Destruction of cyanide and thiocyanate ions by bacterial and chemical methods

This article deals with the destruction of cyanide and thiocyanate ions by bacterial and chemical methods. Adsorption with granulated activated carbon and oxidation with sodium pyrosulfite in the presence of a copper sulfate catalyst were used as methods of chemical destruction. The bacterial method of destruction was carried out using the bacterial consortium *Agrobacterium tumefaciens* AC-1, *Agrobacterium tumefaciens* AC-2 and *Pseudomonas sp.* AC-3. The control solutions were placed under identical conditions without the addition of any reagent. Sodium pyrosulfite was effective reagent at removing cyanide ions with 97 % efficiency in 1–1.5 hours, while the bacterial consortium was the most productive against thiocyanate ions, resulting in 99 % removal at 120 hours. Thus, the significant amount of cyanide ions has been removed chemically in the first stage, to reduce the effect of high cyanide concentrations on the bacterial process. In the second stage, the cyanide ions have been removed until the MPC levels. In addition, thiocyanate ions were neutralized completely. Model experiments with technological solutions from the gold mining plant were carried out. According to the results of the studies, the respective degrees of bacterial destruction of CN- and SCN- ions are 99 % and 99.9 %. The residual concentration of CN- and SCN- ions in the treated solution was ≤0.1 mg/l and ≤4 mg/l, respectively. These amounts correspond to the maximum permissible concentrations approved in the Republic of Kazakhstan and CIS countries.

Keywords: cyanide, thiocyanate, destruction, removal efficiency, degradation, sodium pyrosulfite, granulated activated carbon, bacterial consortium.

**Introduction**

Environmental protection has a big priority not only in the Republic of Kazakhstan, but also at international level. One of the biggest sources of contamination of the environment is a waste from the mining and processing industries. Disposal of cyanide- and thiocyanate containing residues after manufacturing is a big concern for the current waste management companies worldwide [1].

There are sufficient number of techniques for the treatment of waste water with the high content of cyanide- and thiocyanate ions, mainly, they are chemical approaches like hydrogen peroxide treatment [2], adsorption with activated carbon [3], alkaline chlorination [4], iron sulfide and zinc sulfate treatment, ozonation, UV and electrochemical neutralization [5, 6]. Most of them showed an ability to remove cyanide from the liquid phase, but with low efficiency. In addition to the high cost, they might cause secondary toxic compounds.

**Methods**

Quantitative accounting of microorganisms was carried out with the use of methods of serial dilutions [7], Goryaev's counting chamber and «Zeiss Standart 25» microscope with phase contrast device [8]. The pH and redox potential (Eh) were determined through the use of universal analyzer «Mettler Toledo Seven Multi S47-Ks». Thiocyanate concentrations were determined photometrically with appropriate complex [9], while the determination of cyanides was conducted use of photometry with pyridine and barbituric acid [10] and by titration with silver nitrate [11]. The determination of the metal concentration was carried out using the Kvant-2AT atomic absorption spectrometer with atomization in a graphite furnace [11].

**Results**

High concentrations of cyanide and thiocyanate ions are commonly observed in industrial water after cyanide leaching of gold-containing sulfide ores or concentrates. The formation of thiocyanate occurs during gold cyanidation as a product of the reaction between cyanide ions and sulfur compounds (including elemental sulfur) [12]:

\[ S^0 + CN^- \rightarrow SCN^- \]
Depending on the sulfur content and the number of turnover cycles, the concentration of thiocyanates in sewage or circulating waters can range from several milligrams to several grams per liter. Various model solutions containing CN\(^-\) ions with a concentration of 300 mg/l and SCN\(^-\) ions of 2500 mg/l were prepared for experiments. Adsorption with activated carbon and oxidation with sodium pyrosulfite in the presence of a copper sulfate catalyst were used as the methods of chemical destruction. The bacterial method of destruction was carried out by a consortium of bacteria Agrobacterium tumefaciens AC-1, Agrobacterium tumefaciens AC-2, Pseudomonas sp. AC-3. As a control in the above experiments, the control solution was prepared under identical conditions without adding any reagents.

Granular activated carbon (GAC) with a particle size of 2–4 mm and a density of 0.4 g/ml was used as an adsorbent. Chemical treatment of cyanide and thiocyanate was carried out with sodium pyrosulfite in an amount of 5 g/g for CN\(^-\) and 7 g/g for SCN\(^-\) in the presence of a copper sulfate catalyst at pH 9.0–10.0.

Bacterial degradation was carried out by a consortium of bacteria at temperature 30 \(^\circ\)C. pH of the medium was maintained with a solution of sodium hydroxide at a level of 9.0–9.5. The experiments were carried out on an orbital shaker at 200 rpm. The results of the experiments are shown in Table 1 and Figures 1, 2.

**Table 1**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Duration, hours</th>
<th>Control</th>
<th>Bacterial consortium</th>
<th>GAC</th>
<th>Na(_2)S(_2)O(_5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(C_{ini}, \text{mg/l})</td>
<td></td>
<td>CN(^-)</td>
<td>SCN(^-)</td>
<td>CN(^-)</td>
<td>SCN(^-)</td>
</tr>
<tr>
<td>0</td>
<td>300</td>
<td>279</td>
<td>2488</td>
<td>240</td>
<td>2188</td>
</tr>
<tr>
<td>12</td>
<td>279</td>
<td>2488</td>
<td>225</td>
<td>175</td>
<td>1925</td>
</tr>
<tr>
<td>24</td>
<td>270</td>
<td>2481</td>
<td>225</td>
<td>175</td>
<td>1925</td>
</tr>
<tr>
<td>36</td>
<td>262.5</td>
<td>2475</td>
<td>150</td>
<td>156</td>
<td>1800</td>
</tr>
<tr>
<td>48</td>
<td>247.5</td>
<td>2469</td>
<td>112.5</td>
<td>125</td>
<td>1500</td>
</tr>
<tr>
<td>96</td>
<td>217.5</td>
<td>2444</td>
<td>57</td>
<td>125</td>
<td>1300</td>
</tr>
<tr>
<td>120</td>
<td>202.5</td>
<td>2425</td>
<td>57</td>
<td>25</td>
<td>1200</td>
</tr>
<tr>
<td>The removal efficiency, %</td>
<td></td>
<td>32.5</td>
<td>3</td>
<td>81</td>
<td>99</td>
</tr>
</tbody>
</table>

![Figure 1. The removal efficiency of cyanides with chemical and bacterial methods](image_url)
Table 1 and Figure 1 shows that the most effective removal of cyanide ions has been observed when Na$_2$S$_2$O$_5$ was used. The removal of CN$^-$ was 97 % in 1–1.5 hours. When the granular activated carbon was used, the removal of cyanide ions was also intensive and achieved 90 % in 46 hours, while destruction occurred slower with the use of bacterial consortium, achieving 81 % in 96 hours. Probably, this result is related to the inhibition of bacterial growth due to the increased concentration of cyanide ions [13].

Figure 2. The effectiveness of chemical and bacterial methods at thiocyanate removal

The consortium of bacteria was able to remove 99 % of the thiocyanate ions in 120 hours, showing the highest efficiency. However, other reagents have shown less activity at the same time with 52 % and 9 % for GAC and Na$_2$S$_2$O$_5$, respectively. In the control solutions, the decrease in the concentration of cyanide and thiocyanate ions occurred under the influence of atmospheric oxygen (Table 1 and Fig. 2).

Thus, sodium pyrosulfite was the most efficient reagent at removing cyanide ions with 97 % efficiency in 1–1.5 hours, while bacterial consortium was the most productive against thiocyanate ions, resulting 99 % removal at 120 hours.

In this regard, a two-stage combined technology for the removal of cyanide and thiocyanate was developed. At the first stage, in order to reduce the effect of high concentrations of cyanide on the bacterial process, a significant amount of cyanide ions is removed chemically. The second stage is the removal of the cyanide ions until the MPC levels, also full neutralization of the thiocyanate ions.

The results of previous experiments have shown that the most optimal method is combining of the chemical (sodium pyrosulfite treatment) and bacterial methods. In other words, conversion of the cyanides to the less toxic thiocyanates with subsequent bacterial degradation is a solution to the problem of wastewater treatment.

Samples of technological solutions from the Aksu gold extraction plant (Aqmola region, Kazakhstan) were selected for model experiments. The mineral ores and concentrates of the plant are treated with cyanide solutions. Table 2 shows the chemical analysis of the samples taken.

<table>
<thead>
<tr>
<th>Sample</th>
<th>CN</th>
<th>SCN</th>
<th>Cu</th>
<th>Fe</th>
<th>Zn</th>
<th>Mg</th>
<th>As</th>
<th>Ni</th>
<th>Sb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Industrial solution, mg/l</td>
<td>221</td>
<td>3570</td>
<td>15</td>
<td>730</td>
<td>7</td>
<td>2</td>
<td>1,7</td>
<td>3</td>
<td>3</td>
</tr>
</tbody>
</table>
In the first stage, cyanides and metal cyanide complexes were oxidized with sodium pyrosulfite (5 g/g), in the presence of copper ions at pH 9.0–10.0.

The obtained data (Fig. 3) show that the neutralization degree is 97% in 90 minutes, the residual CN– concentration was 6.6 mg/l, whereas the concentration of thiocyanate ions increased due to the oxidation of cyanide ions, which amounted to 3785 mg/l.

In the second stage, the processing solution was passed through a bioreactor for destruction of the residual content of cyanide and thiocyanate by bacterial consortium immobilized on a zeolite. The results of the experiments are shown in Figures 4, 5.

Figure 3. Effectiveness of CN– removal with sodium pyrosulfite

Figure 4. Effectiveness of CN– removal with bacterial consortium *Agrobacterium tumefaciens*AC-1, *Agrobacterium tumefaciens*AC-2, *Pseudomonas sp.* AC-3
Destruction of cyanide and thiocyanate ions …

Based on the results of the experiments, the degree of bacterial destruction of CN⁻ and SCN⁻ are 99 % and 99.9 %, respectively. Moreover, respective residual concentrations of CN⁻ and SCN⁻ ions in the treated solution were ≤0.1 mg/l and ≤4 mg/l, which corresponds to the maximum permissible concentrations approved in the Republic of Kazakhstan and CIS countries.

References

N.K. Жаппар, В.М. Шайхутдинов, О.А. Тен, Д.С. Балпанов, Е.Н. Канафин, Р.А. Ханнанов, Р.Ш. Еркасов, А.А. Бакибаев, А.Т. Кедзикбаева

Бактериальные-химические тезисы – жёл вионанат-ионов деструкции

Макдада бактериальные және химические деструкция меселелері каратырылған. Химическая деструкция меселері ретінде өсім сулуғын калыптастырылыған натрий пиросульфатын тоқтыру және беңесерлірлігін көмірмен адсорбциялау қолданылды.

Деструкцияның бактериальные тәсілі Agrobacterium tumefaciens AC-1, Agrobacterium tumefaciens AC-2 және Pseudomonas sp. AC-3 бактериялар консорциумын қалыптастырып, тұжырымды қадам алау. Максат мағын арқылы шарттарға жауап қоюға қосылған моделдік әрекетпен байланысты. Цианид іондар алған шығару ұсын әр тәсіл натрий пиросульфит қызқылына байлық: 97 % – 1,5 сағ; Agrobacterium tumefaciens AC-1, Agrobacterium tumefaciens AC-2 және Pseudomonas sp. AC-3 бактериялар консорциумы натрий пиросульфит алуға ықпал етеді: 120 сағ — 99 %. Сондықтан, бірнеше кезіндегі бактериальные процессті бюджет болыу мүмкін цианидтің концентрациясын құра аударған және адсорбцияның жұмыс іздейді.

Кізім сөзір: цианид, тиоцианат, деструкция, алған шығару тиімділігі, азу, натрий пиросульфиті, туынды тәсіл, бактериялар консорциумы.

References

Destruction of cyanide and thiocyanate ions ...


