increases REE through complexation reactions (Zhu, 2006; Liu, 2018). OPR can directly affect redox-sensitive REEs such as Ce and Eu and can indirectly affect REEs by influencing the precipitation or dissolution of Fe and Mn oxides/hydroxides (Guo et al., 2010). Fe and Mn oxides/hydroxides adsorb REEs in an oxidizing environment but...
dissolve and release REEs in a reducing environment. As shown in the diagram of Eh versus $\Sigma$REE of the study area (Figure 7B), there is essentially no correlation between the Eh and $\Sigma$REE, reflecting the fact that REEs are barely influenced by Eh. In alkaline water bodies, HCO$_3^-$ occurs in the form of LnCO$_3^+$ and Ln(CO$_3$)$_2^-$ after experiencing complexation reactions with REEs (Wood, 1990). Figure 7C shows that there is no correlation between HCO$_3^-$ and $\Sigma$REE. However, in geothermal water with a high HCO$_3^-$ content, the $\Sigma$REE tends to increase with an increase in HCO$_3^-$. This finding indicates that high HCO$_3^-$ content is favorable for REE enrichment (Xie et al., 2012). In geothermal water with a low HCO$_3^-$ content, however, REEs may be liable to be affected by other factors such as pH. The diagrams of Fe versus $\Sigma$REE and Mn versus $\Sigma$REE (Figures 7D,E) show a certain positive correlation between Fe and $\Sigma$REE and no significant correlation between Mn and $\Sigma$REE. These results indicate that the reductive dissolution of Fe oxides/
hydroxides in geothermal water accounts for the increase in the REE content. Overall, the REE contents depend on the reductive dissolution of Fe oxides/hydroxides, followed by pH.

**Rare Earth Element Fractionation in Geothermal Water and Controlling Factors**

The North American Shale Composite (NASC) (Haskin et al., 1968) normalized REE patterns of the geothermal water in the study area (Figure 8) take the shape of gentle sawteeth (Haley et al., 2004). The LREE(La-Eu)/HREE(Gd-Lu) ratio of geothermal water varies from 5.19 to 14.38. The (La/Yb)\textsubscript{N} ratio is generally used to replace the LREE(La-Eu)/HREE(Gd-Lu) ratio to characterize the relative enrichment of LREEs in geothermal water (Zhao Y. Y. et al., 2019). The (La/Yb)\textsubscript{N} ratio of the geothermal water in the study area is 0.41–2.95, with an average of 1.31, reflecting the enrichment of LREEs.

Figure 9A shows no significant correlation between Eh and (La/Yb)\textsubscript{N}, indicating that Eh does not directly affect the REE contents. Studies show that Fe and Mg preferentially adsorb LREEs in the process of oxidation-induced precipitation, leading to the relative enrichment of HREEs. Meanwhile, Fe and Mg oxides will release LREEs after reductive dissolution, leading to a relative enrichment of LREEs in water bodies (Tang and Johanesson, 2010). Figure 9B shows a significant positive correlation between Fe and (La/Yb)\textsubscript{N}. That is, as the Fe content increases, (La/Yb)\textsubscript{N} increases significantly and gradually and the REE fractionation enhances. In contrast, Figure 9C shows that this is almost no correlation between Mn and (La/Yb)\textsubscript{N}. These occur possibly for the following reasons. Fe and Mg oxides have different absorption capacities of LREEs and HREEs. The Fe concentration is higher than the Mn concentration in the geothermal water, and Fe oxides release more LREEs after reductive dissolution. As a result, the geothermal water is richer in LREEs than in HREEs.

The redox-sensitive Ce and Eu elements are also commonly used to reflect the fractionation of REEs. Ce exists in both +3 and +4 valence states. In an oxidizing environment, Ce\textsuperscript{3+} in water bodies converts into Ce\textsuperscript{4+} after losing electrons, which will be preferentially adsorbed by Fe and Mn oxides and is then separated from other REEs in the +3 valence state, leading to Ce depletion in water bodies. In a reducing environment, adsorbed Ce\textsuperscript{4+} is reduced to Ce\textsuperscript{3+} and enters water bodies due to the dissolution of Fe and Mn oxides, resulting in Ce enrichment (Byrne and Sholkovitz, 1996).
Eu normally occurs in the Eu\(^{3+}\) valence state. In a reducing environment, Eu\(^{3+}\) converts into Eu\(^{2+}\) after extracting electrons, which will separate from the other REEs in the +3 valence state due to chemical differences, thus resulting in Eu depletion. Among various equations used to calculate Ce/Ce* and Eu/Eu* ratios (Zhao Y. Y. et al., 2019), the following equations (Guo et al., 2010) are used in this study:

\[
\frac{\text{Ce}}{\text{Ce}^*} = \frac{\text{CeN}}{(\text{LaN} \times \text{PrN})} \cdot 0.5
\]

\[
\frac{\text{Eu}}{\text{Eu}^*} = \frac{\text{EuN}}{(\text{SmN} \times \text{GdN})} \cdot 0.5
\]

The Ce/Ce* ratio of the geothermal water in the study area is 1.31–5.96, with an average of 2.43, reflecting positive Ce anomalies. The Eu/Eu* ratio of the geothermal water is 2.52–32.93, with an average of 14.56, indicating positive Eu anomalies. Factors affecting Ce anomalies in water bodies usually include OPR and mineral adsorption/dissolution or the weathering of the surrounding rocks (Elderfield et al., 1990). There is a weak negative correlation between Eh and the Ce/Ce* ratio in the geothermal water in the study area (Figure 9D). That is, a lower Eh suggests a higher Ce/Ce* ratio, indicating that the positive Ce anomalies in water bodies are caused by the redox environment. Specifically, Ce\(^{4+}\) converts into Ce\(^{3+}\) with the dissolution of Fe and Mn oxides, resulting in positive Ce anomalies (Xie et al., 2012). In contrast, Figure 9E shows a certain negative correlation between Fe and the Ce/Ce* ratio, and Figure 9F shows a certain positive correlation between Mn and the Ce/Ce* ratio. These results indicate that the dissolution of Mn oxides in water bodies leads to positive Ce anomalies.

To further investigate the REE fractionation in water-rock interactions, this study obtained NASC-normalized REEs patterns of the Tumugou and Qugasi formations and Yanshanian granites (Figure 1C) in the study area. According to the NASC-normalized REE patterns, the rocks of the Qugasi Formation show positive Eu anomalies (Wang, 2017; Figure 10). In contrast, rocks of the Tumugou Formation and especially the Yanshanian granites show negative Eu anomalies (Zhang F. Y. et al., 2018). All types of rocks show slightly negative Ce anomalies, from which it can be inferred that the weathering of surrounding rocks roughly does not cause positive Ce anomalies. The surface water near the geothermal water shows positive Ce anomalies (Figure 8), with a Ce/Ce* ratio of 1.49–4.77, with an average of 3.13. This value is slightly higher than the Ce/Ce* ratio of the geothermal water, indicating that the geothermal water may be influenced by the surface water (Barrat et al., 2000). Therefore, the positive Ce anomalies in the geothermal water result from both the dissolution of Eh and Mn oxides and surface water. Table 3.

The negative correlation between Eh and the Eu/Eu* ratio in the geothermal water (Figure 9G) indicates that the positive Eu anomalies in the geothermal water are caused by the reducing environment. Meanwhile, the correlations between Fe and the Eu/Eu* ratio (Figure 9H) and between Mn and the Eu/Eu* ratio (Figure 9I) indicate that the dissolution of Mn oxides promotes the positive Eu anomalies. As mentioned above, the NASC-normalized

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**FIGURE 10** | Standardized allotment map of North American shales in the study area rocks (Qugasi Formation data (QGSF1, 4, 5) according to Wang H Y, 2017; HZS data according to Zhang F. Y. et al, 2018).

**TABLE 3** | Rare Earth element composition of geothermal well cores in the study area (ug/g).

| Sample   | La  | Ce  | Pr  | Nd  | Sm  | Eu  | Gd  |
|----------|-----|-----|-----|-----|-----|-----|-----|
| BT23-Y1  | 51  | 74.5| 7.96| 32.4| 5.58| 1.08| 4.82|
| BT23-Y2  | 35.7| 58.6| 6.61| 28.4| 5.44| 0.96| 4.67|
| BT23-Y3  | 23.3| 37.3| 4.33| 18.5| 3.66| 0.68| 3.26|
| BT23-Y4  | 29  | 46.9| 5.54| 23.8| 4.62| 0.91| 3.96|

| Sample   | Tb  | Dy  | Ho  | Er  | Tm  | Yb  | Lu  |
|----------|-----|-----|-----|-----|-----|-----|-----|
| BT23-Y1  | 0.74| 4.39| 0.88| 2.36| 0.36| 2.37| 0.31|
| BT23-Y2  | 0.73| 4.3 | 0.85| 2.32| 0.36| 2.35| 0.31|
| BT23-Y3  | 0.53| 3.19| 0.65| 1.77| 0.27| 1.83| 0.25|
| BT23-Y4  | 0.62| 3.62| 0.72| 1.94| 0.3 | 1.94| 0.26|
REE patterns (Figure 10) show that only rocks in the Qugasi Formation show positive Eu anomalies, while the rocks in the Tumugou Formation and Yanshanian granites show negative Eu anomalies. It can be inferred from these results that water-rock interactions between the Qugasi Formation and deep geothermal water also cause positive Eu anomalies. The Eu/Eu* ratio of the surface water is 7.94 and shows significant positive Eu anomalies which indicating positive Eu anomalies in surface water was inherited from the rock of Qugasi Formation (Dong et al., 2017). It is lower than that of the geothermal water, suggesting that the geothermal water is affected by the surface water. In addition, the dissolved feldspar minerals in water bodies also lead to positive Eu anomalies and calcite precipitation (Lee et al., 2003; Liu et al., 2016). The surrounding rocks in the study area are rich in feldspar minerals and the geothermal water in the Batang area is supersaturated with calcite (Zhang, 2020). Therefore, the dissolution of feldspar in the surrounding rocks in the study area promotes the Eu enrichment. Overall, the Eu positive anomalies in the geothermal water result from the combined effects of the reducing environment, surrounding rocks, water bodies, and feldspar dissolution.

Distribution of Rare Earth Element Complexes in Geothermal Water

In this study, the composition and forms of REEs in water sample BT15 from the Rekeng hot springs and the water sample BT25 from the Reshuitang hot springs were determined through
calculation and simulation analysis using the Visual MINTEQ 3.0 software. Specifically, measured temperatures, pH, major anions, and REE concentrations were input to MINTEQ to determine the inorganic complex species of REEs in the water samples. The simulation results show that the complex species in the geothermal water mainly include Ln(CO$_3$)$_2^-$, LnCO$_3$$^{+}$, and LnOH$^{2+}$ (Ln signifies the REE). This finding is roughly consistent with the complex species in partially alkaline water bodies (Liu et al., 2021). As shown in Figure 11, the inorganic complexes in the Rekeng geothermal water are dominated by Ln(CO$_3$)$_2^-$, which accounts for 90.47–99.11% of the total mass concentration of inorganic complexes. They also include LnCO$_3$$^{+}$ and LnOH$^{2+}$, which account for 0.82–9.06% and 0.06–1.14%, respectively, and other complexes account for less than 0.01%. The complexes in the Reshuitang geothermal water mainly include Ln(CO$_3$)$_2^-$ and LnCO$_3$$^{+}$, which account for 62.89–95.22% and 4.47–33.18%, respectively. They also include LnSO$_4$$^{2-}$, which accounts for 0.013–1.31%, and other complexes account for less than 0.1%.

As shown in Figure 11, with an increase in the atomic number of the REEs in the geothermal water, the percentage of Ln(CO$_3$)$_2^-$ gradually increases but the percentages of other complexes such as LnCO$_3$$^{+}$ gradually decrease (Liu et al., 2021). This occurs because the stability constants (lgK) of different REE complexes increase as the atomic order of REEs increases. The lgK of (CO$_3$)$_2^-$ is higher than that of other complexes, and therefore the percentage of Ln(CO$_3$)$_2^-$ increases with an increase in the atomic number of REEs (Figure 12).

CONCLUSION

This study determined the hydrochemical characteristics and REE contents of water samples collected from the Chaluo hot springs and rock samples from the surrounding areas through analysis and tests, obtaining the following conclusions:

1) In terms of hydrochemical types of the Chaluo hot springs, the geothermal water is of Na-HCO$_3$ type, the river water is of mixed Ca-Na-HCO$_3$ type, and the lake water is of Ca-HCO$_3$ type. The cations in the geothermal water are mainly controlled by water-rock interactions and evaporation, the anions are primarily determined by water-rock interactions, and the hydrochemical composition of the surface water is controlled by water-rock interactions. Overall, the hydrochemical processes in the water bodies of the study area are mainly controlled by the dissolution of silicate minerals.

2) The total REE content in the geothermal water is 0.306 ± 0.103 μg/L. Studies reveal that the REE contents are primarily affected by the reductive dissolution of Fe oxides/hydroxides, followed by pH.

3) The LREE/HREE and (La/Yb)$_{NG}$ ratios of the geothermal water are 5.19–14.38 and 0.41 to 2.95, respectively, indicating the relative enrichment of LREEs. They are mainly affected by Fe oxides. In addition, the geothermal water shows positive Eu and Ce anomalies, which are caused by both the dissolution of Eh and Mn oxides and surface water. Furthermore, the water-rock interactions between the Qugasi Formation and deep geothermal water also contribute to the positive Eu anomalies.

4) According to the results of the calculation and simulation analysis using the Visual MINTEQ 3.0 software, the complex species of REEs in the geothermal water mainly include Ln(CO$_3$)$_2^-$, LnCO$_3$$^{+}$, and LnOH$^{2+}$. This finding is roughly consistent with the complex species of REEs in alkaline water and is caused by the stability constants of complexation reactions.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

SW: Drafting the manuscript, analysis and interpretation of data. FL: Acquisition of data. WZ: Analysis and interpretation of data. HZ: Acquisition of data. RY: Verify data. YL: Participate in mapping. XY: Check the manuscript.

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