

Article



Mineralogy and Processing of Hydrothermal Vein Quartz from Hengche, Hubei Province (China)

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Abstract: Quartz occurs in many geological materials, and is used in numerous industrial fields as a raw material. Mineralogy and the processing of hydrothermal quartz were studied by optical microscope, electron probe microanalysis, scanning electron microscope, inductively coupled plasma-optical emission spectrometry, and inductively coupled plasma mass spectrometer. A combination of the geological occurrence of the quartz deposit, mineralogical studies, and the processing technologies of the hydrothermal quartz was accomplished. The results show that impurities within the quartz mainly include muscovite, hematite, apatite, and secondary fluid inclusions. The main chemical impurities are Al (353 μ g·g⁻¹), K (118 μ g·g⁻¹), Fe (61.2 μ g·g⁻¹), P (15.5 μ g·g⁻¹), Na (13.4 μ g·g⁻¹), Mg (11.8 μ g·g⁻¹), Ti (8.31 μ g·g⁻¹), and B (10.8 μ g·g⁻¹). Based on these results, a combined process consisting of calcination and fluoride-free pressure acid leaching was established to effectively decompose and dissolve the quartz, and remove gangue minerals and fluid inclusions. The calcination process not only removed volatile components; it also destroyed the crystal structure of gangue minerals and enhanced their release probabilities. The calcination process has a positive influence on the removal of impurity elements by the fluoride-free pressure acid leaching process. A total of 85.2 wt % and 84.0 wt % of impurity elements was removed using the leaching systems of HCl-NH₄Cl and H₂SO₄-NH₄Cl, respectively.

Keywords: hydrothermal quartz; geological setting; mineralogy; processing technology

1. Introduction

SiO₂ minerals and rocks play an important role in geological processes and industrial applications [1]. Quartz is one of the most important silica minerals, and it is abundant in the Earth's crust in igneous, metamorphic, and sedimentary rocks [1]. SiO₂ minerals and rocks have been formed by primary and secondary magmatic, hydrothermal, or sedimentary processes or during diagenesis and metamorphosis [2].

Hydrothermal quartz is an important substitute of crystal quartz that has been studied in detail because it is rich in SiO₂ [3,4]. The hydrothermal quartz commonly contains a significant amount of associated minerals and fluid inclusions, which leads to difficulties in quartz processing [5]. Some primary and fine-grained mineral inclusions including mica, feldspar, and hematite are difficult to be removed because they are closely included within quartz grains. Unlike metal ores, the quartz can't be freely ground to cause the decomposition of quartz and fine-grained gangue minerals, because there is a special demand for particle size in quartz sand production [6]. Further, some micron-size fluid inclusions are widely distributed in hydrothermal quartz. Although decrepitation methods at low and medium temperatures (below 700 °C) can enhance the removal of volatile materials in quartz fluid inclusions, the micron-size fluid inclusions rarely are removed, especially in high-temperature quartz [7,8].

Quartz processing technologies have been studied in detail, but systemic research on geological occurrence and mineralogy, combined with optimized processing technologies, is warranted [9,10]. The research on the geological formation and mineralogy of hydrothermal quartz provides an important basis for quartz processing, and helps to characterize the quartz ore and its value.

Processing technologies of high-grade quartz mainly include pre-processing, physical processing (magnetic separation and flotation), chemical treatment (acid leaching and hot chlorination), and thermal treatment (calcination) [11–14]. Furthermore, physical processing and chemical leaching are common processing technologies used in industry. Magnetic separation is mainly used for removing iron-bearing magnetic minerals, while flotation is mainly used for removing aluminum-bearing silicate minerals from quartz ore [15]. However, a common weakness of both processes is that they can only remove specific kinds of impurity minerals. This weakness unavoidably leads to a long process flow during the effective purification of quartz. Although the acid leaching process can effectively remove different impurities at the same time, its leaching solution commonly contains HF, NaF, or CaF. Fluoric reagents, especially hydrofluoric acid, can cause environment pollution [16,17]. Leaching techniques are more effective than flotation for removing intergrown impurities of quartz and gangue minerals [18,19], but the selectivity of acid leaching with hydrofluoric acid is typically low, and this process could sharply reduce particle size of quartz sand as it removed impurities as a result [20,21].

The calcination process, as a thermal treatment technology, is commonly used to reduce reaction time and agent consumption by increasing the exposure probability of gangue minerals and destroying their crystal structures [10,11]. In addition, pressure acid leaching with mixed agents consisting of acids and inorganic salts is deemed to be an effective method to dissolve and remove muscovite without using any fluorides [22]. The metallic ions in the inorganic salts are unacceptable in quartz purification because they would unavoidably drag in metallic impurities during processing, but the use of NH₄Cl avoids the problem. Therefore, leaching systems including HCl-NH₄Cl and H₂SO₄-NH₄Cl could be introduced into quartz purification alongside the calcination process to effectively remove impurities from quartz.

This study presents an analysis of the regional geological setting and mineralogy of the hydrothermal quartz from Hengche Town in the Qichun County of the Hubei Province of China, and further develops a suitable process flow for the purification of hydrothermal quartz. In particular, the research is focused on analyzing occurrences and distributions of impurity elements in the quartz. Based on those results of this mineralogy, a combination process consisting of calcination and fluoride-free acid leaching was developed to remove impurities in the hydrothermal quartz. More importantly, the effects of the calcination and pressure acid leaching on quartz sand were evaluated.

2. Materials and Methods

2.1. Materials and Geological Situation

Quartz ore samples studied are from a hydrothermal vein deposit in Hengche Town in the Qichun County of the Hubei Province in China. The hydrothermal vein deposits are located in the south of a fold belt along the Qinling and Dabie mountains, and close to the Yangtze block. Figure 1 shows a sketch of the geology of the southeast of Hubei Province, China. The Qichun County belongs to the fold belt along the Qinling and Dabie mountains (I), which are bordered by the Yangtze block along the Zengjiaba–Qingfeng–Xiangguang (ZQX) fault zone [23,24]. The reentry belt of high pressure and ultrahigh pressure metamorphism from Tongbai to Dabie (I23) [23,24] was formed by the subduction and collision of the area from late Paleozoic to Triassic, and ended in late Triassic and early Jurassic age. In general, the quartz deposits in the area were formed in the combined belt of Shangdan, influenced by the orogenesis of the Qinling Mountains, and activated by the Yangtze block [23].

Subduction and collision from Paleozoic to Jurassic age caused ore-forming conditions, and the high pressure and ultrahigh pressure rarely contribute to the deposit. As a result, the agglutinated

fine-grained quartz likely contains secondary mineral inclusions in assemblages. In late Triassic to Mesozoic age, the northward subduction and reentry of the Yangtze block or Shangdan Ocean resulted in an orogenic belt south of the Qinling Mountains. This indicates that the geological environment provides a good framework for the deposit formation. Since the quartz deposit is close to the block in the south of the Qinling Mountains, some wall rocks rich in Si, Fe, P, S, and Ca unavoidably have an influence on assemblages of quartz and enlarged sides.



Figure 1. Locations of the Qinling and Dabie Mountains, and a sketch map of geological formations in the southeast of the Hubei Province in China (Revised according to Mao et al., 2014) [23]: QL—Qinling Mountain, DB—Dabie Mountain, BJ—Beijing, HQD—the hydrothermal quartz deposit, MC—Macheng, YC—Yingcheng, XZ—Xinzhou, LT—Luotian, HC—Hanchuan, TF—Tuanfeng, YS—Yingshan, WH—Wuhan, HG—Huanggang, EZ—Ezhou, XS—Xishui, HS—Huangshi, QC—Qichun, HM—Huangmei, DY—Daye, WX—Wuxue; Fd—Fault depressions, I—Qinling and Dabie fold belt, II—Yangtze block, I2-Combining belt of Shangdan, I3—Block in the southern of Qinling Mountains, I4—Arc basin system from Maota to Suizhou, I5—Combining belt from Liangzhu to Suinan, I23—Reentry belt of high pressure and ultrahigh pressure metamorphism from Tongbai to Dabie, II11—Passive margin in the north rim of Yangzi block, II21—Forelandbasin in the south rim of Yangzi block, II22—Passive margin in the south rim of Yangzi block.

When compared with the few ultrahigh purity quartz deposits ($c(AI) < 20 \ \mu g \cdot g^{-1}$) in Qichun County [24], the Hengche quartz is from a typical, low-grade and small-scale deposit. In the Luliang age, a hydrothermal solution rich in SiO₂ was split from granite magma. Along the bedding, the vein quartz was formed when the hydrothermal solution invaded into biotite-plagioclase gneiss, granitic gneiss, and amphibolite in the Hongan Group of the Dabie Mountains. The deposit length ranges from 80 m to 115 m, the deposit width ranges from 60 m to 120 m, and the deposit thickness ranges from 5 m to 18 m. The ore mainly contains quartz, and the size of the quartz grain is about 0.8 mm, while a part of the quartz grain is larger than 1 mm.

2.2. Mineralogical Analysis

Examples of the typical bulk hydrothermal quartz were sliced up and polished into sections for microscopy observation. Mineral phases and their distribution in quartz ore were determined by polarizing microscope (DLMP, Leica Microsystems, Wetzlar, Germany). Element components of gangue

minerals were analyzed by electron probe microanalysis (EPMA, JXA-8230, JEOL Ltd., Tokyo, Japan) and X-ray energy spectrometer (EDS).

The electron optical system of EPMA consists of a LaB6 electron gun with a centered cartridge filament; acceleration voltage: $0 \sim 30$ kV; beam ranging: $10^{-5} \sim 10^{-12}$ A; and image magnification: $\times 40 \sim \times 300,000$. Analysis precision of EDS: analysis element: 5B-92U, $2\% \sim 3\%$ for common element, detection limit: 0.1 wt %~0.5 wt %. Electron images were obtained at an acceleration voltage of 20.0 kV and a working distance of 11.2 mm. The GBT15617-2002 standard is used for EPMA [25].

2.3. Chemical Analysis

The chemical composition of the hydrothermal quartz was analyzed by inductively coupled plasma (ICP) techniques according to national standards SJ/T 11554-2015 [26] and GB/T 32650-2016 [27] in China. The analytical method and procedure are as follows.

2.0000 g per sample were dissolved by 12 cm³ hydrofluoric acid (40 wt %, guarantee reagent) in a pressure-tight reaction kettle with a lining of polytetrafluoretyhylene (PTFE, 50 cm³), while digestion temperature and time were 200 °C and 3 h, respectively. The produced fluid was heated and evaporated in a PTFE beaker at 150 °C. When the fluid shrank into droplets, 3 cm³ ultrapure water was added for further evaporation, while the operation was repeated three times, and the remained fluid was commonly diluted to 20 cm³, 500 cm³, or 1000 cm³, according to the different contents of impurity elements.

The element contents of diluted fluids were analyzed by inductively coupled plasma-optical emission spectroscopy (ICP-OES, Prodigy 7, LEEMAN LABS INC., Hudson, NH, USA) and inductively coupled plasma mass spectrometry (ICP-MS, iCAP-Qc, THERMO FISHER SCIENTIFIC, Waltham, MA, USA).

The analytical conditions include:

Wavelength coverage of ICP-OES: 165~1100 nm, optical resolution: 0.007 nm at 200 nm, relative standard deviation: <2%.

Mass spectrum of ICP-MS: $4\sim290$ amu, sensitivity: >150 Mcps·ppm⁻¹ for common elements, detection limit: <0.1 ppt for common element, relative standard deviation: <2% in short time, <3% in long time.

The computational formula to obtain the element contents is as follows:

$$\varphi = \frac{V(\rho - \rho_0)}{m} \tag{1}$$

where φ is the mass of element content in 1 g of quartz sand, $\mu g \cdot g^{-1}$; *V* is the volume of the diluted fluid, cm³; ρ is the mass concentration of the analyzed element by ICP, ρ_0 is the mass concentration of the blank sample, mg·dm⁻³ or $\mu g \cdot dm^{-3}$; m is the mass of the analyzed sample, 2.0000 g.

2.4. Processing and Characterization of Quartz Sand

Mining, washing, crushing, and grinding of quartz ore were accomplished by Kaidi Building Materials Co., Ltd. (Huangshi, China). The mined quartz ore was washed and air-dried. Then, the bulk quartz was crushed by a jaw crusher (PE- Φ 100 × 125), and ground by a Raymond mill (3R2115). Produced quartz sand ranging from 106 µm (140 mesh) to 212 µm (70 mesh) was selected by standard sieve, and used for experiments. The quartz sand was calcinated at 900 °C for 5 h in a muffle furnace (KSY-12-16A), and then cooled by ultrapure water (18.2 MΩ·cm). The calcinated quartz sand was leached by mixed leaching agents in a pressure-tight reaction kettle. Leached quartz sand was washed by ultrapure water, and the leach liquor and washing liquor were mixed to wash the calcinated quartz sand in turn. Quartz and gangue minerals during processing were observed by scanning electron microscope (SEM, JSM-IT300) and EPMA after carbon coating.

Secondary electron images were obtained at 20.0 kV of acceleration voltage, 16.2~16.6 mm of working distance, and 30 nm of spot size.

3. Results and Discussion

3.1. Mineralogy of Hydrothermal Quartz

Bulk quartz is milky-white as a whole, and its size ranges from 20 cm to 40 cm. Scattered red materials are distributed in quartz fractures.

3.1.1. Impurity Elements in Quartz Ore

Elemental concentrations of the quartz sand are shown in Table 1. The main impurity elements determined by ICP are Al (353 μ g·g⁻¹), K (118 μ g·g⁻¹), Fe (61.2 μ g·g⁻¹), P (15.5 μ g·g⁻¹), Na (13.4 μ g·g⁻¹), Mg (11.8 μ g·g⁻¹), B (10.8 μ g·g⁻¹), Ti (8.31 μ g·g⁻¹) and Ca (8.04 μ g·g⁻¹) by ICP analysis, and proportions of these elements in total impurities are 57.0 wt %, 19.1 wt %, 9.89 wt %, 2.50 wt %, 2.16 wt %, 1.91 wt %, 1.74 wt %, 1.34 wt % and 1.30 wt %, respectively.

Table 1. Contents of main impurity elements within the hydrothermal quartz analyzed by inductively coupled plasma (ICP) analysis.

Element	Al	Fe	Na	S	Р	Li	К	Ca	Ti
Content ¹ ($\mu g \cdot g^{-1}$)	353	61.2	13.4	5.64	15.5	2.20	118	8.04	8.31
Element	Mg	Ni	Zr	Zn	As	В	In total		
Content ($\mu g \cdot g^{-1}$)	11.8	1.01	6.46	0.567	3.16	10.8	619		
1 Relative standard deviations (RSD) are less than 4%.									

3.1.2. Optical Microscope Analysis

The hydrothermal quartz mainly contains solid inclusions of muscovite and hematite, as well as fluid inclusions (Figure 2). The muscovite platelets are widely distributed along the quartz grain boundaries (Figure 2a), but a small quantity of muscovite is included in the quartz grains (Figure 2b). Muscovite is commonly thought to be an important source of impurity elements, including Al, K, Mg, Fe and Ti [28].



Figure 2. Transmitted-light microscopy images of muscovite, hematite, and fluid inclusions within quartz thin sections: (**a**) muscovite along a grain boundary (XPL); (**b**) muscovite included in quartz crystal (PPL); (**c**) hematite in compression fractures of quartz (PPL); (**d**) two generations of fluid inclusions (PPL).

Hematite is the main iron carrier in these veins, and mainly occurs filling thin cracks (Figure 2c). In hydrothermal quartz, most of the fluid inclusion generations (Figure 2d) traverse the whole quartz grains, so these secondary fluid inclusions commonly containing gas and liquid phases are a possible source of fluid impurities [29].

3.1.3. Electron Probe Microanalysis

Figure 3 shows secondary electron images of muscovite, hematite, apatite, and quartz sections. Figure 3a is a micrograph of muscovite inclusions within quartz. The analysis (Table 2) shows that Al, K, Mg, and Fe are the major elements obtained in the muscovite. These impurities occurring in aluminum silicate minerals could reduce the quality of quartz production [30]. Figure 3b is an image of hematite (the mineral phase was determined by polarizing microscope within an electron probe microanalyzer) in quartz, and its composition (Table 2) shows that the secondary mineral contains elevated Fe (the Cr is from polishing powder). Generally, the iron in high-grade quartz is typically less than 1 ppm [31]. It is difficult to remove some filmy iron on the surface of quartz particles by conventional physical methods [32]. Figure 3c is an image of apatite, which is associated with hematite. Its chemical analysis (Table 2) shows that P, Ca, Fe, Mg, and Na occur in the mixed minerals. The apatite is included in quartz, and it is a diagnostic mineral of hydrothermal quartz [33].





Figure 3. Details of solid inclusions in quartz veins using EMP (electron microprobe) images in SE (secondary electron) mode: (**a**) muscovite inclusions in quartz; (**b**) hematite in quartz fracture; (**c**) apatite and hematite included in quartz; (**d**) a typical microarea used for area-scan analysis of the electron probe; AP—analyzed point by X-ray energy spectrometer (EDS).

Element	Na	Mg	Al	Si	Р	S	К	Ca	Mn	Cr	Fe	0
Figure 3a (wt %)		0.31	19.16	24.69			8.40				0.28	47.17
RSD ² (%)		2.43	1.10	0.96			1.22				1.96	
Figure 3b (wt %)			2.24	2.48	0.63					2.89	65.91	25.85
RSD (%)			2.23	1.32	2.64					1.08	0.68	
Figure <mark>3</mark> c (wt %)	3.31	5.06	1.57	6.63	10.38	2.01	0.87	14.50	0.91		14.52	40.24
RSD (%)	2.88	2.09	2.11	1.16	1.67	1.34	2.89	1.44	1.57		0.83	

Table 2. Elemental components of analyzed points in Figure 3a-c by EDS analysis of electron probe.

² Relative standard deviations (RSD) are less than 3%.

Besides impurities in these inclusions of the quartz, some lattice elements in the hydrothermal quartz cannot be neglected. As shown in Figure 4, Götze (2009) [34] reported four substitutions between Si and impurity elements in a quartz lattice. In addition, substitution with interstitial charge compensator is suitable to be used for explaining occurrences of impurity elements in the hydrothermal quartz, because P mainly occurs in the apatite, and Ge is not detected by ICP analysis.



Figure 4. Schematic quartz structure showing the configuration of trace elements in the quartz lattice (modified from Götze 2009) [34]. Mclaren et al. (1983) proposed that the substitution of Si^{4+} by four H^+ is also possible (silanol groups) [35]. Since the illustration is two-dimensional, the fourth H^+ is not shown on the figure.

Figure 5 shows the surface distributions of Si, O, Al, and Na in the microarea of Figure 3d. Al and Na are uniformly distributed across the microarea. This indicates that the two elements replace Si in a quartz lattice in substitution with the interstitial charge compensator, although Li and B cannot be effectively detected by electron probe.

Based on a combination of optical microscope analysis and electronic microscope analysis, the main mineral inclusions in the quartz are muscovite, hematite and apatite. The muscovite contains Al, K, Mg, and Fe. The hematite contains Fe, while some hematite is associated with apatite. The apatite contains P and Ca. The associated Na, Mg, and Al might be from the wall rock of biotite plagioclase gneiss. For the main impurities Al, K, and Fe, the microscopic examination and component analysis indicate that Al and K mainly occur in muscovite, and Fe mainly exists in hematite. In addition, some Al and Na possibly occur in the quartz lattice. In addition, fluid inclusions can be possible sources for H, Na, and othe alkali and alkali Earth elements (and probably anions such as Cl). Most of the fluid inclusions contain gas and liquid phases, and are secondary. The quartz ore mainly contains quartz, muscovite, hematite, apatite, and secondary fluid inclusions, which are common features of hydrothermal deposits [36].



Figure 5. Surface distributions of Si, O, Al, and Na in the microarea of Figure 3d (analyzing for 12 min): luminance was enhanced for showing clearly: (**a**) the surface distribution of Si; (**b**) the surface distributions of O; (**c**) the surface distribution of Al; and (**d**) the surface distributions of Na. White spots represent elements at the corresponding positions of Figure 3d, and the scale bars are the same as Figure 3d.

3.2. Quartz Processing

Xiong et al. used pressure acid leaching with hydrofluoric acid as an efficient purification process for purifying the vein quartz [10]. However, in the conventional acid leaching process of quartz ore, quartz purification depends on an acid attack to gangue minerals, which results in high acid consumption and a long leaching time [10,11,22]. The two problems significantly reduce equipment life, increase production costs, and decrease production efficiency. Due to the pollution of the fluoric leaching agent [16,17], it is therefore necessary to develop an efficient fluorine-free leaching system to purify vein quartz. The objective of the recommended process is to realize a fluoride-free technology of pressure acid leaching at the same time as ascertaining quartz quality.

Calcination at 900 °C for 5 h is deemed to be effective to remove organic impurities and H_2O within fluid inclusions, and increase the exposure probabilities of mineral inclusions, including muscovite, hematite, and apatite, according to different thermal expansibilities [10,11,37]. In addition, calcination could destroy the crystal structures of gangue minerals, especially muscovite [37]. When combined with the calcination process, pressure acid leaching has the power to effectively purify quartz sand without any fluorides.

3.2.1. Recommended Process

Figure 6 shows two recommended processes for purifying the hydrothermal quartz. Quartz sand was calcinated at 900 °C for 5 h, and the calcinated quartz sand was washed by leaching and washing liquors. In a pressure-tight reaction kettle [10], the washed quartz sand was either leached by acid solutions containing H_2SO_4 and NH_4Cl , or HCl and NH_4Cl . Leached quartz sand was again washed by ultrapure water 5–10 times, and then dried at 105 °C for 3 h.



Figure 6. Engineering flow sheet of the recommended processes for purifying the hydrothermal quartz.

Following calcination, the first washing process is used to make full use of acids remained in leaching and washing liquors, and cut the cost effluent disposal. Using H_2SO_4 or HCl, calcinated gangue minerals can be efficiently dissolved and removed from quartz at high pressure and temperature. The NH₄Cl is used to provide hydrogen ions with a more stable and higher concentration during the hydrolysis of NH₄⁺.

Based on a combination of calcination and pressure acid leaching, the quartz sand was efficiently purified. The optimal process conditions and contents of impurity elements in quartz concentrates for two leaching processes are shown in Tables 3 and 4, respectively. The effective removal of Al, K, and Fe indicates that the main gangue minerals, including muscovite and hematite, are efficiently removed by the two processes. Parts of P and Ca were removed, but the removal rates were not so high. The possible reason is that some primary apatite is included in the quartz particle, and some P occurs in the quartz lattice. S, Ni, Zn, and As are not detected in quartz concentrates, which indicates that these impurity elements are liberated from quartz particles after ore grinding. Zr occurs in hydrothermal zircon due to the ultra-high pressure action in the Dabie Mountains [38], but it is difficult to be dissolved by acid solution [39], so its removal rates are only 0.155 wt %. The removal rates of some elements, including Li, B, Na, P, and Ti, are not so high because parts of them occur in the quartz lattice. As a result, the fluoride-free leaching processes have little influence on removing them.

Leaching Process	Temperature (°C)	Acid Concentration (HCl or H ₂ SO ₄) (mol·dm ⁻³)	NH ₄ Cl Concentration (mol·dm ⁻³)	Liquid/Solid Ratio (cm ³ ·g ⁻¹)	Leaching Time (h)
$HCl + NH_4Cl$	280	0.8	0.8	10	6
$H_2SO_4 + NH_4Cl$	250	0.3	0.45	5	7

Table 3. Optimal conditions of leaching processes 1 and 2.

In total

Element	Ore ($\mu g \cdot g^{-1}$)	Concentrate 1 ⁴ (µg·g ⁻¹)	Removal Rate 1 (wt %)	Concentrate 2 ⁴ (µg·g ⁻¹)	Removal Rate 2 (wt %)
Al	353	41.5	88.2	44.1	87.5
Fe	61.2	1.14	98.1	1.12	98.2
Na	13.4	11.3	15.7	12.2	8.96
S	5.64	_	_	_	_
Р	15.5	5.00	67.7	5.39	65.2
Li	2.20	2.19	0.455	2.04	7.27
Κ	118	1.21	99.0	2.24	98.1
Ca	8.04	4.54	43.5	4.40	45.3
Ti	8.31	4.89	41.2	5.38	35.3
Mg	11.8	4.88	58.6	7.15	39.4
Ni	1.01	—	—	—	_
Zr	6.46	6.45	0.155	6.45	0.155
Zn	0.567	—	—	—	_
As	3.16	_	—	_	_
В	10.8	8.77	18.8	8.77	18.8

Table 4. Contents and removal rates of major elements at optimal conditions in Table 3.

 4 Relative standard deviations (RSD) of element contents in quartz concentrates are less than 4%, and not shown in the Table; (—) = below the limit of detection.

85.2

99.2

84.0

91.9

3.2.2. Effects of the Calcination Process

619

Muscovite, as a main gangue mineral in the hydrothermal quartz, can be transformed into amorphous phases during the calcination process [37]. Figure 7 shows the surface topographies of muscovite in calcinated quartz sand. The planar water and constitution water were removed from the muscovite during calcination (Formula (2)), and the water evaporation increased the interlayer spacing of the muscovite [37], and further promoted a seperation of different muscovite sheets along the cleavage plane (Figure 7a). Muscovite flakes close to its surface were separated from the host, and some of them were damaged into fragments with micron sizes (Figure 7b). The fragments consisting of active K₂O, Al₂O₃, and SiO₂ (Formula (3)) could be easily dissolved by acid solutions. This demonstrates that calcination is a necessary procedure for removing muscovite by a fluoride-free acid leaching process.

$$K_2 O \cdot 3Al_2 O_3 \cdot 6SiO_2 \cdot 2H_2 O = K_2 O \cdot 3Al_2 O_3 \cdot 6SiO_2 + 2H_2 O^{\uparrow}$$

$$\tag{2}$$

$$K_2O \cdot 3Al_2O_3 \cdot 6SiO_2 = K_2O + 3Al_2O_3 + 6SiO_2$$
 (3)



Figure 7. Surface topographies of muscovite in calcinated quartz sand: calcination temperature 900 °C, calcination time 5 h; (**a**) backscattered electron image obtained by electron probe microanalysis (EPMA); (**b**) secondary electron image obtained by scanning electron microscope (SEM); SMS—seperated muscovite sheets along cleavage plane.

The calcination process is also effective for removing other mineral inclusions, such as hematite and apatite. The exposure probabilities of gangue minerals included in quartz grains have great influences on impurities removal. The impurities included in quartz particles are hardly removed unless a calcination process is used, especially in case of a fluoride-free purification process of quartz sand. The different expansion coefficients between quartz and hematite during calcination promoted the exposure of hematite. The calcination is also effective for apatite when it is associated with hematite.

Calcination is also an effective method for removing fluid impurities within quartz fluid inclusions. Schmidt–Mumm [40] thought that the decrepitation of the fluid inclusions could be classified as several processes: microcracks opening and propagating, the rupture of grain boundaries, transgranular fracturing, intragranular fracturing, and the decrepitation of large/small fluid inclusions. Figure 8 shows a decrepitation pit on the surface of a calcinated quartz particle. The small pit is about 7 μ m long and 1 μ m wide. It indicates that calcination at 900 °C for 5 h can even remove micron-size fluid inclusions. Volatile materials occurring in the fluid inclusions were removed during the calcination process, and inorganic salts dissolved in the fluids could be leached in the following acid leaching process.



Figure 8. Decrepitation pit on the surface of a calcinated quartz particle: calcination temperature 900 °C, calcination time 5 h.

At the optimal leaching conditions of processes 1 and 2 in Table 3, the element contents in quartz sand only after pressure acid leaching (rather than using a calcination process) were compared with those of quartz concentrates so as to describe the effect of the calcination process on the removal of impurity elements. As shown in Table 5, the calcination process can effectively promote the leaching of Al, Fe, S, P, K, and As. The contents of Al and K were respectively reduced from 167 and 188 $\mu g \cdot g^{-1}$ to 41.5 and 44.1 $\mu g \cdot g^{-1}$ by an additional calcination process. The differences indicate that the calcination process by destroying their crystal structures. In addition, the decrease of the Fe and P contents may be caused by exposure of the hematite and apatite during calcination. S and As could easily be removed during calcination due to their volatility. Moreover, the decrease of Ca and Na contents might be caused by the decrepitation of fluid inclusions (compare to Figure 8).

Table 5. Contents of major elements in quartz sand with or without a calcination process.

Element	Ore ($\mu g \cdot g^{-1}$)	C and PHAL ($\mu g \cdot g^{-1}$)	PHAL ⁵ ($\mu g \cdot g^{-1}$)	C and PSAL ($\mu g \cdot g^{-1}$)	PSAL ⁵ ($\mu g \cdot g^{-1}$)
Al	353	41.5	167	44.1	188
Fe	61.2	1.14	7.42	1.12	13.4
Na	13.4	11.3	13.5	12.2	12.9
S	5.64	—	4.00	_	4.52
Р	15.5	5.00	13.6	5.39	10.4
Li	2.20	2.19	2.20	2.04	2.19

K	118	1.21	42.6	2.24	45.9
Ca	8.04	4.54	7.50	4.40	8.01
Ti	8.31	4.89	6.07	5.38	5.91
Mg	11.8	4.88	8.60	7.15	8.12
Ni	1.01	_	_	_	_
Zr	6.46	6.45	6.46	6.45	6.46
Zn	0.567	—	_	_	_
As	3.16	—	1.47	_	2.15
В	10.8	8.77	9.80	8.77	8.47
In total	619	91.9	290	99.2	316

Table 5. Cont.

⁵ Relative standard deviations (RSD) of the element contents in quartz concentrates are less than 4%, and not shown in the Table; C—calcination process (900 °C for 5 h), PHAL—pressure acid leaching (HCl + NH₄Cl), PSAL—pressure acid leaching (H₂SO₄ + NH₄Cl); (—) = below the limit of detection.

3.2.3. Effects of Fluoride-Free Pressure Acid Leaching

Figure 9 shows the surface topographies of muscovite in leached quartz sand. Compared with the surface topographies of muscovite (Figure 7) in calcinated quartz sand, the fragments with micron sizes were dissolved and removed from the muscovite surface (Figure 9a). This process can be described by the dissolution of the active fragments containing K₂O and Al₂O₃ (Formulas (4) and (5)). In addition, the muscovite in calcinated quartz sand was further dissolved along its edge. More importantly, some fractures were developed in the vertical direction of the long side of muscovite when concentrations of leaching agents were doubled (Figure 9b). Moreover, expanded interlayer spacing provided favorable conditions for leaching reactions, and promoted a faster process of muscovite dissolution. The several effects of acid leaching led to the structure collapsing and the dissolution of muscovite.

$$K_2O + 2H^+ = 2K^+ + H_2O$$
(4)

$$Al_2O_3 + 6H^+ = 2Al^{3+} + 3H_2O$$
(5)



Figure 9. Surface topographies of muscovite in leached quartz sand: (a) secondary electron image obtained by SEM, leaching conditions: $0.05 \text{ mol}\cdot\text{dm}^{-3} \text{ H}_2\text{SO}_4$, $0.10 \text{ mol}\cdot\text{dm}^{-3} \text{ NH}_4\text{Cl}$, liquid/solid ratio 5 cm³·g⁻¹, leaching time 7 h, and leaching temperature 200 °C; (b) backscattered electron image obtained by EPMA, leaching conditions: $0.10 \text{ mol}\cdot\text{dm}^{-3} \text{ H}_2\text{SO}_4$, $0.20 \text{ mol}\cdot\text{dm}^{-3} \text{ NH}_4\text{Cl}$, liquid/solid ratio 5 cm³·g⁻¹, leaching time 7 h, and leaching temperature 200 °C.

During calcination, most of the mineral inclusions in quartz sand were transformed into mineral intergrowths of quartz and gangue minerals due to heat stress. Then they were exposed in the leaching solution. With further calcination, the mineral intergrowth was promoted into separated

mineral monomers. Figure 10 shows the surface topographies of calcinated and leached quartz sands. Although associated minerals were liberated from quartz particles in Figure 10a, some impurities unavoidably remained in the pores of the calcinated quartz surface. The impurities that remained on the surface of calcinated quartz were leached by mixed leaching agents. As a result, the surface of the leached quartz sand in Figure 10b became smoother than that of the calcinated quartz sand in Figure 10a. Common intergrowths of quartz–hematite and quartz–hematite–apatite were leached by mixed leaching agents, and the leaching processes are as follows:

$$Fe_2O_3 + 6H^+ = 2Fe^{3+} + 3H_2O$$
(6)

$$Ca_{5}[PO_{4}]_{3}(OH) + H^{+} \rightarrow Ca^{2+} + H_{3}PO_{4} + H_{2}PO_{4}^{-} + HPO_{4}^{2-} + PO_{4}^{3-} + H_{2}O$$
(7)



Figure 10. Surface topographies of quartz sand: (a) calcinated quartz sand (900 °C for 5 h); (b) quartz concentrate of process 2.

By a combined process of calcination and fluoride-free acid leaching, gangue minerals including muscovite, hematite, and apatite were effectively separated from the hydrothermal quartz. The calcination process not only destroyed the crystal structures of the gangue minerals; it also removed volatile compounds. Meanwhile, the exposure probabilities of the impurities were enhanced due to the calcination process. In general, the calcination process had a positive influence on the removal of impurity elements with the fluoride-free pressure acid leaching process.

4. Conclusions

The present study analyzed the geological formation and mineralogy of the hydrothermal quartz deposit from Hengche in Qichun County, of the Hubei Province in China, A combined purification process consisting of calcination and fluoride-free pressure acid leaching was developed based on two fundamental investigations. Moreover, the effects of this processing procedure on the raw quartz material were discussed, and the following conclusions obtained:

(1) Impurity elements in the hydrothermal quartz are related to fluid inclusions and associated gangue minerals such as muscovite, hematite, and apatite. The muscovite not only occurs along the grain boundaries of quartz particles; it is also included in quartz particles. The hematite is mainly of secondary origin, and occurs along crystal fractures in quartz. However, some of the hematite is also associated with apatite.

(2) The gangue minerals such as muscovite, hematite, and apatite were effectively removed by a combined process of calcination and fluoride-free pressure acid leaching, while the removal rates of the most important impurity elements Al, K, and Fe were 88.2 wt %, 99.0 wt % and 98.1 wt %,

respectively, for the HCl-NH₄Cl system, and 87.5 wt %, 98.1 wt % and 98.2 wt % for the H_2SO_4 -NH₄Cl system. The removal rates of some elements, including Li, B, Na, P and Ti were not so high, because they might be structurally incorporated in the quartz lattice.

(3) The calcination process not only destroyed the crystal structure of the gangue minerals, it also removed volatile compounds. The liberation probabilities of impurities were enhanced due to the calcination process. In conclusion, the calcination process had a positive influence on the removal of impurity elements during the fluoride-free pressure acid leaching process.

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