Optimizing soil washing remediation of mercury contaminated soil using various washing solutions and solid/liquid ratios

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Abstract. Most of the artisanal and small-scale gold miners in Indonesia as in the case of those who are in Banten Province, still use the amalgamation process in the gold extraction process. Therefore, mercury contamination could not be avoided. As a result, it was found that the concentration of mercury-contaminated soil in Lebak, Banten Province was detected as high as 136.9 mg/kg. Since the contamination process occurred for a long time, making the formation and mobility of complex mercury in soil increased by time. Soil washing is one remediation technology that can be applied in the recovery of mercury-contaminated soil. This study aims to determine the optimum conditions of the soil washing process: effect of pH, the concentration of washing solutions and ratio of solid/liquid. Furthermore, the value of the distribution coefficient, desorption of mercury in the soil and fractionation of mercury in the soil were also observed. Potassium Iodide was found to be the best washing solution among others used in this study. The optimum condition was obtained at pH 2, the concentration of washing solution at 0.25 M and the solid/liquid ratio of 1:15. Under these conditions, mercury desorption efficiency reached 86.9% with the value of the distribution coefficient of 0.185. Mercury fractionation analysis in the contaminated soil showed that the mobile > semi-mobile > non-mobile fractions. Potassium iodide was able to desorb at about 84% of mobile fractions, 97% of semi-mobile and 25% non-mobile mercury.

Keywords: Soil washing, mercury, desorption, distribution coefficient, fractionation

1 BACKGROUND

Artisanal and small-scale gold miners (ASGM) is one of the anthropogenic sources of mercury pollution in the environment. Based on the report of the Ministry of Environment and Forestry, Government of Indonesia there are as many as 850 ASGM points consisting of more than 250,000 miners spread all over Indonesia [1]. According to Ruslan and Khairuddin [2] the process of gold ore beneficiation that carried out by ASGM is called the amalgamation process. Amalgam formation process is carried out in a drum, called the batch amalgamator that using mercury as gold binding agent. Mercury added to drum is reduced at the end of the process due to the processing stage that is allowing mercury to carry away in the wastewater (tailings).

Mercury is a metal that has the potential to be harmful to the environment, especially in areas affected by the gold mining industry. The soil has an important role in the mercury cycle both acting as a medium and source for biota metabolisms, atmosphere and hydrological cycles. Soil can affect mercury forms (species of mercury) through chemical, physical and biological processes, thus affecting solubility, biological accessibility, ecology and toxicological effects of mercury in the environment. Metal mercury in the form of Hg⁰ is the most non-toxic mercury compound because it is not soluble in water, and it is not bound to animal tissues and cannot be consumed in low-level animals or microbes [3, 4]. Hg⁰ can be oxidized in the atmosphere to inorganic mercury (Hg²⁺) which is found in various forms of salts such as chloride, nitrate or sulfide. Hg²⁺ is a reactive form of mercury that has a high affinity for animal/plant tissue and can be consumed by microorganisms. Besides Hg²⁺ functions as a substrate in the process of bacterial methylation under anaerobic conditions, such as in sediments and inundated soils that will form methyl mercury compounds [5].

The fate, pathway and toxicity of mercury in the soil depend on the forms of mercury. Organic mercury species such as methyl mercury have a higher toxicity level and are more easily biologically accumulated when compared to inorganic mercury. This is because organic mercury has large mobility compared to inorganic mercury. The species of mercury in the soil included in the semi-mobile category is a less toxic species of
mercury. Whereas the non-mobile mercury species are chemically toxic and only mercury sulfide is non-toxic [6, 7]. Therefore, it is necessary to remediate mercury-contaminated soil due to tailing disposal of artisanal and small-scale gold mining activities. One remediation technology that can be applied in the recovery of semi-volatile or volatile compounds mercury-contaminated soils is by washing the soil or so called a soil washing method. Soil washing is a technique to recover contaminated soil which has several advantages, such as the recovery process has a short time, can be used widely, is economically feasible and can be applied to soil contaminated with semi-volatile or volatile compounds to include mercury [8, 9].

2 METHODOLOGY

2.1 SAMPLE PREPARATION

The composite soil samples were homogenized using the cone quarter method, then using a ± 200 mesh sieve to separate rocks and large particles [10]. Homogenized soils were then dried at room temperature for 2-3 days [11]. Soil samples were further analysed to determine the initial characteristics of the soil include physical and chemical characteristics. The physical characteristics were soil texture (silt, clay, and sand), moisture content and soil permeability. While the chemical characteristics included soil pH, Cationic Exchange Capacity (CEC), C-organic, C, N Total, and P Total as well as metal content. The initial concentration of mercury was 136.9 mg/kg. Data provided in result and discussion part was produced from 2 replicates.

2.2 SOIL WASHING SOLUTION

Four (4) washing solutions were used. They were sodium ethylenediaminetetraacetic acid (Na2EDTA), potassium chloride (KCl), potassium iodide (KI), and water as control. Each beaker glass contained 20 grams of washing solutions were mixed with water with a ratio of 1:10 (gr/mL). Batch stirred at a speed of 100 rpm for 3 hours. Then after 3 hours of mixing the sample was taken and centrifuged at a speed of 1000 rpm for 15 minutes. The supernatant was filtered using a vacuum pump filter, to separate the solution from washing and sediment. The filtered supernatant was then destructed before measuring mercury concentration (Hg) using the Atomic Absorption Spectrophotometry (AAS) method.

2.3 OPTIMUM CONDITIONS DETERMINATION

A total of 20 grams of contaminated soil were mixed with a 200 ml solution of the best washing solution (KI) at a concentration of 0.2 M with a variation of pH 2, 3, 5, 7 and 8. The mixture of this soil sample and a solution of KI stirred at a speed of 100 rpm for 3 hours. The mixture was centrifuged at a speed of 1000 rpm for 15 minutes. The supernatant was filtered using a vacuum pump filter and then destructed before measuring mercury concentration (Hg) using the Atomic Absorption Spectrophotometry (AAS) method. Under the optimum pH, the study followed by determining the optimum concentration of KI with variations of 0.1 M, 0.15 M, 0.2 M, and 0.25 M respectively. Finally, after obtaining the optimum pH and KI concentration as washing solution, the best ratio of solid/liquid ratio at 1:10, 1: 12.5, 1:15 and 1: 17.5 (gr/mL) respectively, were determined.

2.4 DESTRUCTION PROCEDURE

The determination of total mercury concentration was obtained after going through the destruction process based on the EPA 7471B method of mercury in solid or semisolid waste (manual cold-vapour technique). The subsequent determination of fractionation of mercury speciation followed the EPA 3200 mercury species fractionation and quantification by microwave-assisted extraction or solid-phase extraction.

2.5 DISTRIBUTION COEFFICIENT & MASS BALANCE

Soil washing performances were carried out by determining the effect of solid/liquid ratio, pH and concentration of potassium iodide washing solution in mercury removal in polluted soil. The removal efficiency was simply determined by the following equation:

\[ Efficiency(\%) = \frac{(C_{\text{initial}})-(C_{\text{effluent}})}{C_{\text{initial}}} \times 100 \]  

Furthermore, to find out the distribution coefficient (Kd) was calculated with the equation as follows:

\[ Kd = \frac{C_{1}}{C_{5}} \]  

Then the mass balance calculation was carried out, this calculation is intended to estimate the concentration of contaminants in the liquid phase at the end of the process in optimum conditions with the following equation:

\[ C_{f} = \frac{(M_{f}C_{2})-(C_{1}V_{s})}{M_{f}} \]  

Where Kd is the distribution coefficient of metal mercury (Hg), C1 is the concentration of Hg in the liquid phase (washwater) and C5 is the concentration of Hg in the solid phase or in the soil, Cf is the final mercury concentration after washing process, Mf is the soil mass and Vf is the volume of washing solution.

3 RESULTS AND DISCUSSION

3.1 INITIAL CHARACTERISTICS OF SOIL

Table 1 indicated the physical and chemical properties of soil sample. Soil distribution used consists of 85% silt, 12.7% fine sand and 2.2% clay with the initial concentration of total Hg reached 136.9 mg/kg. Since the soil has high content of silt and sand (more than 90% in total), soil water content was only 2.46% because the
soil has low water holding capacity. The pH of soil sample was high because the soil was found to have high content of Calcite minerals (CaCO₃). Moreover, it could be seen that the C-organic content was high making it is difficult to desorb mercury from soil due to reaction between mercury with organic compounds.

Table 1. Soil Samples Physical & Chemical Properties

| No | Parameter       | Result   | Unit |
|----|-----------------|----------|------|
|    | Physical Properties |         |      |
| 1  | Water Content   | 2.46     | %    |
| 2  | Specific Gravity| 2.68     |      |
|    | Chemical Properties |        |      |
| 3  | pH              | 10.25    |      |
| 4  | Hg Initial      | 136.9    | mg/kg|
| 5  | C-organic       | 0.617    | %    |
| 6  | N total         | <0.004   | %    |
| 7  | P total         | 0.031    | %    |
| 8  | CEC             | 11.57    | cmol/kg|
| 9  | Organic Matter  | 1.829    | %    |

Metal minerals affect the cation exchange capacity (CEC). According to Pansu and Gautheryou [12] cation exchange capacity is determined by the amount of clay minerals and organic matter present in the soil. The mechanism of mercury adsorption in clays occurs due to the presence of Ca²⁺, Mg²⁺, Na⁺, and K⁺ as shown in table 2. Mercury ions can be replaced with monovalent cations at the site exchange.

Table 2. Metals Mineral in Soil Samples

| No | Element | Value (%) | Compound | Value (%) |
|----|---------|-----------|----------|-----------|
| 1  | C       | 10.92     | C        | 10.92     |
| 2  | O       | 42.47     | -        | 1.46      |
| 3  | Mg      | 0.88      | MgO      | 10.11     |
| 4  | Al      | 5.35      | Al₂O₃    | 64.12     |
| 5  | Si      | 29.97     | SiO₂     | 0.17      |
| 6  | S       | 0.07      | SO₃      | -         |
| 7  | Cl      | -         | -        | -         |
| 8  | K       | 3.78      | K₂O      | 4.55      |
| 9  | Ca      | 2.06      | CaO      | 2.88      |
| 10 | Fe      | 4.5       | FeO      | 5.79      |
| 11 | Hg      | -         | HgS      |           |

3.2 INITIAL FRACTIONATION OF MERCURY SPECIES

In this study, the fractionation process was carried out in stages in determining the total fractionation of organic, inorganic, semi-mobile and non-mobile. Fractionation was determined based on the EPA 3200 procedure on mercury species fractionation and quantification by microwave-assisted extraction, selective solvent extraction and/or solid-phase extraction. The results of initial fractionation before the soil washing process can be seen in Table 3.

Based on the results of mercury fractionation, it could be seen that extractable mercury < non-extractable mercury. This indicated that the concentration of mercury dissolved in the washing solution was expected to be relatively small and would be dominated by vaporized mercury. Fractionation results were dominated by organic and semi-mobile mercury, which were 41.61 mg/kg and 51.71 mg/kg, respectively.

Table 3 Total initial concentration of mercury fraction

|                   | Mercury Fractionation | Mercury Concentration (mg/kg) |
|-------------------|-----------------------|-------------------------------|
|                   | Extractable mercury   | Organic                       | 46.61 |
|                   | Inorganic mercury     | Inorganic                     | 6.00  |
|                   | Non-extractable mercury| Semi-mobile                  | 51.71 |
|                   |                       | Non-mobile                    | 39.46 |

3.3 EFFECT OF THE WASHING SOLUTION ON MERCURY REMOVAL

Basically, water has the ability as a washing solution because water is a monodentate chelating agent. However, it was found that the removal efficiency of mercury using water alone as washing solution was low. Based on the results of the study of soil washing processes using water could only remove metal mercury at around 15.18%. Hg metal is very strongly bound with organic compounds and sulphides in the soil to form complex compounds that are insoluble. Therefore, chemicals washing solution is required to desorb mercury from the soil. The results showed that the ability of Na₂EDTA, KCl and KI in removing mercury from soil at pH 2 was found at around 22.93%, 73.94% and 84.49% respectively. So, In the case of mercury chelating agents, potassium iodide was more efficient for mobilizing mercury from the soil as shown in figure 1. According to Wasay, et al. [13] iodide solution is a better chelation agent than EDTA especially in extracting mercury to form complex compounds.

Figure 1. Washing Solutions Removal Efficiency

3.4 PH EFFECT

Soil washing process by using a solution of potassium iodide at acidic pH can increase the solubility and desorption of metal ions resulting in the removal of mercury in the soil increases. This was inline with the opinion of Peng, et. al. [14] which stated that extraction especially in the fast extraction process depended on pH solution where the lower the pH, the faster the extraction process.
after the soil washing process at pH 2 was hing agent in pes of taminants as pH ETMC and RC EnvE KI Cn c ce ntratio M)

0.00  10.00  20.00  30.00  40.00  50.00  60.00
Mercury in Washed Soil (mg/kg)

Figure 2. Effect of pH on Desorption of Mercury using KI as Washing Solution

Figure 2 above shows the correlation between pH of washing solution (KI) with mercury retention in soil. It could be seen that the pH of washing solution affected the ability of the soil to retain mercury significantly. Basically, the lower the pH value, the greater the desorption process and solubility of mercury from the soil. Potassium iodide (KI) as a chelating agent worked effectively at pH 2 in dissolving mercury to reach removal efficiency up to 85.14%. Mercury concentration in the soil after the soil washing process at pH 2 was found around 20.34 mg/kg. The coefficient of solid/liquid distribution (Kd) is used to estimate the mobility and distribution of elements in the environment [15]. The value of each distribution coefficient can be seen in the following table.

Table 4. Effect of pH on the distribution coefficient value

| Sample | Mercury Concentration | Distribution Coefficient |
|--------|------------------------|--------------------------|
|        | Soil (mg/kg)           | Water (mg/L)             |
| pH 2   | 20.34                  | 2.69                     | 0.1322 |
| pH 3   | 37.56                  | 1.59                     | 0.0423 |
| pH 5   | 40.50                  | 0.98                     | 0.0242 |
| pH 7   | 54.25                  | 0.41                     | 0.0076 |
| pH 8   | 55.61                  | 0.33                     | 0.0059 |

3.5 SOLID/LIQUID RATIO

Solid/liquid ratio is a parameter that contributes greatly to the extraction process and removal of contaminants as a whole on contaminated soils, as well affected the amount of residual water produced [16]. Figure 3 indicates the effect of solid/liquid ratio against mercury desorption when KI used as washing solution at pH 2.0.

Table 5. Effect of solid / liquid ratio on distribution coefficient value

| Sample  | Mercury Concentration | Distribution Coefficient |
|---------|-----------------------|--------------------------|
|         | Soil (mg/kg)          | Water (mg/L)             |
| 1 : 10  | 21.28                 | 1.99                     | 0.093 |
| 1 : 12.5| 34.04                 | 0.74                     | 0.021 |
| 1 : 15  | 17.89                 | 2.69                     | 0.150 |
| 1 : 17.5| 20.02                 | 2.06                     | 0.103 |

3.6 WASHING SOLUTION CONCENTRATION (KI)

Figure 4 demonstrated that the greater concentration of chelating agents used, the greater the removal efficiency of mercury metal from contaminated soil. Mercury removal efficiency using different concentrations of KI was found between 68.0% - 88.8% at solid/liquid ratio of 1:15 and at pH 2.0.

At KI concentration of 0.25 M, soil washing process was able to remove mercury from the soil by 88.8% so that the remaining mercury concentration in the soil was 15.6 mg/kg. The use of high concentrations of KI could reduce the volume of washing solutions required in the soil washing process and form complex compounds with metal mercury because of the presence of excess iodide in the washing solution. Allen, et. al, [18] added that the concentration of chelating agents was influenced by soil characteristics, chelating agent types and types of contaminants. Distribution coefficient was best determined when the KI concentration as washing solution was increased to 0.25 M as shown in Table 6.
Table 6. Effect of KI Concentration on distribution coefficient

| Sample | Mercury Concentration | Distribution Coefficient (L/kg) |
|--------|-----------------------|---------------------------------|
|        | Soil (mg/kg)          | Water (mg/L)                    |                                |
| 0.10 M | 44.45                 | 1.66                            | 0.037                          |
| 0.15 M | 27.47                 | 1.82                            | 0.066                          |
| 0.20 M | 17.89                 | 2.69                            | 0.150                          |
| 0.25 M | 15.60                 | 2.88                            | 0.185                          |

The pH value is the most important factor in determining the distribution coefficient (Kd) because the pH value can increase the solubility of one of the compounds against other compounds that cannot be mixed [19]. Furthermore, the higher the concentration of KI resulted in the greater the distribution of mercury in the solution that is equal to 0.185 L/kg. According to Gil, et al. [15] the Kd value is highly dependent on pH, redox conditions, particle size distribution, organic matter content, biological activity, and temperature. However, having examined the distribution coefficient value (d) obtained in this study, it showed that the amount of metal mercury distributed in potassium iodide (KI) solution had low concentration. This was probably due to the mercury contaminated soil used in this study were dominated by volatile compounds so that some of metal mercury vaporized.

3.7 MERCURY FRACTIONATION OF WASHED SOIL

Table 7 indicated that the mercury-contaminated soil was dominated by organic mercury fractionation of 46.61 mg/kg and semi-mobile at 51.71 mg/kg. This was because the sampling site was a former artisanal and small-scale gold mining area that has not been operated since 2016.

Table 7. Mercury Fractionation Concentration Before and After Soil Washing

| Mercury Fractionation | Mercuric Concentration (mg/kg) | Before Soil Washing | After Soil Washing |
|-----------------------|--------------------------------|---------------------|-------------------|
|                       | KI                             | KCl                 | EDTA              | H2O |
| Extractable Mercury-  | Organic                        | 46.61               | 7.90             | 17.03 |
|                       |                                | 37.92               | 48.37            |
| Extractable Mercury-  | Inorganic                      | 6.00                | 0.29             | 3.14 |
|                       |                                | 20.12               | 10.72            |
| Non-extractable       | Semi-mobile mercury            | 51.71               | 1.14             | 2.73 |
|                       |                                | 4.59                | 18.48            |
| Non-extractable       | Non-mobile mercury             | 39.46               | 29.32            | 23.85 |
|                       |                                | 60.21               | 51.82            |

Inorganic mercury (Hg\(^{0}\)) has been involved in the soil microorganism metabolisms to form organic mercury and other complex compounds. Organic mercury fractionation has a higher toxic level compared to other fractions because it is easily mobilized and can accumulate in organisms [20]. After the soil washing process, KI solution was able to remove organic mercury fraction to 7.9 mg/kg, semi-mobile to 1.14 mg/kg, inorganic at 0.29 mg/kg and non-mobile fractions to reach 29.32 mg/kg. These results demonstrated that KI was very suitable to be applied as washing solution in the absorption of mercury in the soil as compared to others chelating agents.

Furthermore, based on kinetics model using Lagergren equation, the soil washing process using KI 0.25 M as washing solution at pH 2 and solid/liquid ratio of 1:15 was found to be fit with pseudo-second-order kinetics model and was suitable kinetics model in describing the desorption kinetics of mercury in the soil. This result indicated that the entire mechanism process was dominated by chemical process. This was following the role of potassium iodide (KI) as chelating agents that can form complex compounds with mercury in the desorption process. Vadivalen and Kumar [21] stated that if the process adsorption/desorption follows the pseudo second order kinetics model, for the whole adsorption/desorption process, the chemical mechanism is an adsorption/desorption rate controller that is greater than that of physical adsorption/desorption. Desorption rate under this optimum condition was determined at a rate of 0.0025/minute as shown in the following table.

Table 8. Soil Washing Adsorption and Desorption Rate

| S/L Ratio | Rate (1/minute) |
|-----------|-----------------|
|           | \(K_{d}\)        | \(K_{ads}\)       |
| 1:10      | 0.0016          | 2.985 \(10^{6}\)  |
| 1:12.5    | 0.0010          | 2.649 \(10^{6}\)  |
| 1:15      | 0.0025          | 5.844 \(10^{6}\)  |
| 1:17.5    | 0.0008          | 5.806 \(10^{6}\)  |

Overall, mercury retention in soils is controlled by the chemical composition of the soil (sulphur and organic matter), but the rate of desorption controlled by the physical properties of the soil (particle size) [22].

4 CONCLUSION

Based on this study, it can be concluded that potassium iodide (KI) was a suitable washing solution to be applied in the absorption of mercury in the soil. The optimum conditions were determined at pH 2, a concentration of washing solution of 0.25 M and a solid/liquid ratio of 1:15. With these optimum conditions, potassium iodide could remove mercury more than 88% with the distribution coefficient value of 0.185 L/kg. Contaminated soil was dominated by organic mercury that was equal to 46.61 mg/kg and semi-mobile mercury fraction of 51.71 mg/kg, inorganic mercury fraction of 6 mg/kg and non-mobile mercury fraction of 39.46 mg/kg. Potassium iodide (KI) solution was able to remove mercury of organic fraction to 7.9 mg/kg, semi-mobile fraction to 1.14 mg/kg, inorganic fraction to 0.29 mg/kg and non-mobile fraction to 29.32 mg/kg. These results demonstrated that potassium iodide was very suitable to be applied in the absorption of mercury in the soil with a desorption rate of 0.0025/minute.
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