Manganese and limestone interactions during mine water treatment

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Abstract

Manganese removal from mining-affected waters is an important challenge for the mining industry. Addressed herein is this issue in both batch and continuous conditions. Batch experiments were carried out with synthetic solutions, at 23 ± 2 °C, initial pH 5.5 and 8.3 g limestone/L. Similarly, continuous tests were performed with a 16.5 mg/L Mn²⁺ mine water, at 23 °C, initial pH 8.0 and 20.8 g limestone/L. Calcite limestone gave the best results and its fine grinding proved to be the most effective parameter for manganese removal. In either synthetic solutions or industrial effluents, the final manganese concentration was below 1 mg/L. A change in limestone surface zeta potential is observed after manganese removal and manganese carbonate formation was suggested by IR spectroscopy. The conclusion is that limestone can remove manganese from industrial effluents for values that comply with environmental regulations.

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1. Introduction

The removal of manganese and sulfate from mining-affected water is a important challenge for water management in the mining industry, as these specie are not solely removed by pH control. Manganese is an important trace element for the functioning and activation of many enzymes (manganese superoxide dismutase, kinases, decarboxylases, among others) in the human body, but at high levels, it can cause damage to the brain, liver, kidneys and nervous system [1]. As a consequence, many environmental agencies worldwide have regulated limits for manganese concentration in water. For instance, the USEPA has set the level of manganese in drinking water at 0.05 mg/L. Similarly, its maximum level, according to the Brazilian legislation, is 0.01 and 1 mg/L for surface water and wastewater, respectively.

Manganese removal from wastewater depends on: the concentration and speciation encountered in the effluent; the volume of wastewater to be treated and economical aspects. In addition to manganese, mine water usually contains a series of other elements such as iron, aluminum, copper, and zinc [2] and most of these metals can be removed by increasing the pH; while iron is readily precipitated through oxidation to the trivalent state using either active or passive systems. Manganese can also be removed by oxidation, since Mn(IV) compounds are insoluble. Notwithstanding, manganese (II) is soluble over a wide pH range. Its chemical oxidation is kinetically slow [3], and due to the high reduction potential of Mn(IV), parallel reactions, such as organic matter and Fe(II) oxidation, induces excessive oxidant consumption. Therefore, iron removal is required prior to manganese oxidation [4]. The use of biological processes involving manganese (II) oxidation [3,5] is an alternative treatment for manganese-affected water.

Carbonate sources such as limestone can be successfully applied for manganese removal, since they are inexpensive, effective [6,7] and widely available. This method has been widely used as an alternative for treating contaminated drinking water [8–10] and also for treating acid mine drainage (AMD) [11]. More than 80% of the heavy metals, such as: copper, iron, manganese, cadmium and others can be removed using either batch or continuous-flow limestone filtration processes [12]. Aziz and Smith [8,13] studied manganese removal (1 mg/L) from drinking water using different solid media such as limestone, brick powder and gravel in both batches and in columns. Out of the three materials, limestone showed the best results with more than 95% Mn(II) removal. Similarly, Thornton [10] and Rose et al. [14] have shown that limestone can be applied with relative success to remove manganese from acid mine drainage, although limestone beds are not the best option because they are usually clogged with organic matter (algae, leaves), silt, Fe- and Al-hydroxides and gypsum, among others.

Although metal carbonate formation seems obvious when applying limestone to the treatment of manganese-affected water, such compounds have seldom been identified, especially during the treatment of AMD. One of these species was kutnahorite, CaMn(CO₃)₂, which is thermodynamically stable in such waters, provided Ca²⁺ is available [4,15]. Due to the complexity of the different processes occurring during the passive treatment of
AMD (wetlands), manganese removal as oxyhydroxides derived from either abiotic or biotic Mn(II) oxidation has been proposed. Manganese oxyhydroxides (MnOOH) were noticed in reactors containing different substrates (dolomite, magnesite, limestone, quartzite) [16], while nearly 90% of the manganese present in a carbonate precipitate from an AMD pond was in the form of oxides [15].

Seeking to shed new light on the role of limestone in manganese removal, the use of fine ground limestone powder as a cost-effective, simple alternative to remove manganese from mining-affected water was studied. An attempt to characterize the manganese removal mechanism was also carried out using zeta potential and infrared spectroscopy.

2. Materials and methods

Calcite and dolomite limestone samples were kindly provided by Brazilian producers. These samples were dry-sieved or alternatively dry-ground in a vibrating cup mill (Pulverisette 9, Fritsch) to produce particle size distributions with different d50 values. These particle size distributions were assessed by laser diffraction (Mastersize, 2000E). Chemical analysis (ICP-OES, Spectro) was also performed on all limestone samples to determine their nature. Table 1 presents the chemical analysis of three different samples showing their calcium and magnesium content, so that the composition of the limestone could be defined, i.e. dolomite limestone presents higher than 5% magnesium, while calcite limestone has less than 5% magnesium content. In addition, the limestone nature was confirmed by XRD analysis (Shimidzu XRD 6000).

The effects of both calcite and dolomite limestone as well synthetic CaCO3 on manganese removal were assessed in both batch and continuous experiments. Batch manganese removal was assessed with 120 mL synthetic Mn2+ solutions (MnCl2 · 4H2O, Synth) stirred with limestone powder in capped erlenmeyer (250 mL capacity) flasks in an orbital shaker at 350 m-1 and 23 ± 2 °C, for 60 or 90 min. Unless otherwise stated, a 8.3 g/L pulp density and initial pH = 5.5 were applied throughout the experiments. At the end of the experiment, the pH was measured (it reached values around 9.0), the pulp was filtered and the manganese concentration in the aqueous phase was determined by atomic absorption spectrometry (AAS) (Perkin Elmer AAnalyst 100). The manganese removal efficiency was determined by mass balance.

Continuous experiments were carried out in a 1.64 L capacity MSMPR reactor, at 23 ± 2 °C. The reactor was fed separately with (i) mine water and (ii) a pulp made up of limestone and distilled water to achieve a solid content of 20.8 g/L limestone. After proper pH adjustment, both flows were pumped at 16.0 mL/min (mine water) and 3.0 mL/min (limestone pulp) into the reactor. Residence time was set at 60 min and the experiments ran for at least 12 residence times to ensure steady state conditions. After every 60 min, an aliquot was withdrawn from the reactor, filtered and its zeta potential and infrared spectroscopy determination. The mine water was stored in plastic containers at room temperature and filtered prior to each experiment. Its pH was adjusted with a 6 mol/L NaOH solution and the manganese content was determined by AAS (16.5 mg/L).

Scanning electron microscopy (SEM) was used to examine the morphology of limestone particles. The samples were coated with graphite by electro-deposition, using a Jeol JEE 4C instrument and analysed by a JEOL JSM 5510 scanning electron microscope (SEM), with an accelerating voltage 0.5–30 kV, equipped with a spectrometer for micro-analysis based on an energy dispersive X-Ray spectroscopy system (EDS).

FTIR spectra were registered on a Nicolet Nexus 470 model spectrophotometer equipped with a Centaurus microscope, an attenuated total reflectance (ATR) apparatus (Thermo) and a ZnSe internal reflection devise. Medium infrared measurements were produced with a KBr bean splitter and an HgCdTe detector. The spectra were pressed against a flat glass surface and mounted under the ATR apparatus. The spectra were collected in the 400–4000 cm−1 region with a minimum of 32 scans at 4 cm−1 resolution.

After pulverizing the limestone samples (d50 < 10 μm), zeta potential was assessed in a Zetazizer Nano ZS zetameter (Malvern). Prior to the readings, a 0.5% solid pulp was prepared with deionised water and limestone samples (one, before, and another, after mixing with a 60 mg/L Mn solution). Diluted HCl and NaOH solutions were used to control the pH.

3. Results and discussion

The effect of limestone composition on manganese removal from a 60 mg/L Mn(II) synthetic solution is presented in Fig. 1, for <45 μm particle size. The dolomite limestone removed 18.6% Mn(II), whereas the calcite limestones I and II removed 100% and 13.5% Mn(II), respectively. The results achieved with these limestone samples are compared with that produced by a chemically precipitated CaCO3, where 100% removal was observed. When comparing dolomite and calcite limestone, the latter shows better performance. This is because powdered dolomite limestone is less reactive [6] than the calcite limestone and this is consistent with the results achieved for drains, designed for AMD treatment, constructed from limestone of different origins [11] and composition [17]. Furthermore, the particle size distribution of calcite limestone shows a higher proportion of fine particles (75% < 20 μm) as compared to calcite limestone II (20% < 20 μm) which facilitates manganese removal (Fig. 2). Similarly, the high extraction observed with synthetic CaCO3 can be also ascribed to its low particle size (100% < 20 μm) and therefore higher surface area (9.7 m2/g). These results are corroborated by those observed by Letterman et al. [18],

<table>
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<th>Table 1 Chemical analysis of the limestone samples.</th>
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<td>Limestone</td>
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<td>Calcite I</td>
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<tr>
<td>Calcite II</td>
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<td>Dolomite</td>
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Fig. 1. Effect of limestone type on the removal of manganese in batch experiments. 600 mg/L initial manganese concentration; 8.3 g/L limestone; initial pH 5.5; 23 ± 2 °C, particle size: <45 μm.
Potgieter-Vermaak et al. [19] and Sun et al. [20]. For the material that enabled high manganese removal, a pulp pH of 9.0 was the highest value attained, therefore suggesting that to some extent, limestone dissolution accounts for manganese removal [8]. As calcite limestone I showed the best results, it was selected for further experiments, being hereafter referred to as limestone.

After this selection, a series of experiments was performed to determine the effects of time, pulp density and initial Mn(II) concentration on manganese removal from a synthetic solution, at an initial pH of 5.5 (Fig. 3). Manganese removal with limestone is relatively fast with 100% extraction in 60 min (Fig. 3a) which is consistent with the work of Aziz and Smith [8]. In the present work, experiments carried out at 70 °C removed manganese slightly slower than those at 23 °C, i.e. time and temperature have significant effects on manganese removal, as also observed by Ghaly et al. [16]. As expected, the higher the limestone content in the pulp, the higher the pH and the greater the manganese removal [16]. From a solution containing 60 mg/L Mn(II), the removal increased with an increase in the limestone content up to 8.3 g/L and levelled out above this value (Fig. 3b). Additionally, this 8.3 g/L pulp density was able to remove manganese in the range of 0–155 mg/L Mn(II) [14] (the latter represents the highest concentration usually found in mine waters) producing final effluents with manganese concentrations below 1 mg/L. However, these results were not reproduced when experimenting with the mine water sample. No appreciable removal was observed at an initial pH of 5.5 and 42.6% manganese removal was accomplished using 8.3 g/L limestone from a solution containing 16.5 mg/L Mn(II), at pH 8.0 (initial), producing a final effluent containing 8.0 mg/L Mn^{2+} (Table 2). Therefore, the limestone content had to be increased to 20.8 g/L to produce a final solution with manganese concentration below 1 mg/L and pH 8.8.

Table 2
<table>
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<th>Limestone content (g/L)</th>
<th>Final Mn^{2+} concentration (mg/L)</th>
<th>Effluent final pH</th>
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<tr>
<td>8.3</td>
<td>8.0</td>
<td>7.7</td>
</tr>
<tr>
<td>16.7</td>
<td>0.3</td>
<td>8.5</td>
</tr>
<tr>
<td>20.8</td>
<td>&lt;Q.L.</td>
<td>8.8</td>
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Since powdered limestone is not suitable for fixed bed column applications, continuous experiments were performed to assess manganese removal from the mine water (Fig. 4) on a 1.64 L MSMPR reactor with 60 min residence time, at 23 ± 2 °C. In these experiments, the effect of the initial solution’s pH was also assessed, since it was observed that low pH waters affected the performance of manganese drains which were treating acid mine drainage [10,21]. Fig. 4 shows that 20.8 g/L limestone removes only 13% (average) of the manganese present in the mine water at its natural pH (3.3).
Manganese removal increases to 56% when the mine water’s initial pH is increased to 5.0 before the addition of limestone. The manganese removal increases further to 97% when the initial pH is 8.0, resulting in less than 1.0 mg/L residual manganese concentration in the final effluent. These results confirm improved manganese removal when pH increases as most manganese compounds are not stable at low pH [14]. Fig. 4 also shows that the process is relatively fast with metal removal in the first residence time (60 min) since the manganese concentration is already below 1.0 mg/L, remaining throughout the experiment, while the final pH reaches the 8.3–8.5 range. These results are consistent with the work of Thornton [10], who studied the treatment of mine water having an average Mn(II) concentration of 3.5 mg/L. The experiments were run continuously for 495 days, being the manganese concentration reduced to 0.49 mg/L and the pH increased from 6.4 to 7.6. A black film coated the limestone which was ascribed to the likely presence of manganese-oxidizing microorganisms in that system. Discussing the effectiveness of limestone beds to remove manganese, Rose et al. [14] analyzed a series of experiments treating AMD with manganese concentrations ranging from 5 to 30 mg/L and observed that most of these systems were able to reduce the manganese concentration to less than 1 mg/L. These results clearly show that increasing the pH of manganese-affected wastewater has a beneficial effect on manganese extraction, regardless if synthetic solutions or mine water is treated. This reinforces the idea that manganese removal should follow a previous step where the pH is increased; some metals are precipitated before limestone addition.

One of the reasons for the poor performance observed with the mine water would be the armoring effect [22] of the limestone surface as several works have shown that precipitation of iron and aluminum compounds on the limestone surface reduces its dissolution. However, as shown in Table 3, the mine water, after pH
correction and before limestone addition, has a low concentration of these elements. It also shows that the manganese concentration was reduced from 16.5 to 0.3 mg/L and the other heavy metals concentration is even lower after contacting with limestone. Interestingly, the calcium concentration did not increase suggesting a small amount of limestone dissolution as observed elsewhere [10], which is supported by a decrease in the particle size (Fig. 5). The sample d_{50} was reduced from 8.0 to 5.5 μm with a concomitant increase of the specific surface area from 3.4 to 4.3 m^{2}/g. The stability of the Mn carbonate system is currently being addressed.

To gain further insights on the limestone interaction with the mine water, X-ray energy dispersive spectra of manganese, calcium and sulfur were produced with the residues of the continuous experiments (Fig. 6). For the particle shown in Fig. 6a, Fig. 6b depicts the manganese mapping (dots); while those of calcium and sulfur are presented in Fig. 6c and d. It can be noticed that calcium covers the whole surface of the calcite limestone, as expected, while manganese is spread over the limestone grains and it is not concentrated in any region of the different limestone particles. Although sulfur concentration in the limestone surface (Fig. 6d) is lower than that of calcium (Fig. 6c), calcium sulfate precipitation on the limestone surface would likely account for the lower manganese removal observed in the experiments with the mine water as compared to those performed with synthetic solutions. Notwithstanding, the armoring effect produced by the presence of sulfur on the limestone surface requires further investigation.

3.1. Manganese species on the calcite surface

The interaction of manganese and calcium carbonate was studied with the material produced in the batch experiments using synthetic solutions, since a higher removal of manganese was observed. Firstly, SEM-EDS was applied to characterize the limestone particles before and after manganese removal. It showed no heavy metal on the solid limestone before the experiment, since only carbon, oxygen and calcium were observed in the EDS spectrum (Fig. 7a). In the batch experiments, the presence of manganese on the surface of the calcite limestone is readily seen (Fig. 7b) after contact with the manganese-containing solution. Following, these materials were characterized by infrared spectroscopy. Due to the low manganese concentration on the limestone surface, attenuated total reflectance (ATR) was applied so that the identification of the manganese species could be accomplished. ATR permits the identification of functional groups on the surface of the mineral particle, especially surface complexation and changes on the polar functional groups [23]. Fig. 8 presents the spectra of synthetic manganese carbonate, the limestone sample, and the residues of both batch and continuous manganese removal experiments. Characteristic absorption bands ascribed to calcite limestone are observed at 712–713 cm\(^{-1}\) in all samples, irrespective of the presence of manganese. Manganese carbonate presents an absorption band at 724 cm\(^{-1}\), which is consistent with the findings of Nassrallah-Aboukais et al. [24] who observed bands at 725 and 727 cm\(^{-1}\) assigned, respectively, to synthetic and natural MnCO\(_3\) (rhodochrosite) during manganese precipitation on calcite carbonate. Fig. 8 also shows a weak band at 729 cm\(^{-1}\) (which is 2 cm\(^{-1}\) higher than that observed for rhodochrosite [24]) that is proposed to be ascribed to an manganese carbonate phase formed on the surface of the calcite limestone. The shift on band position as compared as the bulk MnCO\(_3\) would be derived from the effect of the calcium atoms on the manganese carbonate bonds. This type of shift has also been observed in different adsorption systems [25,26].

The mechanisms of manganese removal may include occlusion, surface adsorption, or trace phase formation [27]. An insight on that may be offered by the analysis of the surface charge of the limestone particles. Fig. 9 depicts zeta potential measurements of both the raw and manganese-containing limestone. It can be seen that the pH_{pzc} of the raw sample is 9.8, which is consistent with the range observed in literature [28,29]; a result of the nature of the potential determining ions, Ca\(^{2+}\) and CO\(_3^{2-}\) [30]. Many researchers have
proposed models for the calcite mineral–aqueous solution interface and a review in this topic is beyond the scope of the present work. Nevertheless, the presence of manganese on the limestone surface decreases the pH_{pzc} to 7.2 suggesting chemical adsorption. Considering the pH of the system at the end of the experiment (around 9.0), it is expected that the limestone surface would be positively charged. Additionally, a speciation diagram using manganese hydroxy–complex formation constants available in the NIST database [31] shows the predominance of aqueous Mn^{2+} and MnOH^{+} in the pH range of manganese removal. Therefore, a physical adsorption mechanism would not explain the removal process, supporting the chemical adsorption mechanism. It is interesting that a study on rhodocrosite (MnCO_{3}) surface properties has determined the pH_{pzc} of that compound as 7.4 [32] which is similar to the value observed in the present work for the Mn-containing limestone. Comparing the removal efficiency of limestone-sandstone mixtures with that of limestone alone, Ghaly et al. [16] stated that the former performs better than the latter for the removal of iron from AMD, due to the more negative surface charge of sandstone [20]. However, the authors did not notice any improvement in the removal of manganese with that mixture, which is consistent with the findings of the present work.

The interaction between manganese and limestone is complex. Although manganese carbonate would be expected in the presence of the anion, the formation of oxides is the rule. Even under those conditions where the water presented a significant concentration of carbonates, manganese oxides were suggested as the predominant species, favored by highly dissolved oxygen concentrations (forced aeration) as well as high pH [33], while carbonates were frequent only at high activities of dissolved carbon dioxide [15]. In this regard, when carbonate dissolution provided high calcium concentrations, kutnahorite (CaMn(CO_{3})_{2}) was precipitated [4] as geochemical modeling predicts that some mine water would be supersaturated with respect to this mineral. The formation of oxides over carbonates can also be partially ascribed to the presence of manganese-oxidizing bacteria in many studies [4,11,16]. Likely, the time scale of the experiments may play a key role, since manganese removal experiments in biotic systems last longer (3 months, for instance [16]) than those performed in the present work. Previous studies of Mn^{2+} adsorption on calcite indicate that manganese was adsorbed at concentrations below those defined by the solubility product of MnCO_{3}. However, McBride [34] as well as Franklin and Morse [35] studied the interaction of Mn^{2+} with the surface of calcite in dilute aqueous solutions and proposed a precipitation reaction at high manganese concentrations, supported by electron spin resonance (ESR) studies. The results of the present work suggest manganese was not electrostatically adsorbed but instead was substituting for calcium on the limestone surface as proposed by Pingitore et al. [27].

4. Conclusions

Limestone powder was effective in removing manganese from both synthetic and mine waters. The results showed that both fine grinding as well as the initial water pH defines the effective manganese removal. For synthetic solutions, up to 155 mg/L manganese could be removed in 60 min for an initial pH of 5.5 and 8.3 g CaCO_{3}/L reaching final concentrations below 1.0 mg/L. The removal of manganese from mine waters (16.5 mg/L) was more difficult, requiring an initial pH of 8.0 and a higher limestone concentration (20.8 mg/L), likely to the deleterious effect of calcium sulfate armoring on the limestone surface. Manganese was chemically adsorbed without significant oxidation as suggested by AFT-IR. Manganese removal with limestone should therefore follow a neutralization step to increase the effluent pH so that efficient metal removal can be accomplished.

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References