Preparation and characterization of gum karaya hydrogel nanocomposite flocculant for metal ions removal from mine effluents

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Abstract This research paper reports the removal of heavy metal ions from mine effluents using the gum karaya (GK)-grafted poly(acrylamide-co-acrylic acid) incorporated iron oxide magnetic nanoparticles (Fe$_3$O$_4$ MNPs) hydrogel nanocomposite [i.e., GK-cl-P(AAm-co-AA)/Fe$_3$O$_4$ hydrogel nanocomposite] and inorganic coagulants such as polyferric chloride (af-PFCl), Al$_2$(SO$_4$)$_3$, FeCl$_3$ and Mg(OH)$_2$. The Fe$_3$O$_4$ MNPs were incorporated in the matrix of the hydrogel polymer of Gk-cl-P(AAm-co-AA) through the free radical graft co-polymerization technique using $N,N'$-methylene-bis-acrylamide as the cross-linking agent. The graft co-polymerization of the P(AAm-co-AA) with Gk and the successful incorporation of the Fe$_3$O$_4$ MNPs within the hydrogel polymer matrix was evidenced using different characterization techniques such as FTIR, XRD, SEM and TEM. The performance of coagulants was evaluated by considering parameters such as turbidity removal, pH correction, metal removal and settling time. It was observed that the monomeric inorganic coagulants had a relatively poor performance compared to the organic coagulant, i.e., GK-cl-P(AAm-co-AA)/Fe$_3$O$_4$ hydrogel nanocomposite. Most of the coagulants achieved maximum turbidity removal in the range of 67–99.5 %, but the hydrogel nanocomposite showed the greatest reactivity by achieving the fastest floc formation rate and shortest optimum sedimentation time of 5 min (100 % removal in 5 min). The removal of metal followed the order Pb$^{2+} >$ Cr$^{6+} >$ Ni$^{2+}$ with an optimum settling time of 15 min; more often, Ni$^{2+}$ was poorly removed (≤23.2 % removal after 15 min) from acidic mine water samples. Therefore, the synthesized hydrogel nanocomposite has shown great potential as a flocculant and adsorbent for the removal of suspended particles as well as heavy metal ions and can be used to improve the quality of mine effluents prior to discharge in the environment.

Keywords Suspended particles · Heavy metal ions · Mine water · Inorganic and organic polymers · Flocculation

Introduction

A host of techniques making use of inorganic coagulants are often applied for the removal of suspended solids from mine water and control of the pH; however, problems related to the large amount of coagulants needed, excessive accumulation of sludge, the instability and breakage of aggregates, as well as the need of water conditioning have motivated investigation of alternative flocculants or coagulants (Rodriguez et al. 2003; Brostow et al. 2009; Yang et al. 2013). The efficiency of the flocculation process depends on the ability of coagulants and/or flocculants to adsorb on the fine residue particles bringing them together to form larger flocs which are likely to settle at a faster rate. Depending of their source, suspended particles may have different structures or properties and the selection of suitable coagulant will depend on the level of understanding of interactions between factors such as composition charge,
particle size, shape and density. Suspended particles found in mine ponds often derived from streams of mill rejects or flotation tailings in the form of slurry and are likely to contain free metal ions which could contaminate the water system and affect aquatic life as well as pose a risk to human health. Therefore, water authorities around the world have tightened regulations regarding disposal of effluent loaded with metal ions. The removal of heavy metal ions from solution is conventionally achieved using chemical and physical methods which have a number of limitations including sludge formation and relatively high cost (Kefala et al. 1999; Ahalya et al. 2003; Fosso-Kankeu et al. 2009).

The inorganic coagulants commonly used are aluminum and iron salts. In comparison with monomeric form of salts, pre-polymerized inorganic coagulants have shown distinctive advantages including wide range of pH applications, relatively lower doses required for effective treatment, lower sensitivity to low water temperature and lower residual metal ion concentrations (Jiang and Graham 1998). Pre-polymerized inorganic coagulants of aluminum [polyaluminum chloride (PACl)] and ferric [polyferric sulfate (PFS) and polyferric chloride (PFCl)] ions have been intensively used recently for wastewater treatment (Dewolfe et al. 2003; Zhao et al. 2011). Coagulants based on the macromolecular compounds have a lot of advantages over the inorganic coagulants because their molar mass, chemical structure, charge density and the nature of their functional groups can be easily modified (Mittal et al. 2015a, b). The polymer-based coagulants can be further divided into the synthetic and natural polymer-based coagulants. Synthetic polymer-based coagulants have better property profile as compared to natural polymer-based coagulants, but at the same time, they are also associated with some serious drawbacks like toxicity and non-biodegradability (Singh et al. 2000; Gaudreault et al. 2009). Moreover, it is very difficult to dispose the synthetic polymer-based coagulants after use so they cause a huge environmental burden. Therefore, in order to overcome this environmental problem, the natural biopolymer-based biodegradable coagulants/flocculants have been used by a number of researchers all over the world (Ghimici and Nichifor 2010, 2012; Ali 2010, 2012; Ghimici et al. 2012; Mittal et al. 2014a, b). Polysaccharide-based flocculants are environment-friendly and can be easily modified according to the end-use application. Recently, a great deal of effort has been made toward the development of flocculants with the advantages of both natural and synthetic polymers (Wan et al. 2007; Rani et al. 2010; Ho et al. 2010; Ghimici and Nichifor 2010, 2012; Ghimici et al. 2012; Mittal et al. 2013a). Different polysaccharides like pullulan-, dextran-, gum ghatti-, gum karaya- and gum xanthan-based polymeric materials and different nanomaterials have shown excellent potential as coagulants for the removal of different suspended particles from polluted waters. Furthermore, they have been also used as adsorbents for the removal of different metal ions and dyes from the aqueous solutions (Saleh et al. 2011; Gupta et al. 2011; Ghimici et al. 2012; Mittal et al. 2013a, b; Ghorai et al. 2014; Saravanan et al. 2013, 2014, 2015). Nanomaterials have also shown their potential as photo-catalysts (Saleh and Gupta 2011, 2012; Gupta et al. 2012; Saleh 2015). In our previous studies, the biodegradable graft co-polymers of gum ghatti showed good potential as flocculants and adsorbents for the adsorption of metal ions from the aqueous solutions (Mittal et al. 2013a, b, 2014a, b, 2015a, b).

Gum karaya (GK) is an anionic polysaccharide obtained from trees such as Sterculia urens, Sterculia villosa and Sterculia setigera growing in the Indian subcontinent and the North African countries, respectively (Weiping 2000). It is a highly branched complex polysaccharide consisting of D-galacturonic acid, D-galactose, L-rhamnose and D-glucuronic acid. The carbohydrate structure has a rhamnogalacturonan main chain consisting of α-(1→4)-linked D-galacturonic acid and α-(1→2)-linked-L-rhamnosyl residues. The side chain is made up of (1→3)-linked β-D-glucuronic acid or (1→2)-linked β-D-galactose on the galacturonic acid unit where half of the rhamnose is substituted by (1→4)-linked β-D-galactose (Anderson et al. 1982; Anderson and Wang 1994).

Therefore, keeping in view the structural diversity of the gum polysaccharides and their potential in water treatment applications, this study investigates the flocculation potential and adsorption behavior of GK-cl-P(AAm-co-AA)/Fe3O4 hydrogel nanocomposite for the simultaneous removal of suspended particles and heavy metal ions from mine water in comparison with the performance of monomeric and pre-polymerized (af-PFCl) inorganic coagulants.

Materials and methods

Methodology

Sample collection

The sampling points were located at the underground and surface ponds within the premises of the mine. The water samples were carefully collected at the subsurface in polyethylene bottles, 1 L each of capacity; parameters such as pH, temperature, electrical conductivity and turbidity were measured at the sites using portable Lovibond SensoDirect 150 multi-parameter water quality meter. The instruments were calibrated using original buffer solutions.
prior to the use in the field. The water samples were transported in cooler boxes containing ice packs and stored in the fridge overnight prior to the experiments. The mine water characteristics are reported in Table 1.

### Preparation of coagulants

#### Inorganic coagulants

The chemicals used were of analytical grade; the stock solutions of Al\(_2\)(SO\(_4\))\(_3\), FeCl\(_3\) and Mg(OH)\(_2\) were prepared in order to obtain 1 g/L of Al, Fe and Mg, respectively; this was done by diluting the salts of these metals into the demineralized water. The inorganic polymeric af-PFCl coagulant was synthesized by slowly adding Ca(OH)\(_2\) solution of 0.5 mol/L to a 0.1 mol/L solution of FeCl\(_3\) under constant stirring speed at 350 rpm at room temperature. A specific volume of Ca(OH)\(_2\) was added to FeCl\(_3\) solution till the predetermined basicity (\(B = [\text{OH}^-]/[\text{Fe}^{3+}]\)) was reached.

#### Organic coagulant

To synthesize the organic coagulant, i.e., GK-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite, cross-linked network of poly(AAm-co-AA) with GK was prepared through free radical graft co-polymerization technique using KPS and ABC redox pair as an initiator and MBA as a cross-linking agent. The details of the synthesis of the Gk-cl-P(AAm-co-AA) hydrogel polymer are given in our previous publication (Mittal et al. 2015a, b). Briefly, prior to grafting, a varied amount of the Fe\(_3\)O\(_4\) MNPs (25–100 mg) were added in 20 mL deionized water and sonicated for 4 h for a better dispersion of nanoparticles. After the sonication, GK (1 g) was added in the reaction mixture and stirred vigorously for the complete dissolution of gum in water. At this stage, a mixture of 30 mg KPS and 20 mg ABC was added in the reaction mixture and the temperature was maintained at 60 °C. Fifty milligrams of MBA was added in the reaction mixture followed by the addition of the mixture of AAm (1 g) and AA (2.5 mL) with constant stirring. After 3 h, the reaction was stopped and cooled at room temperature. Homopolymers were removed by successive soxhlet extraction using acetone as solvent. Finally, the product was washed with 250 mL of acetone and dried in hot air oven at 40 °C till a constant weight was achieved. The selection of the best hydrogel nanocomposite for the rest of the studies was done on the basis of the maximum percentage swelling (\(P_s\)). Initially, 100 mg of each hydrogel nanocomposite was added to 100 mL deionized water. After 24 h, the samples were removed from the water, wiped gently to remove the surface water and weighed. The \(P_s\) value was calculated using the following equation:

\[
P_s = \frac{W_f - W_i}{W_f} \times 100
\]

where \(W_f\) and \(W_i\) are the weights of the swollen and dry hydrogel nanocomposites, respectively.

It was observed that the hydrogel nanocomposite with 50 mg concentration of Fe\(_3\)O\(_4\) MNPs gave the maximum swelling, so it was used for the rest of the studies (Table 2).

#### Characterization of the organic coagulant

The incorporation of Fe\(_3\)O\(_4\) MNPs in the cross-linked network of GK-cl-P(AAm-co-AA) hydrogel was confirmed using different techniques such as BET, FTIR, SEM, TEM and XRD. X-ray diffraction study of the samples was done by using Rigaku Ultima IV X-ray diffractometer employing Cu Kx radiation of the wavelength of 1.5406 Å with visible lights at 45 kV/40 mA. The structural determination and the confirmation of grafting of P(AAm-co-AA) onto GK and incorporation of Fe\(_3\)O\(_4\) MNPs within the polymer matrix were done through FTIR (PerkinElmer Spectrum 100 spectrometer) using KBR pellet method in the spectral range of 4000–400 cm\(^{-1}\) with a resolution of 4 cm\(^{-1}\). The changes in the morphology of GK after grafting and the polymer matrix after the incorporation of Fe\(_3\)O\(_4\) MNPs were studied using scanning electron microscope (TESCAN, VEGA SEM) under a 20-kV electron acceleration voltage coupled with energy-dispersive (EDX) for elemental analysis. As the samples were non-conducting in nature, these were coated with carbon for conductive

### Table 1

Recorded values of the physicochemical quality of water samples

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Turbidity (NTU)</th>
<th>pH</th>
<th>EC (µs/cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>56.31</td>
<td>2.24</td>
<td>4.6</td>
</tr>
<tr>
<td>2A</td>
<td>54.26</td>
<td>7.75</td>
<td>2</td>
</tr>
<tr>
<td>3A</td>
<td>2209.41</td>
<td>7.37</td>
<td>1.9</td>
</tr>
<tr>
<td>4A</td>
<td>333.07</td>
<td>7.69</td>
<td>1.8</td>
</tr>
<tr>
<td>5A</td>
<td>150.27</td>
<td>2.71</td>
<td>2.3</td>
</tr>
</tbody>
</table>

### Table 2

Effect of the Fe\(_3\)O\(_4\) MNPs concentration on the \(P_s\)

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Amount of Fe(_3)O(_4) MNPs (mg)</th>
<th>(P_s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>25</td>
<td>1625</td>
</tr>
<tr>
<td>2</td>
<td>50</td>
<td>2049</td>
</tr>
<tr>
<td>3</td>
<td>75</td>
<td>1768</td>
</tr>
<tr>
<td>4</td>
<td>100</td>
<td>1426</td>
</tr>
</tbody>
</table>
metal coating and the scanning was concurred with the position of beam on the specimen for maintaining small size over a large distance relative to the specimen. Successful incorporation and variation in the size of Fe₃O₄ MNPs within polymer matrix was examined through JEOL JEM-2100F, field emission electron microscope with an accelerating voltage of 90 kV. Small pinch of sample was added in few milliliters of ethanol and sonicated for 30 min. Two drops of sonicated suspension was transferred to the TEM grid and allowed to dry in ambient air. The increase in the BET surface area and porosity of the polymer matrix after the incorporation of Fe₃O₄ MNPs was studied by nitrogen adsorption–desorption measurements using Micromeritics, ASAP 2020, surface area and porosity analyzer.

**Jar test procedures**

The coagulation experiment was conducted using a jar test apparatus equipped with flat paddle impellers and was based on the conventional jar tests method with two mixing stages: a rapid mixing at 200 rpm for 1 min 30 s after dosing followed by a slow mixing at 50 rpm for 15 min. The jar test was carried out with beakers of 500 mL capacity, and each beaker was filled with 300 mL sample. A fixed dosage considered in this study was 50 mg/L of the inorganic coagulants and 100 mg/L of the organic polymer. After the slow mixing, the samples were left to settle for a given period of time. After sedimentation, the sample was collected at 1 cm below the surface for water quality analysis.

**Settling time**

The settling time was an important variable considered to evaluate the rate of settling of flocs formed during the reaction of inorganic and organic coagulants with suspended particles. Samples were carefully withdrawn at 1 cm below the supernatant surface after 5, 10, 15 and 60 min of sedimentation.

**Analysis**

The residual metal ions in the supernatant were quantified using the inductively coupled plasma optical emission spectrometer (ICP Expert II, Agilent Technologies 720 ICP-OES). To detect significant difference between metal removal rate at different settling times, Mann–Whitney test was performed and the difference was considered significant for values of $P < 0.05$.

**Results and discussion**

**Characterization of the organic coagulant**

**FTIR**

The successful graft co-polymerization of P(AAm-co-AA) onto Gk and then the cross-linkage of the co-polymer [Gk-cl-P(AAm-co-AA)] to the surface of Fe₃O₄ MNPs were evidenced using FTIR spectroscopy. FTIR spectrum of Gk, Gk-cl-P(AAm-co-AA) hydrogel polymer, uncoated Fe₃O₄ MNPs and Gk-cl-P(AAm-co-AA)/Fe₃O₄ hydrogel nanocomposite is shown in Fig. 1. The FTIR spectrum of Gk reveals absorption peaks at 3332 cm⁻¹ assigned to the –OH stretching of the carbohydrates; 2938 cm⁻¹, attributed to the C–H stretching modes of CH₂ groups; and 1715 cm⁻¹ attributed to the C=O stretching of amides.
and 1605 cm\(^{-1}\), due to the C=O stretching vibrations of the free carboxylic acids and the methylated galacturonic acid esters of the gum. The absorption peak observed at 1148 cm\(^{-1}\) is attributed to the pyranose rings of the polysaccharides, and the peak at 1414 cm\(^{-1}\) is due to the \(-\text{COO}^-\) and the –C–C stretching of \(-\text{CH}_2\) scissoring (Fig. 1).

In the FTIR spectrum of the Gk-cl-P(AAm-co-AA) hydrogel polymer, the following peaks and assignments were observed: 3445 cm\(^{-1}\), due to –OH stretching vibrations; 2944 and 2641 cm\(^{-1}\), due to aliphatic –CH\(_2\) stretching vibrations; 1725 cm\(^{-1}\), due to the C=O stretching in AA; 1457 cm\(^{-1}\), due to the amide –NH in-plane bending; 1246 cm\(^{-1}\), due to the amide –CN stretching vibration; and 893 cm\(^{-1}\), due to the amide –OCN deformation (Fig. 1). In the FTIR spectrum of uncoated Fe\(_3\)O\(_4\) MNPs, characteristic adsorption peak of Fe–O bond was observed at 555 cm\(^{-1}\), whereas after graft co-polymerization in Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite, this particular peak was shifted at 585 cm\(^{-1}\). FTIR spectrum of Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite showed peaks at 3417.87 cm\(^{-1}\) (–OH stretching vibration of carbohydrate), 2928 cm\(^{-1}\) (–OH stretching of acrylic acid), 1723 cm\(^{-1}\) (C=O stretching of acrylic acid), 1431 cm\(^{-1}\) (–CH, –CH\(_2\) and –OH in-plane bending in carbohydrates) and 1071.05 cm\(^{-1}\) (–CO stretching regions as complex bands resulting from C–O and C–O–C stretching vibrations).

**XRD**

The XRD patterns of uncoated Fe\(_3\)O\(_4\) MNPs and Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite are shown in Fig. 2. The XRD pattern of uncoated Fe\(_3\)O\(_4\) MNPs (JCPDS 75-0033) showed the standard pattern with the diffraction peaks at (220), (311), (400), (422), (511) and (440) at 2\(\theta\) scale. The XRD patterns of Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite showed all the peaks observed in case of uncoated Fe\(_3\)O\(_4\) MNPs which confirm the incorporation and uniform distribution of Fe\(_3\)O\(_4\) MNPs within Gk-cl-P(AAm-co-AA) polymer matrix with a little bit of reduced intensity. The peaks became slightly wider because of the coating of non-magnetic and amorphous Gk-cl-P(AAm-co-AA) hydrogel polymeric shell (Mittal et al. 2014a, b).

**Scanning electron microscopic studies**

The SEM images of Gk, Gk-cl-P(AAm-co-AA) and Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite are shown in Fig. 3. Very smooth, granular and homogeneous surface morphology was observed in the SEM image of Gk (Fig. 3a). Whereas this smooth surface became heterogeneous and very irregular and rough surface morphology was observed in case of the SEM of Gk-cl-P(AAm-co-AA) hydrogel, this might be due to the cross-linking between different polymeric chains (Fig. 3b). The surface morphology of the polymeric matrix was also found to change after the incorporation of Fe\(_3\)O\(_4\) MNPs, and a better matrix coherence was achieved (Fig. 3c). The surface morphology of Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite was more coherent and near co-continuous than the fibrillar surface of Gk-cl-P(AAm-co-AA). The EDX spectra of the Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite also showed peaks of Fe which further confirmed the incorporation of Fe\(_3\)O\(_4\) MNPs within the cross-linked polymer matrix, i.e., Gk-cl-P(AAm-co-AA) (Fig. 3d).

**Transmission electron microscopy**

Figure 4 shows the typical TEM images of uncoated Fe\(_3\)O\(_4\) MNPs and Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite. TEM image of uncoated Fe\(_3\)O\(_4\) MNPs showed well-shaped ellipsoidal and spherical magnetic nanoparticles (Fig. 4a). TEM of Gk-cl-P(AAm-co-AA)/Fe\(_3\)O\(_4\) hydrogel nanocomposite (Fig. 4b) also showed the presence of ellipsoidal and spherical magnetic nanoparticles which confirmed the incorporation of Fe\(_3\)O\(_4\) MNPs in the three-dimensional cross-linked structure of Gk-cl-P(AAm-co-AA). The mean diameter of coated nanoparticles (60 nm) in hydrogel nanocomposite was found to be higher than that of uncoated Fe\(_3\)O\(_4\) MNPs (35 nm).
Brunauer–Emmett–Teller (BET) surface area

The effect of the incorporation of Fe$_3$O$_4$ MNPs within the cross-linked polymer matrix of Gx-cl-PAA on its surface and pore volume was studied using BET analysis. BET surface area, pore volume and pore diameter of Gk-cl-P(AAm-co-AA) hydrogel polymer were found to be 0.4658 m$^2$/g, 0.002458 cm$^3$/g and 183.6915 Å,
respectively, and the values of these parameters were increased to 0.9845 m²/g, 0.008473 cm³/g and 216.7254 Å, respectively, after the incorporation of Fe₃O₄ MNPs in the polymer matrix.

**Characteristics of water samples**

The pH of mine water samples ranged from acid to alkaline (from 2.24 to 7.75) as shown in Table 1; the ground water samples (1A and 5A) were mostly acidic, while the surface water samples (2A, 3A and 4A) were mostly alkaline. The surface water samples were more turbid than the underground water, the turbidity ranging from 54.26 to 2209.41 NTU. The electrical conductivity values of the ground water samples were relatively higher than those of the surface water samples.

**Flocculation potential of coagulants**

A series of jar tests were conducted using five coagulants to reduce the turbidity in five mine water samples of different characteristics (Table 1). Initial experiments were conducted to estimate the optimum coagulant dosage; the fixed coagulant dosage was added to mine water solutions without amendment of pH. The result in Fig. 5 shows unequal performance of coagulants, the turbidity removal ranging from 0 to 99.43 %. In general, alum (Al₂(SO₄)₃) and FeCl₃ showed a very poor turbidity removal capacity. The inorganic and the organic [GK-cl-P(AAm-co-AA)/ Fe₃O₄] polymers achieved higher turbidity removal, but the organic polymer was most effective in acidic mine water, while the inorganic polymer performed better in alkaline mine water. The monomeric coagulants were generally very ineffective in the removal of turbidity from acidic mine water; this dependence on pH was previously reported by other researchers (Jiang and Graham 1998; Gao et al. 2011). Co-polymers of acrylamide and sodium acrylate are reported to be non-anionic and therefore are likely to show the greatest activity at pH of 4 and below (Connelly and Richardson 1984).

An advantage of the treatment of water with the inorganic polymer (af-PFCl) is that the branches of the polymer are adsorbed onto different colloidal particles, interlocking with one another and thereby leading to the formation of a settleable mass which improves turbidity removal (Peavy et al. 1985). The pH range in the batch samples falls within a range 2.35–8.43, which is suitable for the development of various hydrolysis species. Each species has an area of dominance with some overlap as explained by Peavy et al. (1985). Fe(OH)⁺² is the first species to form due to decreasing concentration of Fe³⁺ covering an area in the pH range 1–5. It is followed by Fe(OH)⁻² which falls within pH range 2–7 and follows Fe(OH)₃(aq) which covers pH of 4–8. The last species is Fe(OH)₄⁻ which covers pH range 4–14. According to turbidity removal and corresponding pH values in the raw samples 1A and 5A, it is evident that adsorption occurs through charge neutralization. Charge neutralization occurs when the pH is acidic where excess positive charges neutralize negative charges in colloidal particles. Alkaline batch samples are conducive for the existence of ferric hydroxide species as they have a minimum solubility in a pH range of 7–10, which gives them more flexibility during hydrolysis (Binnie et al. 2003).
Correction of pH by coagulants

The treatment of waste water especially acid mine drainage (AMD) focuses among other on the correction of pH; hence, in most cases two or more coagulants are used to address turbidity removal and pH correction; it will be ideal to have a coagulant that can do both in a single step saving, therefore, time and money. The divergence of pH

Fig. 7 Effect of aging on the turbidity removal efficiency of (a) (GK-cl-P(AAm-co-AA)/Fe₃O₄), (b) FeCl₃, c Al₂(SO₄)₃, d Mg(OH)₂, e af-PFCl
in our mine water samples offers a good opportunity to investigate the potential of our coagulants to adjust the pH to acceptable level. The results in Fig. 6 show that our coagulants had little or no effect on pH in acidic mine water; in sample 5A, Mg(OH)\textsubscript{2} contributed to increase the pH from 2.71 to around 4. Most of the coagulants especially FeCl\textsubscript{3} contributed to change alkaline pH to acidic; however, Mg(OH)\textsubscript{2} increased the pH by around 0.5 under such conditions, while af-PFCl did not affect the pH.

**Optimization of flocculation**

Flocc formation, growth and sedimentation are time dependent; the potential of flocc formation and growth is mostly affected by the coagulant type, ionic strength, water pH, suspended particles, coagulant dilution, shear, molecular weight and process conditions. The reactivity of the coagulant through adsorption and neutralization promotes formation of bigger and stable flocs likely to settle at faster rate. The settling time is an important parameter to consider for the optimization of the flocculation process.

Figure 7a shows the turbidity removal by the organic polymer [GK-cl-P(AAm-co-AA)/Fe\textsubscript{3}O\textsubscript{4}] over a period of time (5, 10, 15 and 60 min); it is observed that except for the samples 2A and 3A where the turbidity removal remains stable after 5 min, there is improvement in turbidity removal with the increase in time in samples 1A, 4A and 5A. However, the little increase in turbidity removal over time suggests that our organic polymer has a quick reactivity with suspended particles resulting in flocc growth at faster rate and hence the rapid sedimentation; in this case, an optimum sedimentation time could be considered to be 5 min. The effect of time on the turbidity removal by FeCl\textsubscript{3} was inconsistent in samples 4A and 5A (Fig. 7b), but the general trend also suggests a rapid sedimentation achieved after 5 min.

The coagulant alum (Al\textsubscript{2}(SO\textsubscript{4})\textsubscript{3}) exhibited a relatively slow and poor reactivity during the treatment of samples 1A, 2A and 5A judging by the low turbidity removal after each sedimentation time (5, 10, 15 and 60 min), corresponding to 68.99–74.84–83.21–89.75 % (1A) and 35.88–72.779–72.963–88.35 % (2A), respectively (Fig. 7c). Although the quicker reactivity was achieved in samples 3A and 4A, the performance of alum was mostly poor. The coagulant Mg(OH)\textsubscript{2} exhibited almost a similar performance as alum with optimum turbidity removal only after 10 min in samples 1A and 3A and 15 min in sample 2A.

Except for samples 3A and 4A, it was observed that the turbidity removal efficiency of af-PFCl was improved with aging especially in acidic mine water samples (1A and 5A) (Fig. 7e); in this case, prolonged retention time for settling enables the formation of larger flocs during nucleation to effectively reduce turbidity. Studying the impact of pH on floc structure with polyferric chloride (PFCI) as coagulant, Cao et al. (2011) came to realize that the formation of flocs was much slower when coagulation condition was acidic.

**Simultaneous removal of metals from mine water**

**Effect of settling time**

The removal of metals can be easily predicted when using synthetic solutions; it is, however, more complex for turbid solutions, as more than one mechanisms including charge neutralization, enmeshment and possibly hydroxylation–precipitation may be involved in the removal of metals. The type of coagulant and metal is therefore likely to determine the flocculation mechanism as well as the rate of agglomeration of flocs and settling; in this study, the metal removal capacities of four coagulants were evaluated by considering the settling speed. Figure 8a–e shows that there was an increase in removal rate with time, but the optimum time recorded using each coagulant was 15 min after which no significant removal occurred. However, the removal capacity did vary among the coagulants. Better coagulants are likely to effectively remove the metal at a faster rate; it was observed that the organic polymer nanocomposite [GK-cl-P(AAm-co-AA)/Fe\textsubscript{3}O\textsubscript{4}] performed better than the other coagulants. The pH of the solution is likely to affect the net charges of the elements present, but most importantly a high pH may lead to metal hydroxide precipitation which is different to the removal as a result of coagulation and flocculation; removal of metals from acidic and non-acidic mine solutions was therefore important to evaluate the exact mechanism of metal removal. The results did show that nickel was much soluble in acidic solutions (samples 1A and 5A). According to the hydroxocomplex stability, Ni\textsuperscript{2+} is more soluble than Cr\textsuperscript{6+} and Pb\textsuperscript{2+}, and in previous works Ni\textsuperscript{2+} removal was achieved at relatively high pH; using tannin-based flocculant agent, Heredia and Martin (2009) required an increase in pH to 9 for optimum removal of Ni\textsuperscript{2+} while in separate study, using polyampholyte chitosan derivatives (–N-carboxyethylated chitosans), Bratskaya et al. (2009) observed a 60 % removal
of Ni$^{2+}$ at pH 7.5, but better removal of Ni$^{2+}$ from wastewaters was achieved at pH 9 which was ascribed to increase in complexation degree of the metal in hydroxycyplexes. In this study, the removal of Cr$^{6+}$ and Pb$^{2+}$ was, however, not affected by the pH, and they were effectively removed even below their pH precipitation zone; the removal in this case was likely due to a combination hydroxylation and colloids entrapment in the case of inorganic flocculants, while in the case of the organic polymer which has a net negative charge, surface charged neutralization and bridging may have also played a role. The metals of interest being of positive charge

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Fig. 8 Influence of settling time on metal ions removal from various mine effluents samples: a sample 1A, b sample 2A, c sample 3A, d sample 4A, e sample 5A
Fig. 9 Schematic diagram of the adsorption of metal ions onto the organic flocculant

Where, $M^{2+} = Pb^{2+}, Ni^{2+}$

Table 3 Compliance of residual metals concentrations to SABS (2005) drinking water quality guideline

<table>
<thead>
<tr>
<th>Flocculants</th>
<th>Metals</th>
<th>Sample 1A</th>
<th></th>
<th></th>
<th>Sample 2A</th>
<th></th>
<th></th>
<th>Sample 3A</th>
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<th>Sample 4A</th>
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<td>GF</td>
<td>P</td>
<td>%R</td>
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<td>P</td>
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</table>

GF guideline fitness, R removal, P p value
which is opposite sign to the charge of the organic polymer could be attracted to the surface of the latter according to the principle of surface charged neutralization (Fig. 9). On the other hand, the long chain of the organic polymer tends to form a loop that attaches to other particles in water, resulting in bridging of particles (Kolya and Tripathy 2013). Among all the metals, Pb$^{2+}$ was easily removed from solutions; it has been reported (Fosso-Kankeu and Waanders 2014) that higher atomic weight leads to preferential binding of metal to the adsorbent material, and furthermore, binding affinity of metal to the adsorbent increases with the electronegativity of the metal (Fosso-Kankeu et al. 2014). In this case, Pb$^{2+}$ has a higher atomic weight and electronegativity compared to Cr$^{6+}$ and Ni$^{2+}$, which may partly explain the higher removal of Pb$^{2+}$ compared to other metals.

Achievement of acceptable level of metal ions in water

Evaluation of the optimum time of settling was done by statistical comparison with metal removal rate at the various settling time considered, and it was observed that higher metal removal was achieved between 15 and 60 min, while there was no significant difference [p values 0.126 (sample 1A), 0.139 (sample 2A), 0.748 (sample 3A), 0.076 (sample 4A) and 0.382 (sample 5A)] between the two settling times (Table 3). Therefore, the optimum time of 15 min was considered to evaluate the water quality fitness to the South African Bureau of Standard guideline (2005) for drinking water.

Heavy metal ions such as Cr$^{6+}$, Pb$^{2+}$ and Ni$^{2+}$ are generally highly toxic and can be responsible for the disequilibrium of the ecosystem in the aquatic environment as well as impacting human life. It is therefore important to ensure effective removal of metals from aqueous solution to a safer level, meeting the available guideline. After 15-min settling time, it was observed that in most of the cases the recommended values were met; however, as observed above, Ni$^{2+}$ was hardly removed from the water and therefore remains in solutions at concentrations above the acceptable value.

Conclusion

The use of the real solutions from mine sites allowed to investigate in this study the impact of pH on the coagulation efficiency; this enabled to distinctively highlight the behavior of inorganic as well as organic coagulants during turbidity removal. It is therefore important to mention that the solution pH significantly influences turbidity removal efficiency as well as the sedimentation rate, but depending of the type of coagulant, the response to this influence varied. It is observed that the organic coagulant [GK-cl-P(AAm-co-AA)/Fe$_3$O$_4$] and the inorganic polymer performed better in a wide range of pH, while monomeric inorganic coagulants performed well under alkaline condition. The coagulants used in this study did not correct the acidic pH to recommended value. The removal of metal ions during flocculation process was achieved after an optimum settling time of 15 min reaching the SABS’s (2005) recommended values in drinking water for Pb$^{2+}$ and Cr$^{6+}$, but not Ni$^{2+}$ which was poorly removed especially in acidic solutions.

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