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Ammonia-Ca-K Competitive Ion-Exchange on Zeolites in Mining Wastewater Treatment: Batch Regeneration and Column Performance

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Abstract

This manuscript addresses the treatment of explosives-impacted mining wastewaters (EIMWW) using ion-exchange to remove elevated levels of ammonia. Repeated batch loading-regeneration cycles were conducted for two commercially available zeolite media used in the treatment of ammonia-laden EIMWW to establish the effects of competing ions and regeneration solution composition. The Northern Ontario EIMWW tested contained 3.87 meq/L total ammonia (TA) as well as 2.85 mg/L K⁺ and 3.9 meq/L Ca²⁺. The media studied were a natural clinoptilolite and a modified clinoptilolite (SIR-600). Five regenerant solutions with different NaCl and KCl concentrations were evaluated using batch tests. The presence of potassium in the regenerant was found to hinder the TA exchange capacity of both zeolites. The SIR-600 and the natural clinoptilolite used in conjunction with the 10% NaCl solution featured the best TA exchange capacities, 0.46 ± 0.02 meq TA/g and 0.36 ± 0.05 meq TA/g, respectively. The batch tests showed that both media had a slight preference for K⁺ over TA. The continuous flow column tests performed using SIR-600 media greatly accentuated the selectivity of K⁺ over TA. In reaching the same 0.55 meq TA/L breakthrough level, the same modified zeolite column was able to treat five times more volume of a synthetic TA solution than EIMWW.

Keywords: ion exchange, ammonia, clinoptilolite, mining wastewater, competitive ion exchange, potassium

1. Introduction

The mining industry extensively uses nitrogen-based explosives, with ammonium nitrate mixed with fuel oil (ANFO) being the primary blasting agent. Cyanide destruction, transformation of amines in flotation circuits, pH regulation agents, ammonium sulphate as eluent of uranium from ion exchange resins, ammonium hydroxide used in uranium precipitation, and ammonia used to recover copper and nickel in hydrometallurgical processes are other sources of ammonia in mining water [1–4]. These result in mining effluents containing elevated concentrations of ammonia. Ammonia discharges can be toxic to fish and can result in substantially lower dissolved oxygen levels in receiving waters that also greatly impact aquatic life. Accordingly there are an increasing number of ammonia discharge regulations around the world to curb the discharge of ammonia into the environment [5–8]. This has led to the research and development of multiple treatment options for the removal and/or transformation of ammonia, while aiming to minimize subsequent environmental consequences. Biological nitrification-denitrification, which is quite common in domestic wastewater treatment, is not the best choice for mining wastewater treatment because it is significantly impacted by low temperatures, it does not perform as well when there are large fluctuation in the influent ammonia
concentrations, and it is impacted by ammonia toxicity [6,9–11]. These conditions make physicochemical treatment methods particularly suitable, ammonia removal has been achieved through adsorption/ion-exchange, air stripping, microwave radiation, reverse osmosis, sonification, and chemical processes such as chemical precipitation or breakpoint chlorination [6,7,9,10,12–17]. Air stripping, which may be economical in warm climates, is not efficient in the cold temperatures encountered in Northern Canadian mines. This is because under normal conditions most of the ammonia is present as ammonium (NH$_4^+$), at low temperatures an even smaller fraction of the total ammonia (TA) is present in the strippable NH$_3$ form, and the mass transfer of NH$_3$ is slower at these very low temperatures. It also requires chemicals to raise the pH but their transportation costs are high due to the remoteness of these mine sites. On the other hand, ion exchange removes the ionized form of ammonia (NH$_4^+$) so it is naturally favored for cold weather applications.

Among the physicochemical methods, ion exchange (IE) is a good candidate for the removal of ammonia from explosives-impacted mining wastewaters (EIMWW). IE is an effective technology because at near neutral pH conditions ammonia is primarily present as ammonium (NH$_4^+$) ions [5]. It also offers the advantages of fast start-up times, low-cost, simplicity of application and operation.

Removal of ammonia by IE, particularly zeolites due to their high affinity for ammonia and low cost, is a well-documented process [7,14,15,17–22]. Ames proposed the following general order of selectivity for clinoptilolite, a common zeolite [23]:

\[
\text{Cs}^+ > \text{Rb}^+ > \text{K}^+ > \text{NH}_4^+ > \text{Ba}^{2+} > \text{Sr}^{2+} > \text{Na}^+ > \text{Ca}^{2+} > \text{Fe}^{2+} > \text{Al}^{3+} > \text{Mg}^{2+} > \text{Li}^+
\]

This order of selectivity has been studied and confirmed by numerous studies [20,24]. It should be noted that the majority of the NH$_4^+$-IE literature deals with the IE treatment of synthetic and/or single solute ammonia solutions rather than real wastewaters that contain numerous other ions. In addition, the ammonia exchange capacities reported in the literature vary widely in part due to the many differences amongst different zeolite types, the differences among the solutions being treated, and operational conditions such as pH and temperature [17–19,22].

A literature search did not identify studies on the IE treatment of this type of explosives impacted wastewater. A critical characteristic for the IE treatment of this type of wastewater is that it has significant concentration of ammonia in addition to other ions, such as Ca$^{2+}$ and K$^-$. As zeolites generally have a slightly higher selectivity for K$^+$ than NH$_4^+$, the treatment of multi-solute solutions has often resulted in lower NH$_4^+$ removals than those reported for single solute ammonia solutions due to the competition for exchange sites [17,20,21,25]. However, one study reported the effect of the other cations upon uptake of ammonium ion being relatively small [7]. According to [26] the simultaneous uptake and elution behaviour of NH$_4^+$ and K$^+$ on clinoptilolite are poorly documented.

The overall objective of this study was to evaluate the effectiveness of two IE media in the removal of TA from a real EIMWW. The specific objectives of this study were: a) to study the effects of regeneration solution composition on the removal of ammonia from EIMWW using a natural clinoptilolite and a modified clinoptilolite using batch tests, and b) to study the competitive ion exchange within a continuous flow column, which is the more common IE treatment system configuration. Competitive ion exchange is believed to be a significant factor on the feasibility of an IE system for treating EIMWW. The majority of studies have focused on the conventional sodium or acid-based regenerations, but it is believed that the selectivity order can be partially circumvented by greatly increasing the concentration of a less preferred ion [14]. Accordingly, this study will evaluate the impact of using both sodium and potassium regeneration solutions.

2. Experimental Methods and Materials

The experimental plan consisted of performing batch and continuous flow column experiments using real EIMWW in an attempt to assess the performance of an IE process, and to establish the effects of ionic competition. A key objective of the batch experiments was to identify the most suitable regenerant among several combinations of NaCl and KCl solutions. The column experiments used the best regeneration solution identified during the batch tests.

2.1. Materials

The EIMWW was collected from a blasting operation in a Northern Ontario mine, and filtered in the lab prior to the testing. For simplicity, the ammonia concentrations will be quantified in terms of total
ammonia (TA) as the sum of ammonium ion (NH₄⁺) and unionized ammonia (NH₃). The EIMWW contained approximately 3.87 meq TA/L (70 mg TA as NH₄⁺/L) and a complex blend of dissolved metals and anions. Full chemical analysis was accomplished via Inductively Coupled-Plasma (ICP) analysis. TA, potassium and calcium were determined to be the major three cations in EIMWW that are most relevant based on the clinoptilolite selectivity, therefore their interactions were the central focus of the study. For comparison of the impact of competition among ions, one column test was performed with a single-solute synthetic wastewater feed, which was prepared by dissolving technical grade (NH₄)₂SO₄ (Fisher Sci., Fair Lawn, NJ) in distilled water (230 mg in 1 L) for the desired concentration of approximately 3.87 meq TA/L.

The two IE materials considered were both zeolitic in nature. Clinoptilolite is inexpensive and known for its molecular sieving abilities as well as its high affinity for ammonia [14,27]. A natural zeolite (mineral clinoptilolite) was obtained from The Seed Supply co. (Saint Anthony, ID). According to the manufacturer, this product was 85% clinoptilolite with a cation exchange capacity of 1.4-1.65 meq/g and a maximum water retention of 55% of its weight. The bulk density was approximately 881-961 kg/m³ [28]. The rock elemental composition was reported by the supplier [28], it consisted primarily of SiO₂ (67.4%), Al₂O₃ (10.6%), K₂O (4.19%), CaO (2.23%) and Fe₂O₃ (1.7%). The material had a surface area of 24.2 m²/g, based on Brunauer-Emmet-Teller (BET) analysis at the Chemistry Department, University of Ottawa. The second IE material studied was a modified clinoptilolite (SIR-600, Resintech, West Berlin, NJ, USA). This modified clinoptilolite is typically utilized in the removal of cesium and ammonia from wastewaters [29]. Both materials were dried in an oven at 55 °C to remove excess moisture and cooled in desiccators. SIR-600 and the clinoptilolite were selected for this study because an earlier work showed that they had better TA removals from the same EIMWW than three different synthetic ion exchange media tested [30].

The purpose of the regenerations studies was to establish the best regeneration solution for the subsequent continuous flow column tests. The regeneration solutions were prepared using analytical grade KCl and NaCl (BDH, Mississauga, Canada). The ion exchange media was washed prior to use, but not preconditioned.

2.2. Analytical Methods

TA measurements were accomplished by nesslerization as per Standard Method 4500-NH₃ [31] and a spectrophotometer (DR6000, HACH, Loveland, CO, USA). pH was recorded on a pH probe (PHC201, HACH, Loveland, CO, USA) and electrical conductivity (EC) was measured with a probe (40PCID, VWR, Radnor, PA, USA) and a benchtop meter.

Potassium and calcium concentrations were measured by direct air-acetylene flame atomic absorption (AA) spectroscopy as per Standard Methods 3111B [31] with an AA spectrometer (PinAAcle 55, PerkinElmer, Waltham, MA, USA).

2.3. Experimental Setup

2.3.1. Batch regeneration study

This study consisted of several cycles of the sequential batch loading of the exchange media with EIMWW, separation of the media through vacuum filtration, batch regeneration of the media using different regenerants, separation of the media from the regenerant and finally reloading of the regenerated media with EIMWW. A schematic of the steps performed during the loading experiments is presented in Fig. 1.

For the loading, 0.5 g of dry IE media (natural or modified SIR-600 zeolite) were placed in 125 mL glass bottles which were filled with the EIMWW until the solution was overflowing, to avoid any head space and TA volatilization. Teflon tape was wrapped twice around the threaded portion of the 125 mL glass bottles prior to filling; after filling, they were sealed with a Teflon lined cap and electrical tape wrapped around the cap to prevent loosening. Once sealed, the bottles were placed in an end-over-end tumbler that rotated the bottles at 10 rpm to ensure gentle mixing and thorough solid-liquid contact, thus promoting IE. Preliminary batch experiments showed that equilibrium was reached in approximately 2 hrs; to be conservative and to simplify the logistics a 24 hour loading period was utilized. The bottles were then removed from the tumbler, and the content vacuum filtered through a glass microfiber filter (Whatman plc, Maidstone, Kent, England) to separate the solid and liquid phases. The filtrate was analyzed following the procedures cited above. The IE materials captured in the filtration were rinsed with distilled water and stored within
a desiccator until they were regenerated. A mass balance between the initial and final conditions of the loading step was used to determine the equilibrium solid phase concentrations (or loadings or capacity) of TA, potassium and calcium. The equation is as follows:

\[
q_{eq} = \frac{(C_i - C_{eq}) V}{M}
\]  

(1)

where \(q_{eq}\) is the equilibrium IE material capacity (meq ion/g IE media); \(C_{eq}\) and \(C_i\) are the equilibrium and initial pollutant liquid phase concentrations, respectively (meq/L); \(V\) is the volume of solution (L) within the bottles and \(M\) is the mass of IE material (g). All batch experiments were performed in triplicate at room temperature (approximately 23 °C).

In the regeneration step, illustrated by Fig. 2, the loaded IE media from each bottle was transferred to a new, clean 125 mL bottle; these bottles were then filled with the regeneration solution and sealed as described above. As desorption is a slower process, the regeneration contact time in the end-over-end tumbler was 48 hrs. Finally, the bottle contents were vacuum filtered to recover the IE media.

The five regeneration solutions evaluated were identified by their Na and K percentage by weight ratio. Only two contained potassium; a 5% KCl solution, and a solution of 5% KCl plus 5% NaCl. The sodium-based regeneration solutions consisted of 2.5%, 5 and 10% NaCl solutions. All results were referred to by the ion-exchanger regenerating salt (NaCl or KCl) and their percent concentration. The exception being the 5% NaCl + 5% KCl regenerations which were simply referred to as the NaCl + KCl. The pH of each batch regeneration solution was increased to approximately 10 by titration using 0.2 M NaOH or 0.2 M KOH solutions. Any pH adjustments for the NaCl + KCl solution were accomplished prior to combining respective K and Na regenerants. The purpose of the potassium centric regenerations was to assess the impact of other regeneration solutions on the TA removal. The preloading with K may help predict the long-term performance of IE systems in the treatment of EIMWW, as it is analogous to the continuous exposure to K. Once regenerated, the ion exchange media was reloaded following the same procedure as the original loading step. This study regenerated the ion exchange materials using NaCl solutions as this is the standard approach. This approach was selected because the study is primarily focused in determining the feasibility of the ammonia removal. It is recognized that to make the regeneration more sustainable the NaCl solutions should be recyclable, and there is a need for further investigation on this topic.

2.3.2. Column experiments

The second set of experiments investigated the competitive ion exchange in the practical column set-up, three column runs were performed. The experimental set-up is shown in Fig. 3, it included a 28.75 mm I.D., 150 mm long acrylic column. A total of 40 g of SIR-600 was placed within the column. The SIR-600 media was selected for the column experiments because it had a higher TA uptake capacity than clinoptilolite in the batch loading tests, as well as a lower Ca\(^{2+}\) uptake capacity. To prevent loss of material, glass wool was inserted into the bottom of the column, a perforated acrylic disk was placed overtop the glass wool, and the IE media was placed

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![Fig. 1. Schematic of the Batch Loading Experiments (a) Bottle preparation; b) 24 hr equilibration while mixing in tumbler; c) Separation by vacuum filtration; and d) Chemical analysis.](image-url)
above the disk. To distribute the flow equally and reduce channeling, a second perforated disk was placed above the IE material. The total bed depth of the zeolite was 71.0 mm, which resulted in an empty bed volume of 46.1 mL. Flow control was accomplished by placing a clamp on the rubber tubing used for the column effluent. The feed to the column was introduced into the column by a 2.0 L plastic reservoir, which helped ensure a constant head above the IE material. This was a gravity flow system, where the water level within the column was always above the media, ensuring an adequate contact time.

A flow rate of approximately 29 mL/min was maintained for the loading cycles and effluent was collected after various predetermined intervals for chemical analysis. The column was then regenerated using the regeneration solution that was identified as the best in the batch loading/regeneration tests; the regenerant flowrate was approximately 1 mL/min.

During the first experiment, the column was loaded with the EIMWW, run until exhaustion and subsequently regenerated. In the second run, the regenerated SIR-600 zeolite was again loaded with the EIMWW until a breakthrough concentration of 0.55 meq TA/L (10 mg/L as NH₄) was reached. The same media was regenerated and used to conduct a third column run with the 3.87 meq TA/L (70 mg/L as NH₄) synthetic wastewater as feed to quantify the impact of the competition; this column was once again run until a breakthrough concentration of 0.55 meq TA/L.

To compare the performance of the column with regard to potassium, calcium and TA, the respective capacities were calculated using a simplified numerical integration of the mass of the species removed [32]:

\[
q_{\text{column}} = \frac{1}{M} \sum Q (C_{\text{feed}} - C_{\text{eff}}) \Delta t
\]

where \(q_{\text{column}}\) is the capacity for a given species that has been achieved by the IE material within a column at a given time (meq contaminant/g ion exchange media); \(Q\) is the flow rate for a given time interval (L/min); \(M\) is the mass of IE material present in the column (g); \(C_{\text{feed}}\) and \(C_{\text{eff}}\) are the influent and effluent pollutant concentrations respectively (meq/L); and \(\Delta t\) is the given time step (min).

3. Results and Discussion

3.1. EIMWW Characteristics

To characterize EIMWW, it was analyzed for TA, K, Ca, and pH in-house and by ICP at the University of Ottawa’s Advanced Research Centre and confirmed at a commercial lab. The results are presented in Table 1. EIMWW had a pH of approximately 5.7 and the main cations were TA (3.87 meq/L), Ca (3.9 meq/L), K (2.85 meq/L), and sodium (11.8 meq/L). The main anions were sulfate and bicarbonate. So EIMWW had approximately the same concentration of Ca and TA in meq/L and they were 35% higher than the K concentration. Given the general selectivity order of clinoptilolite presented in the introduction, the ions that will impact TA removal the most will be K, and Ca.

3.2. Batch Loading and Regeneration Studies

The first phase of this study focused on the effects of repeated loading of natural and modified SIR-600
zeolites with real EIMWW. Subsequent regeneration cycles were performed with various regeneration fluids to establish the effects of regenerant concentration and composition, as well as to determine the long-term performance of both materials.

The impact of the multiple EIMWW loading and regeneration cycles on the average TA exchange capacities for SIR-600 and the clinoptilolite are shown in Figs. 4 and 5 respectively. The error bars represent one standard deviation. For SIR-600, the two KCl regenerations solutions (Fig. 4) show a decrease in TA exchange capacities of approximately 0.1 meq TA/g over three cycles, or approximately 25%. These reductions in TA capacity were likely caused by a higher affinity for potassium and by the greater quantities of K⁺ present after the KCl regeneration. This would lead to a higher portion of exchange sites occupied by K⁺. Due to lower TA uptake capacities for both IE materials, the regenerations using KCl based solutions did not seem promising and were abandoned after 3 cycles and only the NaCl based regenerants were tested for a 4th cycle.

Previous studies [18,24] suggest that a noticeable increase in exchange capacity is possible due to the conditioning of the ion-exchange as it undergoes regeneration cycles. This was the case for the 5% and 10% NaCl regenerated IE, but not for the 2.5% NaCl regeneration. The 2.5% NaCl regenerant did not contain enough sodium to increase the SIR-600 TA exchange capacities in later cycles, which remained between 0.34-0.38 meq TA/g over the four cycles. The 10% NaCl had the highest capacities ranging from 0.38-0.50 meq TA/g over the four cycles for SIR-600.

Fig. 5 presents the repeated averaged TA uptake capacities for the same regeneration solutions applied to the clinoptilolite. The clinoptilolite regeneration cycles featured similar trends as those presented for the SIR-600 but had greater variability. By comparing Figs. 4 and 5 it was immediately evident that the SIR-600 results were more consistent. Not only are the error bars for the SIR-600 smaller in most cases, but the clinoptilolite’s results are more variable. This was attributed to clinoptilolite’s natural origins that give it a lower product consistency than the manufactured SIR-600. As can be seen in Figs. 4 and 5, SIR-600 and clinoptilolite had lower TA capacities when regenerated with 2.5% NaCl solutions compared to 5 and 10% NaCl solutions.

The impact of the multiple EIMWW and regeneration cycles with 10% NaCl on the calcium exchange capacity is shown in Fig. 6. The first loading cycle yielded a negative Ca²⁺ exchange capacity for the clinoptilolite. This indicated that calcium was initially present as an exchangeable cation, and some Ca²⁺ leached during the first loading cycle. It has been reported in literature that clinoptilolite may initially contain several counter-ions, including calcium and potassium [7,21,33]. Furthermore, the clinoptilolite featured a larger affinity for calcium than the SIR-600, as the second through fourth loadings resulted in larger calcium uptake capacities than the SIR-600. The clinoptilolite had an equilibrium capacity of 0.16 ± 0.01 meq Ca²⁺/g. The SIR-600 featured a slight increase in capacity with loading cycles and stabilized at approximately 0.12 meq Ca²⁺/g. The lower SIR-600 affinity for Ca²⁺ is likely linked...
The higher Si/Al ratio in this study, may explain the which have ratios ranging from 2.7 to 5.7 [34]. Thus, 22.4, which is much larger than typical clinoptilolites. The clinoptilolite used in this study had a Si/Al ratio of one such material. According to the supplier[28], the K

\[ \text{Species} \times \text{Concentration (mg/L)} \times \text{Concentration (meq/L)} \]

\[ \begin{array}{c|c|c}
\text{Species} & \text{Concentration (mg/L)} & \text{Concentration (meq/L)} \\
\hline
\text{Cs} & \text{N.D.} & \text{N.D.} \\
\text{K} & 111.5 & 2.85 \\
\text{TA} & 70.0 & 3.87 \\
\text{Ca}^{2+} & 78.05 & 3.90 \\
\text{Fe} & 0.70 & 0.04 \\
\text{Al} & 0.08 & 0.01 \\
\text{Mg} & 12.67 & 1.04 \\
\text{Si} & 3.56 & 0.51 \\
\text{B} & 0.07 & 0.02 \\
\text{Ca}^{2+} & 78.05 & 3.90 \\
\text{Sr} & 2.10 & 0.05 \\
\text{Na} & 270.29 & 11.76 \\
\text{K} & 111.5 & 2.85 \\
\text{SO}_4 & 1050.0 & 21.86 \\
\text{HCO}_3 & 300.0 & 4.92 \\
\end{array} \\
\]

to this media’s somewhat higher TA exchange capacities shown in Fig. 4.

The potassium exchange capacities over four loading cycles with 10% NaCl as the regeneration solution are presented in Fig. 7. Both IE materials experienced increased K\(^+\) loadings following the first exchange cycle. The leaching of preloaded calcium is believed to be partially responsible [20,21] for the lower first K\(^+\) loadings. The increase in potassium capacities during the second and subsequent cycles could be attributed to the leaching of preloaded potassium. Prior to the first test, the zeolites were only rinsed with distilled water to remove any dirt or impurities, therefore the second factor which may have caused the increase of K\(^+\) uptake capacities was the conditioning of the zeolites. The clinoptilolite respectively leached 0.2, 0.24 and 0.32 meq K\(^+\) for the 2.5, 5 and 10% regenerants whereas the SIR-600 leached 0.26, 0.27 and 0.36 meq K\(^+\). Several studies have stated that repeated loading/regeneration cycles can increase the capacity of a material, by creating a uniformity of counter-ions [14,18,24].

The structure of zeolites plays an important role in defining the selectivity of materials. It has been stated that zeolitic structures containing high Si to Al ratio, in fact, have a higher selectivity towards lower charge density elements [34]. Clinoptilolite is one such material. According to the supplier[28], the clinoptilolite used in this study had a Si/Al ratio of 22.4, which is much larger than typical clinoptilolites which have ratios ranging from 2.7 to 5.7 [34]. Thus, the higher Si/Al ratio in this study, may explain the higher capacities for potassium. In contrast, the SIR-600 featured a smaller calcium capacity than the clinoptilolite and a higher potassium capacity relative to its calcium (a high charge density cation) capacity. Table 2 summarized the averaged calcium, potassium and TA capacities for the 4th exchange cycle for 5% and 10% NaCl regenerations. As the potassium based regenerants and the 2.5% NaCl regenerator resulted in lower TA capacities, they were omitted from Table 2. The main findings of Table 2 are as follows. First, increasing the concentration of regenerant from 5% to 10% did not result in a significant increase of the TA exchange capacity of both materials. For clinoptilolite the final capacities were 0.30 ± 0.02 meq TA/g and 0.34 ± 0.02 meq TA/g, while for SIR-600 they were 0.44 ± 0.03 meq TA/g and 0.46 ± 0.01 meq TA/g, respectively. Second, for the 10%NaCl regenerant the SIR-600 media had resulted 35% higher TA exchange capacities from EIMWW than clinoptilolite (0.46 ± 0.01 meq TA/g versus 0.34 ± 0.02 meq TA/g), thus it was selected for the subsequent column studies. Third, for both materials the average capacities indicated a preference for potassium over TA. The literature for the selectivity of zeolites such as clinoptilolite [26,33] tends to be in agreement, suggesting that there is a defined order of selectivity, with a slight preferential exchange of K\(^+\) over TA. Clinoptilolite's average potassium exchange capacity was higher than its TA exchange capacity for both the 5% NaCl and the 10% NaCl, 20% and 16.7% higher, respectively. However, for the 10% NaCl-clinoptilolite experiments the 95% confidence limits of the K\(^+\) and TA loadings overlap, therefore, the difference is not statistically significant. In contrast, for SIR-600 with 10% NaCl regeneration the 95% confidence limits of the K\(^+\) and TA uptake capacities do not overlap so they are significantly different; while for SIR-600 with 5% NaCl regeneration the K\(^+\) and TA loadings are not statistically different. Thus, for EIMWW the media tested have a slight preference of K\(^+\) over NH\(_4\). The pH of all regeneration solutions was raised to a value between 9.5 and 10, and monitored before and after contact with IE materials (i.e., the regeneration process). It was noted that after contact with the ion exchange material the pH decreased by a similar amount for both IE materials, approximately 1-1.5 pH units. All regeneration cycles featured similar pH reductions regardless of the regenerant used. This reduction in pH was
attributed to the elution of $\text{NH}_4^+$ during the regeneration step which caused the shift in the TA concentration from ammonium ($\text{NH}_4^+$) to ammonia ($\text{NH}_3$) due to the elevated pH. The shift is described by the equation

$$\text{NH}_4^+ \leftrightarrow \text{NH}_3 + \text{H}^+$$  \hspace{1cm} (3)

The shift results in the generation of $\text{H}^+$ and a decrease in the pH. Similar results have been reported elsewhere [5].

3.3. Column Studies

Full-scale IE applications generally consist of continuous flow column systems for practical
reasons as well as to maximize operational efficiency. Therefore, it was important to determine if the EIMWW could be treated in a column system, particularly because in these systems the contact times are only in the order of minutes and the kinetics of the exchange may impact the selectivity. A second goal of this column study was to quantify the reduction in TA capacity of a continuous IE column resulting from the presence of multiple competing species by comparing the performance when treating the EIMWW versus a synthetic TA solution. Based on the higher TA capacities and lower Ca$^{2+}$ uptakes achieved from the batch regeneration studies, the SIR-600 modified zeolite was selected for the column studies.

Fig. 8 illustrates the first column run's effluent concentrations of TA versus the volume of water treated in terms of bed volumes (BV) using a flow rate of approximately $29 \pm 2$ mL/min. This figure shows that Ca$^{2+}$ broke through almost immediately and at approximately 30 BV the media became saturated with Ca$^{2+}$ (i.e., $C_{\text{eff}}/C_{\text{feed}} = \sim 1$ meq/L). The effluent TA and K$^+$ concentrations were almost negligible up to that point, they increased significantly between 30 and 35 BV and gradually thereafter. During this latter part of the run there was no Ca$^{2+}$ removal (i.e., $C_{\text{eff}} = C_{\text{feed}} = 4.17$ meq/L), the TA concentration increased gradually and the K$^+$ concentrations were consistently lower than those of TA. This reinforces the SIR-600 batch test results in Table 2, which demonstrated a preference for potassium and TA over calcium. The column run was stopped after treating 91 BV of EIMWW, as the column was approaching saturation with respect to TA ($C_{\text{eff}}/C_{\text{feed}} \sim 0.85$). At that time, the effluent potassium concentration was approximately 50% of the feed concentration, suggesting a higher ultimate potassium capacity. It is believed that this higher potassium capacity caused the TA exchange zone to move ahead of the potassium exchange front, resulting for a time in primarily single component TA ion-exchange in the portion of the column ahead of the potassium exchange front. It would seem that the potassium front begun to displace previously sorbed TA ions at 30 BV, where the TA effluent concentration became significantly higher than that of K$^+$.

The resulting exchange capacities for calcium, potassium and TA (Fig. 9) were calculated using the Fig. 8 data and equation (2). This figure shows that SIR-600 had a slight preference for TA over potassium, up until approximately 90 BV of EIMWW treated. This preference for TA may be due to the lower K$^+$ feed concentration (2.75 versus 3.75 meq/L). The TA and K$^+$ capacities are consistent with the batch test results presented in Table 2, which reported similar capacities for potassium and TA. The horizontal dotted/dashed lines in Fig. 9 represents the K$^+$, TA and Ca$^{2+}$ exchange capacities obtained for batch studies treating the same source of EIMWW. This dotted line (TA batch) helps illustrate that the TA uptake of the continuous flow column system is approximately 48% smaller than of the batch set-up (0.24 meq TA/g vs 0.46 meq TA/g). It is possible that this was due to a drastic reduction in contact times; the batch tests had
a contact time of 24 hrs, whereas during lab-scale column test the contact time was 1.6 minutes. The differences between the two types of tests are more likely explained by the nature of the tests. During the batch tests there was a limited mass of the stronger sorbing species (i.e., $K^+$) which limited the extent of the reduction in the exchange capacity exhibited for TA, while in the column system, the feed continuously introduced additional mass of the $K^+$ and resulted in even greater competitive exchange and a reduction in the TA capacity. The top dashed grey line (Batch K) shows that the $K^+$ loadings achieved in the column tests were significantly lower than those achieved in the batch tests. If the column would have been operated longer, the $K^+$ loadings would have increased further possibly reaching the levels observed during the batch tests (i.e., the bottom dashed line in Fig. 9). The column was regenerated several times with a 10% NaCl solution. The column experiment was repeated with EIMWW, and for this second column run an effluent breakthrough concentration criterion of 0.55 meq TA/L (10 mg TA/L) was used. This breakthrough concentration was selected as it is a potential effluent discharge criterion; the actual criteria is likely to depend on the degree of dilution upon discharge into a receiving water. This column run was followed by another regeneration and a third column run using a single solute TA.

Table 2. Average of the 4th Exchange Cycle Uptake Capacities of Clinoptilolite and SIR-600 for 5 and 10% NaCl Regeneration (± one SD).

| Regenerant | 5% NaCl | 10% NaCl |
|------------|---------|----------|
| Clinoptilolite |
| $Ca^{2+}$ Uptake Capacity (meq $Ca^{2+}$/g) | 0.14 ± 0.02 | 0.16 ± 0.01 |
| $K^+$ Uptake Capacity (meq $K^+$/g) | 0.36 ± 0.01 | 0.39 ± 0.06 |
| TA Uptake Capacity (meq TA/g) | 0.30 ± 0.02 | 0.34 ± 0.02 |
| SIR-600 |
| $Ca^{2+}$ Uptake Capacity (meq $Ca^{2+}$/g) | 0.13 ± 0.06 | 0.12 ± 0.01 |
| $K^+$ Uptake Capacity (meq $K^+$/g) | 0.46 ± 0.04 | 0.52 ± 0.01 |
| TA Uptake Capacity (meq TA/g) | 0.44 ± 0.03 | 0.46 ± 0.01 |

Fig. 7. Successive Potassium Uptake Capacities in the Batch Tests with 10% NaCl Regeneration.

Fig. 8. Breakthrough Curves for $Ca^{2+}$, $K^+$ and TA in the Treatment of EIMWW/SIR-600 Column Run.
synthetic wastewater solution. The results of the second and third column runs are presented in Fig. 10 where the effluent TA concentration versus treated bed volumes for both wastewaters are compared, thus allowing to determine the extent to which competing ions affect the exchange process. The presence of competing ions resulted in a much earlier TA breakthrough than for the synthetic single component wastewater (50 BV versus 274 BV). When treating the EIMWW using the above breakthrough level, the column’s IE material was only able to treat 18% of the volume of wastewater that it would treat when the feed was a single solute TA solution. These results clearly confirm that the feasibility of TA ion exchange applications need to be assessed with the wastewater in question and not a single solute TA solution. Furthermore, during the second column run the 0.55 meq TA/L breakthrough level occurred at ~50 BV, which is later than in the initial column run (~20 BV in Fig. 8). The smaller quantity of BVs treated prior to breakthrough is attributed to the lack of conditioning prior to the initial column run.

Using the data in Fig. 10 and equation (2), TA uptake capacities were obtained for both the single component synthetic wastewater and the second EIMWW run during the 0.55 meq TA/L breakthrough experiments (Fig. 11). When the TA capacities were considered during the first 50 BV, the synthetic wastewater’s exchange capacities were nearly identical to the EIMWW TA capacities. This implies that the impact of competition had not yet taken effect. It is believed that exchange capacity for potassium was somewhat higher than that of TA, resulting in the potassium ion exchange front within
the IE bed to trail behind the TA ion exchange front. This could be in part due to the lower influent K$^+$ concentration (~26%) which resulted slower relative loading rates.

In the treatment of the synthetic TA wastewater, the SIR-600 column treated 274 BV of wastewater (Fig. 10) before reaching the breakthrough concentration of 0.55 meq TA/L, so it achieved a TA uptake of 1.19 meq TA/g. At the same breakthrough level, the column treating EIMWW only achieved 20% of this value (0.24 meq TA/g). The potassium and calcium capacities were respectively 0.21 meq K$^+$/g and 0.17 meq Ca$^{2+}$/g for the second loading run. Thus, the column’s K$^+$ uptake capacity at the breakthrough (i.e., non-saturation) was 23% higher than that of TA, while for the batch tests for the same sorbent-regenerant combination, the K$^+$ capacity was only 11.3% higher. This indicates that the dynamic conditions in the column further favour the exchange of K$^+$ over TA. At the TA breakthrough in the second column run, the summation of the loadings of the three species was a total of 0.62 meq/g, approximately half of the total measured capacity for the synthetic wastewater (i.e., 1.19 meq/g). The higher value for the synthetic wastewater was likely due to the greater than five-fold longer column run which permitted greater degree of ion exchange. Finally, the effluent pH was 6.14 so the 10 mg TA/L breakthrough concentration corresponds to an unionized ammonia of 0.007 mg N/L, which is lower than the Canadian NH$_3$ industrial effluent guideline [5].

It is also of note that for the breakthrough level of the second column run, the TA uptake by the SIR-600 was approximately 0.23 meq TA/g, while the batch test loadings of SIR-600 regenerated with 5% KCl achieved an uptake of 0.27 meq TA/g. The similar capacities probably arise from the fact that the KCl regenerated media contained substantial quantities of the main competing cation (K$^+$) and was also exposed to a large mass of K$^+$. Thus, using K$^+$ preloaded media may be useful as a simple estimator of EIMWW column TA removals. The similarities may be coincidental, so this needs to be verified.

4. Conclusion

Two zeolites, a natural clinoptilolite and a modified clinoptilolite (SIR-600) were evaluated for their regenerative abilities and long-term performance when removing TA from an explosive impacted mining wastewater. The batch exchange/regeneration/ion-exchange experiments showed that regenerating both zeolites using KCl solutions resulted in smaller TA uptake capacities from EIMWW compared to the NaCl solutions. The regeneration with the 5% and the 10% NaCl solution resulted in the highest TA batch test uptakes, and the modified clinoptilolite had higher TA uptake from EIMWW than the clinoptilolite.

The removal of TA was greatly impacted by the presence of K$^+$ and Ca$^{2+}$ as competing ions in EIMWW. The batch loading tests showed that both media had a slight preference for K$^+$ over TA. In the EIMWW column tests, TA broke through before potassium confirming that the media had a higher selectivity for potassium. The most dramatic demonstration of the competitive interactions by the other ions in EIMWW was that the same column was able to treat five times more volume of a single solute ammonia solution than EIMWW for the same 0.55 meq TA/L breakthrough level. It is noteworthy that for the initial period of SIR-600 column treatment had nearly identical TA exchange capacities for both the EIMWW and the synthetic solution, which suggests that competition from various ions, including potassium had not yet taken effect and would only become prominent if the column was operated for a longer duration. This indicated that potassium exchange capacity is higher than TA capacity, resulting in a slower K$^+$ adsorption front and the increase in TA concentration around breakthrough was caused by the K$^+$ adsorption front catching up.

Because the regeneration with a 5% KCl solution incorporates many K$^+$ ions into the media, the batch test loadings using KCl regenerated media may be a simple and reasonable estimator of the column TA capacities. This requires verification.

The current study was performed using an explosives impacted mining wastewater secured from a mine, and the ion exchange material performed well in our limited length studies. Longer term testing needs to be performed using different EIMWWs which may include other contaminants that may impact the life of the ion exchange material. It is envisioned that the ion exchange material may have to be replaced every year or two. Based on the current results the EIMWW treatment system is expected to consist of two ion exchange columns, one that is in operation while the other is off-line being regenerated with a NaCl solution or waiting to go back on-line. It is expected that ion exchange cycles will last at least six hours while the regeneration will be completed in about two hours. Alternative schemes to treat the regeneration solution to
permit continuous reuse need to explored, however in the current study is concerned primarily on the TA removal.

Ethical statement

Authors state that the research was conducted according to ethical standards.

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Conflict of interest

None declared.

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