Abstract: Massive amount of highly contaminated mining residual materials (MRM) has been left unattended and has leached heavy metals, particularly lead (Pb) and zinc (Zn) to the surrounding environments. Thus, the performance of three immobilizers, raw dolomite (RD), calcined dolomite (CD), and magnesium oxide (MO), was evaluated using batch experiments to determine their ability to immobilize Pb and Zn, leached from MRM. The addition of immobilizers increased the leachate pH and decreased the amounts of dissolved Pb and Zn to different extents. The performance of immobilizers to immobilize Pb and Zn followed the following trend: MO > CD > RD. pH played an important role in immobilizing Pb and Zn. Dolomite in RD could slightly raise the pH of the MRM leachate. Therefore, the addition of RD immobilized Pb and Zn via adsorption and co-precipitation, and up to 10% of RD addition did not reduce the concentrations of Pb and Zn to be lower than the effluent standards in Zambia. In contrast, the presence of magnesia in CD and MO significantly contributed to the rise of leachate pH to the value where it was sufficient to precipitate hydroxides of Pb and Zn and decrease their leaching concentrations below the regulated values. Even though MO outperformed CD, by considering the local availability of RD to produce CD, CD could be a potential immobilizer to be implemented in Zambia.

Keywords: mine waste; contamination; batch experiments; lead; zinc; immobilization; remediation; Kabwe; Zambia

1. Introduction

Kabwe District was one of the most important mining regions in Zambia for almost a century (1902–1994). It was regarded as Southern Africa’s principal lead (Pb)-zinc (Zn) producer, producing
over 1.8 and 0.8 Mt of Zn and Pb, respectively [1]. While in operation, no pollution laws were enforced to regulate the discharge from wastes of the mine; therefore, operations of the mine have left Kabwe with a massive amount of unattended mining residual materials (MRM), which still contain elevated amounts of heavy metals, particularly Zn, Pb, and iron (Fe). Weathering of MRM causes heavy metals to transport from the contaminated sites to the surrounding environments (groundwater, surface water, and soil) [2]. In particular, the redistribution of heavy metals through solute transport processes has been reported to be one of the most dangerous pathways, which invokes harmful effects on water sources of nearby ecosystems and health-threatening to the nearby residents [3–9]. Therefore, the remediation of heavy metals in and around the mine is necessary.

In our recent studies, several potential remediation techniques have been investigated to remediate the contaminated site in Kabwe. Silwamba et al. (2020) [10,11] have proposed the concurrent dissolution and cementation method. The method shows promising results in terms of Pb removal and recovery. However, Zn could not be effectively recovered from the extraction solution, and further investigation is needed. Biocementation by locally available bacteria has been studied by Mwandira et al. (2019) [12,13]. The results indicate that the biocemented material can be effectively used as a covering layer to prevent airborne contamination of metallic dust and infiltration of water into the waste. In the present study, chemical immobilization is introduced as an alternative and practical method to remediate the site.

Remediation techniques for heavy metals polluted sites can be classified into two main categories, in-situ and ex-situ. In general, ex-situ treatment has high efficiency; however, it is less cost-effective than in-situ remediation due significantly to the costs for excavation and transport of large quantities of contaminated materials. In-situ remediation avoids the excavation and transportation costs because of on-site treatments of contaminants. Various kinds of in-situ remediation techniques have been developed to immobilize or extract the heavy metals in the contaminated sites. Among them, chemical immobilization is cheap, easy to implement, and quick in execution [14,15]. Thus, this is the most promising technique, especially to be applied in one of the developing countries.

In in-situ chemical immobilization, the leaching potential of heavy metals from contaminated soils is reduced via sorption and/or precipitation processes by adding chemical agent (immobilizer) into the contaminated area. The performance of a variety of immobilizers, including carbonates, phosphates, alkaline agents, clay, iron-containing minerals, and organic matters, has been evaluated [14–23]. However, most of the studies have been conducted to remediate contaminated soil samples in which they generally contain much lesser metals contents than those in MRM. Moreover, because of the complex interactions between solutes and immobilizers, the definite efficiency of the immobilizer remains site-specific. In other words, there is no guarantee on the effectiveness of a particular immobilizer implemented on different contaminated sites. Therefore, the objective of this study was to evaluate and compare the performance of selected potential immobilizers (e.g., locally available, low-cost) to reduce the mobility of Pb and Zn leached from the highly contaminated sample (MRM).

It is necessary to apply immobilizers that is low cost and abundant in nature for remediating contaminated areas. Hence, in this study, raw dolomite (RD) was selected as one of the potential candidates because it is naturally available in a large quantity in Zambia [24]. It is a carbonate mineral; therefore, it can increase and buffer pH of MRM, leading to more adsorption and precipitation of cationic heavy metal ions [25–27]. In the present study, calcined dolomite (CD) was also used as an immobilizer. The heat treatment was performed to change the carbonate property of dolomite to be more alkaline [28]. As a result, the immobilizer was expected to strongly increase the pH of MRM, favoring the immobilization of heavy metals by hydroxide precipitation in addition to adsorption and precipitation of other secondary minerals. At the same time, commercially available alkaline-based agent, magnesium oxide (MO), was also tested to compare the ability of RD and CD on immobilizing heavy metals in MRM. The current study will provide meaningful information for the development of chemical immobilization to remediate heavy metals contaminated sites in Zambia.
2. Materials and Methods

2.1. Solid Sample Collection, Preparation, and Characterization

MRM was collected from the dumping site of Pb-Zn mine wastes in Kabwe, Zambia. The sampling was done using shovels at random points within the area shown in Figure 1. This leaching residual was selected as one of the representative wastes because the leaching concentrations of Zn and Pb from the waste were higher compared with the other wastes. Moreover, the storage size of this waste occupies more than 50% of the total dumping area. The sample was stored in vacuum bags and transported to the laboratory in Japan with permission by the Ministry of Agriculture, Forestry, and Fisheries of Japan. In preparation, it was air-dried under ambient conditions, lightly crushed, sieved using a 2 mm aperture screen, and kept in a polypropylene bottle before use. Particle sizes of less than 2 mm were chosen to follow the Japanese standard for the leaching test of contaminated soils [29].

![Figure 1. Dumping site in Kabwe; (□) sampling area (top left: 14°27′39″ South, 28°25′42″ East; down right: 14°27′55″ South, 28°26′16″ East).](image)

Three types of immobilizing agents were selected to immobilize Zn and Pb in the waste: raw dolomite, calcined dolomite, and magnesium oxide denoted as RD, CD, and MO, respectively. RD was taken from a dolomite quarry source near the MRM storage site, while CD was prepared by burning RD with particle sizes of less than 2 mm in a furnace at 700 °C for 2 h. MO was commercially available, purchased from Ube Industry, Japan. The same preparation procedure as that for MRM was also applied to these materials.

Chemical and mineralogical properties of all solid samples were characterized on the pressed powder of finely crushed samples (<50 µm) using an X-ray fluorescence spectrometer (XRF) (Spectro Xepos, Rigaku Corporation, Tokyo, Japan) and X-ray diffractometer (XRD) (MultiFlex, Rigaku Corporation, Tokyo, Japan), respectively. Sequential extraction was conducted to evaluate solid-phase heavy metals speciation of MRM. The procedure used in this study was modified from two well-known procedures, Tessier et al. (1979) [30] and Clevenger (1990) [31]. The modification was done by Marumo et al. (2003) [32], and it was widely used to extract the tailings sample, mineral processing wastes, and leaching residues [10,33–36]. The process can divide solid-phase heavy metals bonded to solid into five different phases, including exchangeable, carbonates, Fe-Mn oxides, sulfide/organic matter, and residual. The details of the extraction procedure are summarized in Table 1. The extraction was done on 1 g of the <2 mm MRM sample. Between each step, the extractant solution of the previous step was retrieved by centrifugation of the suspension at 3000 rpm for 40 min to separate the residue out of the leachate. The residual was then washed with 20 mL of deionized water. Finally, the washing and extractant solutions were mixed, diluted to 50 mL, filtrated through 0.45-µm Millex® filters, and kept in polypropylene bottles prior to chemical analysis.
Table 1. Sequential extraction for heavy metals speciation.

| Step | Extractant | pH | Liquid to Solid Ratio (mL/g) | Temperature (°C) | Duration (h) | Mixing Speed (rpm) | Extracted Phase |
|------|------------|----|-----------------------------|------------------|--------------|-------------------|----------------|
| 1    | 1 M MgCl$_2$ | 7  | 20/1                        | 25               | 1            | 120                | Exchangeable    |
| 2    | 1 M CH$_3$COONa | 5  | 20/1                        | 25               | 5            | 120                | Carbonates      |
| 3    | 0.04 M NH$_2$OH·HCl in 25% acetic acid | 20/1 | 50                           | 5               | 120          |                   | Reducible       |
| 4    | 0.04 M NH$_2$OH·HCl in 25% acetic acid; 30% H$_2$O$_2$; 0.02 M HNO$_3$ | 36/1 | 85                           | 5               | 120          |                   | Oxidizable      |
| 5    | Calculated  |    |                             |                  |              |                   | Residual        |

2.2. Batch Leaching Experiments

Batch leaching experiments were performed using 250 mL polypropylene Erlenmeyer flasks with a lateral reciprocating shaker (EYELA Multi Shaker MMS, Tokyo Rikakikai Co., Ltd., Tokyo, Japan). All batches were conducted under ambient conditions by mixing 15 g of solid sample to 150 mL of deionized water (1:10 solid-to-liquid ratio) at 200 rpm for 6 h. Five replications of the leaching tests of MRM were conducted, while a single run of every immobilization experiment was done. The reason is that in immobilization tests, we adjusted the addition of immobilizers and did not control the pH of the suspension. In other words, pH is determined by the complex chemical and physical interactions between immobilizer and MRM. Thus, a variation of pH can be easily observed even though the same mixing ratio between MRM and immobilizer and the conditions is employed. To avoid the uncertainty of the variation of pH at the same immobilizer:MRM mixing ratio, we varied the immobilizer:MRM mixing ratios at 1:100, 3:100, 1:20, and 1:10 to evaluate the performance of immobilizers. After 6 h of shaking, the pH, electrical conductivity (EC), redox potential (ORP), and temperature were immediately measured. The leachates were collected by first centrifuging the mixtures at 3000 rpm for 40 min to separate the suspended particles. The supernatants were then filtered with 0.45-µm Millex® filters (Merck Millipore, Burlington, MA, USA) and kept in air-tight polypropylene bottles prior to chemical analysis.

2.3. Chemical Analysis of Liquid Samples

Inductively coupled plasma atomic emission spectrometer (ICP-AES) (ICPE-9000, Shimadzu Corporation, Kyoto, Japan) and inductively coupled plasma atomic emission mass spectrometry (ICP-MS) (ICAP Qc, Thermo Fisher Scientific, Waltham, MA, USA) were used to quantify the dissolved concentrations of heavy metals and coexisting ions. The analyses were performed on the pretreated liquid samples in which 1% by volume of 60% nitric acid (HNO$_3$) was added to the liquid samples. The acidification was done to make sure that all target elements were in a dissolved form. The non-acidified samples, on the other hand, were used to determine alkalinity or acid resistivity. This parameter is generally reported as bicarbonate concentration (meq/L), quantified by titration of a known volume of sample with 0.01 M sulfuric acid (H$_2$SO$_4$) until pH 4.8. All chemicals used were reagent-grade.

2.4. Geochemical Modeling

An aqueous geochemical modeling program, PHREEQC (Version 3, U.S. Geological Survey, Sunrise Valley Drive Reston, VA, USA) [37], was used to aid in the interpretation of the experimental results. The program can determine the parameters that may affect the mobility of heavy metals from MRM, such as stability of minerals and chemical species. The input data included temperature, pH, ORP, and concentrations of heavy metals and other coexisting ions. Thermodynamic properties were taken from the WATEQ4F database.
3. Results and Discussion

3.1. Properties of Solid Samples

The mineralogical and chemical compositions of MRM and immobilizers, including RD, CD, and MO, are listed in Tables 2 and 3, respectively. MRM was composed of anglesite (PbSO₄) as a primary mineral; zinkosite (ZnSO₄) and quartz (SiO₂) as the second-highest; and goethite (FeOOH), hematite (Fe₂O₃), and gypsum (CaSO₄·2H₂O) as the minor minerals. Anglesite and zinkosite are commonly found as the weathering products of Pb- and Zn-sulfides under natural oxygenated environments [38,39]. Therefore, the presence of these two minerals indicates that MRM has already been exposed to the surface environment for a long time before the sampling was done. It can also be expected that other than goethite and hematite, MRM also contained amorphous iron-(hydr)oxides and iron-sulfate salts (e.g., melanterite, coquimbite) since goethite and hematite were found as the miner minerals but Fe₂O₃ content was the highest among all compositions detected. The contents of Pb and Zn in MRM were 10.9% and 8.1%, respectively. The values of both metals were extremely high and exceeded the permissible limit in soil, 600 mg/kg for Pb and 1500 mg/kg for Zn [40]. However, this does not guarantee that MRM can release significant amounts of Pb and Zn since their mobility also depends significantly on the chemical speciation. Sequential extraction was then performed, and the result showed that around 40% of the total contents of Pb and Zn were in mobile fractions (exchangeable, carbonates, and oxidizable) under surface environments (Figure 2). This confirms that MRM could be a potential source contaminating the surrounding environment with Pb and Zn.

Table 2. Mineralogical composition of solid samples.

|        | MRM | RD  | CD  | MO  |
|--------|-----|-----|-----|-----|
| Quartz | ++  | -   | -   | -   |
| Gypsum | +   | -   | -   | -   |
| Anglesite | +++ | -   | -   | -   |
| Zinkosite | ++ | -   | -   | -   |
| Hematite | +   | -   | -   | -   |
| Goethite | +   | -   | -   | -   |
| Dolomite | -   | +++ | +++ | -   |
| Calcite | -   | -   | +   | -   |
| Magnesia | -   | -   | +   | +++ |

+++: Strong; ++: Moderate; +: Weak; -: None.

Table 3. Chemical composition of solid samples (the unit is in wt%).

|        | MRM | RD  | CD  | MO  |
|--------|-----|-----|-----|-----|
| SiO₂   | 20.9| 0.39| 0.37| <0.01|
| TiO₂   | 0.35| <0.01| <0.01| <0.01|
| Al₂O₃  | 1.91| 0.24| 0.11| <0.01|
| Fe₂O₃  | 45.8| 0.52| 0.68| <0.01|
| MnO    | 1.59| 0.46| 0.43| <0.01|
| MgO    | <0.01| 36.9| 33.3| 100  |
| CaO    | 4.64| 60.7| 62.4| <0.01|
| Na₂O   | <0.01| <0.01| <0.01| <0.01|
| K₂O    | <0.01| 0.15| 0.16| <0.01|
| P₂O₅   | <0.01| 0.14| <0.01| <0.01|
| SO₂    | 2.71| 0.15| 0.2  | <0.01|
| Pb     | 10.9| <0.01| <0.01| <0.01|
| Zn     | 8.1 | <0.01| <0.01| <0.01|

The most dominant mineral found in RD was dolomite (CaMg(CO₃)₂). Magnesium (Mg) and calcium (Ca) oxides contents accounted for more than 95% with a molar ratio of Ca to Mg of 1.2.
This indicates that RD adequately consisted of pure dolomite. Burning RD at 700 °C for 2 h generated the new type of immobilizer, CD. Calcite (CaCO₃) and magnesia (MgO) were detected in addition to dolomite. With almost the same molar ratio of Ca to Mg in CD compared with that in RD, it clearly indicates that the calcination process transformed dolomite into calcite and magnesia. MO composed only of magnesia with 100% MgO content, which shows pure magnesia.

Figure 2. Solid-phase Pb and Zn speciation of mining residual materials (MRM); (□) exchangeable, (□) carbonates, (□) sulfide/organic matter, (□) Fe-Mn oxides, and (□) residual.

3.2. Leaching Characteristic of MRM

Table 4 shows the leaching characteristic of MRM. The experimental values were reproducible with high precision since the standard deviations of all parameters were quite small. Four heavy metals, Pb, Zn, cadmium (Cd), and copper (Cu), were leached at the concentrations falling within the instrument detection limits of ICP-AES and -MS. Among these heavy metals, the leaching concentrations of only Pb (2.1 mg/L) and Zn (365 mg/L) exceeded the effluent standard in Zambia (0.5 mg/L for Pb and 1 mg/L for Zn) [41]. Therefore, this study focused only on the mobility of these two metals. PHREEQC simulation on the saturation indices of all possible Pb- and Zn-minerals showed that with the given conditions and components in MRM leachate, only saturation index of anglesite fell within a common error interval used to indicate saturation equilibrium (±0.2) (Table 5). The result indicates that the low solubility of anglesite restricted the dissolved concentration of Pb, while no restriction by means of precipitation was observed on the leaching of Zn. This can explain why 4.51% of the total Zn content was leached from MRM, while only 0.02% was observed in the case of Pb leaching.

Table 4. Leaching characteristic of MRM (n = 5).

|          | MRM      | Reg. Value * (mg/L) |
|----------|----------|---------------------|
| pH       | 5.26 ± 0.04 | -                   |
| ORP (mv) | 300 ± 36  | -                   |
| EC (mS/cm)| 2.7 ± 0.09  | -                   |
| Pb (mg/L)| 2.1 ± 0.008 | 0.5                 |
| Zn (mg/L)| 365 ± 18  | 1                   |
| Cd (mg/L)| 0.21 ± 0.009  | 0.5                 |
| Cu (mg/L)| 0.08 ± 0.01  | 1.5                 |
| Ca²⁺ (mg/L)| 547 ± 48.2 | -                   |
| Mg²⁺ (mg/L)| 27.7 ± 3.1 | -                   |
| SO₄²⁻ (mg/L)| 1907 ± 36.8 | -                   |
| Si (mg/L)| 13 ± 0.5 | -                   |

* Regulated value in mg/L specified by the Environment Management Act (2013) [40].
Table 5. Calculated saturation indices of possible Pb- and Zn-minerals in MRM leachate.

| Mineral     | Saturation Index |
|-------------|------------------|
| Anglesite   | −0.2             |
| Cerussite   | −2.25            |
| Pb(OH)\(_2\) | −3.4             |
| Smithsonite | −2.45            |
| Willemite   | −3.62            |
| Zn(OH)\(_2\) | −3.82           |
| Hydrocerussite | −8.41         |
| Hydrozincite | −3.08         |

Calcium ion (Ca\(^{2+}\)) and sulfate ion (SO\(_{4}^{2−}\)) were the major ions in the leachate, accounting for more than 85% of the total dissolved ions. These ions were likely to be enriched by the dissolution of soluble phase minerals, such as gypsum and zinkosite [42,43]. However, the dissolution of these two minerals might not only be the sources of SO\(_{4}^{2−}\) since the molar ratio of Ca\(^{2+}\) and Zn to SO\(_{4}^{2−}\) was lower than one. The sulfide fractions of both metals in MRM (Figure 2), together with the slightly acidic pH (5.2) and positive ORP (300 mV) of the leachate, suggest that the oxidation of sulfide minerals (e.g., pyrite, galena, sphalerite) and dissolution of iron-sulfate salts (e.g., melanterite, coquimbite) also occurred and attributed to the enrichment of SO\(_{4}^{2−}\), even though they were not detected by XRD. This could also partly contribute to the enrichment of Pb and Zn in the leachate.

3.3. Potential of Immobilizers

3.3.1. Effects of Addition of Immobilizers on pH and Coexisting Ions

Changes in the pH of the leachate as a function of the amounts of addition of RD, CD, and MO are shown in Figure 3. The pH increased from 5.2 to 6.7, 8.2, and 9.8 with increasing RD, CD, and MO addition from 0 to 10%. When the same amount of immobilizer was added, the performance of the immobilizers to increase the leachate pH followed the order: MO > CD > RD. The results clearly showed the improvement of the alkaline property of CD over RD. The variation in pH could mainly be attributed to the liming effect(s) of dolomite (Equation (1)) in RD treatments, of dolomite (Equation (1)), of calcite (Equation (2)), and of magnesia (Equation (3)) in CD treatments, and of magnesia (Equation (3)) in MO treatments [44–47].

\[
\text{CaMg(CO}_3\text{)}_2 + 2 \text{H}^+ \rightleftharpoons \text{Ca}^{2+} + \text{Mg}^{2+} + 2\text{HCO}_3^−, \quad (1)
\]

\[
\text{CaCO}_3 + \text{H}^+ \rightleftharpoons \text{Ca}^{2+} + \text{HCO}_3^−, \quad (2)
\]

\[
\text{MgO} + \text{H}_2\text{O} \rightleftharpoons \text{Mg(OH)}_2 \rightleftharpoons \text{Mg}^{2+} + 2\text{OH}^−. \quad (3)
\]

Figure 4a–c illustrates the leaching concentrations of major coexisting ions, Ca\(^{2+}\), magnesium (Mg\(^{2+}\)), and SO\(_{4}^{2−}\), as a function of the amount of immobilizer added. The dissolved concentrations of Ca\(^{2+}\) and Mg\(^{2+}\) in RD treatments were higher than those in MRM and increased with the increasing addition of RD, indicating the simultaneous leaching of Ca\(^{2+}\) and Mg\(^{2+}\) from the dissolution of dolomite (Equation (1)). In CD treatments, Ca\(^{2+}\) and Mg\(^{2+}\) were also leached at higher concentrations than those in MRM leachate. At the same amount of CD and RD addition, the leaching concentration of Mg\(^{2+}\) in CD treatment was higher than that in RD treatment, while almost the same dissolved concentration of Ca\(^{2+}\) was observed in both treatments. This, together with the result that CD contained less dolomite and more calcite compared to those in RD, suggest the occurrence of hydration of magnesia (Equation (3)) in addition to the dissolution of carbonate minerals (Equations (1) and (2)) in CD treatments. Moreover, the difference in the leaching concentration of Mg\(^{2+}\) between CD and RD treatments became more significant as the addition of immobilizers increased, which indicates that as pH increased, the hydration of magnesia (Equation (3)) played a more important role in controlling
pH. In MO treatments, the concentration of Mg$^{2+}$ increased with higher addition of MO, while almost no change in the concentration of Ca$^{2+}$ from that in MRM was observed. This result suggests the occurrence of hydration of magnesia (Equation (3)) in MO treatments.

\[
\text{MgO + H}_2\text{O} \rightleftharpoons \text{Mg(OH)}_2 \rightleftharpoons \text{Mg}^{2+} + 2\text{OH}^-. (3)
\]

Figure 3. pH of leachate upon addition of immobilizers.

Figure 4a–c illustrates the leaching concentrations of major coexisting ions, Ca$^{2+}$, magnesium (Equation (1)), and sulphate concentrations upon addition of immobilizers: (a) Ca$^{2+}$, (b) Mg$^{2+}$, and (c) SO$_4^{2-}$.

At the same amount of immobilizer added, the leaching concentration of SO$_4^{2-}$ was the highest when treating MRM with MO, followed by CD and RD (Figure 4c). In consideration of the trace amount of SO$_3$ content in all immobilizers, the results suggest that MRM should be the source of SO$_4^{2-}$, and the leaching of SO$_4^{2-}$ might be caused by the change in the parameter(s) of the leachate, triggered
by the addition of immobilizer. Figure 5 illustrates the correlation between the leaching concentration of \( \text{SO}_4^{2-} \) vs. pH. The \( \text{SO}_4^{2-} \) concentration exhibited a strong positive correlation with pH (correlation coefficient \( r = 0.92, p < 0.01 \)), suggesting that \( \text{SO}_4^{2-} \) level could mainly be influenced by pH due possibly to the following mechanisms: (1) desorption, (2) production by oxidation of sulfide minerals, (3) common-ion of between calcite, dolomite, and gypsum, and (4) dissolution of anglesite. The pH increase led to a higher negative surface potential of MRM, thereby decreasing the affinity of \( \text{SO}_4^{2-} \) toward the surface of MRM. However, Tabatabai (1987) [48] reported that since the adsorption of \( \text{SO}_4^{2-} \) was only flavored under strongly acidic conditions, the amount of adsorbed \( \text{SO}_4^{2-} \) became almost negligible under weakly acidic pH. This means that the desorption might not be a viable explanation in this study since all leachates’ pH ranged from weakly acidic (5.2) to moderately alkaline (9.8). Therefore, the fact that the oxidation rate of sulfide minerals, such as pyrite and galena, increases with pH becomes the potential reason contributing to the higher leaching concentration of \( \text{SO}_4^{2-} \) [49,50]. However, the enrichment of \( \text{SO}_4^{2-} \) could be restricted by the solubility of gypsum because the saturation index of gypsum was within the equilibrium condition range of \( \pm 0.2 \) in all leachates. This could explain why the leaching concentration of \( \text{SO}_4^{2-} \) slightly increased with pH at lower pH region where the dissolution of dolomite containing in RD and calcite and dolomite containing in and CD tended to control the pH (Equations (1) and (2)). In other words, \( \text{Ca}^{2+} \) produced from the dissolution of calcite and dolomite precipitated with \( \text{SO}_4^{2-} \) to form gypsum, thereby restricting \( \text{SO}_4^{2-} \) concentration. Meanwhile, as pH became more alkaline, the concentration of \( \text{SO}_4^{2-} \) increased rapidly. This could be attributed to the less contribution of calcite and dolomite dissolution (Equations (1) and (2)) to control the leachate pH in the case of CD addition, conjointly in MO treatments, only hydration of magnesia (no production of \( \text{Ca}^{2+} \)) (Equation (3)) was found to control the pH of the leachates. The pH-dependent solubility of anglesite could also be attributed to the rapid increase of \( \text{SO}_4^{2-} \) concentration under alkaline conditions since anglesite was originally contained in MRM and is unstable under alkaline pH [51].

![Figure 5. Leaching concentration of \( \text{SO}_4^{2-} \) vs. pH.](image)

3.3.2. Effects of the Addition of Immobilizers on Mobility of Heavy Metals

In this study, the performance of immobilizers was evaluated based on the solubility of Pb and Zn. Figure 6a,b shows the changes in Pb and Zn concentrations as a function of the amount of addition of immobilizers. In general, the leaching concentrations of Pb and Zn exponentially decreased with an increase in the dose of immobilizers. Figure 7a,b illustrates the correlation between leaching concentrations of Pb and Zn vs. pH. The mobility of heavy metals was strongly influenced by pH, indicated by significant negative correlations of Pb-pH \( r = -0.92 \) and Zn-pH \( r = -0.87 \) at the 0.01 significance level (2-tailed). Coupled with the results of the leaching concentration of \( \text{SO}_4^{2-} \) and the characteristic of immobilizers, as well as the leaching condition used in the current study, the major modes of Pb and Zn attenuation could be either one or more of the following...
mechanisms: precipitations of metal-sulfate, -carbonate, and/or -hydroxide, co-precipitation of metal with iron-(oxy)hydroxides, and metal ion adsorption to immobilizer. The formation of low soluble anglesite under acidic conditions was expected since the leaching concentration of $SO_4^{2-}$ was high and increased with pH (Figure 5) \cite{52,53}. The carbonate property of RD and CD might result in the precipitation of cerussite ($PbCO_3$), hydrocerussite ($Pb_3(CO_3)_2(OH)_2$), smithsonite ($ZnCO_3$), and hydrozincite ($Zn_3(CO_3)_2(OH)_2$) \cite{35,54-56}. The carbonate precipitations of Pb and Zn are also expected to occur in MO treatments since the experiments were done under atmospheric conditions in which carbon dioxide ($CO_2$) in the atmosphere was freely dissolved \cite{57}. However, the simulation results by PHREEQC showed that except hydrozincite in 3% addition of MO treatment, the precipitations of anglesite, cerussite, hydrocerussite, smithsonite, and hydrozincite were thermodynamically unfavorable (saturation index $< -0.2$) regardless of the type and amount of immobilizer added (Table 6). Therefore, the possible immobilization mechanisms of Pb and Zn in all types of immobilizers could be narrowed down to the hydroxide precipitation, adsorption, and co-precipitation.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure6.png}
\caption{Leaching concentrations of heavy metals upon addition of immobilizers: (a) Pb and (b) Zn (dashed lines represent the effluent standards of Pb and Zn in Zambia).}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure7.png}
\caption{Leaching concentrations of heavy metals vs. pH: (a) Pb vs. pH and (b) Zn vs. pH (dash lines represent the effluent standards of Pb and Zn in Zambia).}
\end{figure}

To verify the dominant mechanism(s), pH-dependent solubility diagrams of Pb- and Zn-hydroxides were plotted (Figure 8a,b). Points in the figures represent the relationship between the logarithmic activity of divalent heavy metal and pH in each batch test. The solid lines demonstrate the solubility of heavy metal hydroxides. Therefore, any point located on or close to the line implies the hydroxide precipitation-controlled sequestration process.
This means that at pH from weakly acid to neutral, hydroxide precipitation was not the main mechanism of pH. Theoretically, as pH increases under acidic region, more of free Pb(II) ion tends to complex with OH⁻ and CO₃²⁻, generating larger ion with lower charge (Pb(OH)⁺ and PbCO₃), which lowers the affinity of Pb to the surface of the potential adsorbents and inhibits the co-precipitation [69–71].

In the case of RD addition, a discrepancy from the equilibrium line for Pb and Zn was observed. This means that at pH from weekly acid to neutral, hydroxide precipitation was not the main mechanism controlling the mobility of Pb and Zn. Therefore, Pb and Zn were suspected to be immobilized by sorption and co-precipitation with iron-(oxy)hydroxides. The sorption was likely to occur since dolomite, major mineral in the immobilizer, can adsorb Pb and Zn [58,59]. Besides, adding more of this immobilizer induced the leachate pH increase. This could change the surface charge of goethite, hematite, and iron-(hydr)oxide compounds in MRM to more negative, thereby increasing their adsorption ability against cationic divalent Pb and Zn [58–63]. During the neutralization process under ambient conditions, iron-(oxy)hydroxides precipitate from the oxidative dissolution of pyrite and dissolution of iron-bearing salts [63–66]. The precipitations of iron-(oxy)hydroxides have been reported by many studies to induce co-precipitation of divalent metals, including Pb(II) and Zn(II) [50,67,68], and thus the co-precipitation of Pb and Zn with iron-(oxy)hydroxides was also expected in RD treatments. From the above explanations, adding more RD should reduce the leaching concentration of both metals. However, leaching concentration of Pb increased from 0.98 mg/L to 1.4 mg/L when RD rose from 5% to 10% (Figure 6a). This could be attributed to the stability of Pb(II) species as a function of pH. Theoretically, as pH increases under acidic region, more of free Pb(II) ion tends to complex with OH⁻ and CO₃²⁻, generating larger ion with lower charge (Pb(OH)⁺ and PbCO₃), which lowers the affinity of Pb to the surface of the potential adsorbents and inhibits the co-precipitation [69–71].
On the other hand, adding CD and MO made most of the logarithmic leaching activities of Pb\(^{2+}\) and Zn\(^{2+}\) to approach their solubility product lines. This means that hydroxide precipitation is the dominant mechanism of attenuating Pb and Zn. Regardless of the type of immobilizer, at low pH, the logarithmic activities of both metals were slightly lower than their equilibrium lines and then tended to stay on or be slightly higher than the lines afterward. This probably indicates that at low pH, adsorption and co-precipitation with iron-(oxy)hydroxides also occurred in addition to the precipitation, but as pH got higher, they diminished. There are two probable explanations for this phenomenon as follows: (1) competition with strong competing ion (Mg\(^{2+}\)) and (2) change in specification of the dissolved metals. The pH alteration mechanisms of CD and MO appeared to generate Mg\(^{2+}\) as a by-product (Equations (1) and (3)). Because of this, the concentration of Mg\(^{2+}\) significantly increased with pH with a correlation coefficient of 0.97, \(p < 0.01\) (Figure 9). Therefore, as pH increased, high concentration of Mg\(^{2+}\) could compete for Pb and Zn for adsorption sites and for co-precipitation with iron-(oxy)hydroxides, attributing to the less contribution of the adsorption and co-precipitation on the immobilization process. Increasing pH could also result in the redistributions of Pb(II) and Zn(II) species. The fraction of free Pb(II) and Zn(II) reduces as pH increases since they are thermodynamically preferable to be hydrolyzed forming \(-\text{(OH)}^+\), \(-\text{(OH)}_2\), and \(-\text{(OH)}_3^-\) \([69–72]\). Moreover, since the systems contained high dissolved carbonate, the formation of carbonate complexes of Pb(II) and Zn(II) was also expected \([69,73]\). Once these complexes are formed, their abilities to get adsorbed and co-precipitated are inhibited by the larger size and lower positive potential they become.

![Figure 9. Leaching concentration of Mg\(^{2+}\) vs. pH.](image-url)

### 3.3.3. Performance of Immobilizers

When the same amount of immobilizer was added, the dissolved concentrations of Pb and Zn were the highest in RD treatment, second highest in CD treatment, and the lowest in MO treatment (Figure 6a,b). As previously mentioned, adsorption, co-precipitation, and hydroxide precipitation were the major sink of Pb and Zn, and their mobilities depended strongly on pH. Because of the carbonate property of dolomite, RD could not raise the pH of MRM leachate to the value favoring the precipitations of Pb- and Zn-hydroxides. Therefore, RD treatments remediated Pb and Zn by adsorption and co-precipitation in which it was insufficient to reduce the leaching concentrations of Pb and Zn down below their regulated values. On the other hand, magnesia in CD and MO played a significant role in increasing the leachate pH of MRM into the alkaline region. Lead and Zn were then mainly immobilized by precipitation as hydroxides. Thus, metal concentrations as high as 368 mg/L for Zn and 2.1 mg/L for Pb released from MRM were reduced to the values below their regulated concentrations. Efficiency-wise, MO was the most effective immobilizer in immobilizing Pb and Zn since it contained the highest MgO content. However, CD could be the immobilizer of choice since it can be produced from the naturally abundant material in Zambia (RD), and its performance was almost the same as that of MO.
4. Conclusions

The leachate of MRM was slightly acidic (pH 5.2) and contained high concentrations of Pb (2.1 mg/L) and Zn (365 mg/L), exceeding those of Zambian regulation. When immobilizers were introduced, the leachate pH increased, and the leaching concentrations of Pb and Zn decreased. Lead and Zn immobilized by RD were interpreted by the adsorption and co-precipitation mechanisms. On the contrary, chemical immobilization using CD and MO suppressed Pb and Zn leaching mainly by the hydroxide precipitation. Of the immobilizers investigated, only CD and MO decreased the dissolved Pb and Zn concentrations to below their regulated values, in which MO had a higher performance than CD. The results show that heat treatment on RD to produce CD drastically improved the immobilizing performance of Pb and Zn. Even though MO provided the highest efficiency, Pb and Zn could also be effectively immobilized by giving an adequate amount of CD. Therefore, by considering the availability of CD in the local area, CD could be the most promising chemical agent to be implemented in Zambia.

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