Heavy Metal Pollution Assessment by Partial Geochemical Extraction Technique

Samuel Estifanos
Department of Earth Science, CNCS, P.O. Box 231, Mekelle University, Mekelle, Ethiopia (sambersih@yahoo.com)

ABSTRACT

Mn and Fe oxides are powerful absorbents of heavy metal cations in soils and stream sediments therefore considered in this environmental geochemical investigation. Steam sediment samples were collected from Au-Ag and Pb-Zn-Cu mineralized areas of the Rodalquilar old gold mine, located in the southeastern part of Spain. Total and partial chemical analyses were done, using aqua regia and hydroxylamine-hydrochloric acid solutions, respectively to examine preferential concentration of Zn, Pb and Cu in amorphous and ore mineral phases. The results show scavenging association of Pb, Zn and Cu with Mn that is shown by their concentrations in amorphous phases as detected by partial extractions.

Key words: Partial and total extraction, Amorphous phase, Scavenging, Heavy metal pollutants.

1. INTRODUCTION

Normally heavy metals are not harmful to our environment because they are only present in very small amounts. The environmental pollution concern rises when they show up in higher concentration due to natural mineralizing processes and/or human activities. The heavy metals threatening the ecosystem include mercury, arsenic, copper, barium, cadmium, antimony, chromium, lead and zinc. The effects caused by heavy metals vary in that dissolved copper can affect lower trophic levels such as phytoplankton while lead, cadmium and mercury become accumulated through the food-chain, posing a toxic risk to species higher in the food chain, and eventually to humans. The degree of fixation of trace elements by Mn and Fe oxides ranges from adsorption at the surface, through co-precipitation, to relatively strong binding within the oxide structure (Hall, 1997).

Dissolution techniques of samples prior to chemical analysis can be classified as partial extraction and sequential (selective) multiple extraction (Fig. 1). The basic assumption of sequential extraction is that the reagents used are able to dissolve one mineral phase, selectively without solubilizing the others. Partial extraction involves dissolution of all mineral phases which requires milder reagents. The use of partial extraction techniques in the enhancement of soil and stream sediments geochemical anomalies has been applied with success in temperate and Mediterranean climates for Cu, Zn, Pb and U (Cardoso and Ferreira, 1998). Partial and sequential (selective)...
ex extractions have been adopted in studies related to both exploration geochemistry and environmental geochemistry (Cornu et al., 2002; Kelley et al., 2003; Rodrigues and Formoso, 2005; Nada et al., 2011). The requirements of sample decomposition in environmental geochemistry are less exacting than those in exploration works because minerals that require rigorous conditions to be solubilized are unlikely to have environmental impact.

![Diagram of leachant type (reagent for total, partial and selective leaching) and component leached in a sample](image)

**Figure 1.** Relationship of leachant type (reagent for total, partial and selective leaching) and component leached in a sample (Modified after Cohen et al., 2007).

The degree with which the potentially toxic metals impact the ecosystem depends on their solubility in solutions of varying pH conditions. This is in turn related to the phases of occurrence of the metals. A toxic heavy metal is said to have the highest impact on the ecosystem when it exists in a mineral form that dissolves easily with water; otherwise, it has the lowest impact when the host mineral needs rigorous acidic conditions to dissolve, a rare case in nature such as acid mine drainage (AMD). Thus, based on the analytical results, the scavenging degree of the three potentially toxic metals (Pb, Cu and Zn) by Mn and Fe on the study area is assessed.

Manganese is not found as a free metal in nature, however its minerals consisting of oxides, silicates and carbonates are the most common (Web Elements, 2005). The iron and steel industry and acid mine drainage release a large portion of manganese found in the environment. Manganese is very similar to iron in its chemical behavior and so is frequently found in association with iron.
Mn is essential micronutrient with certain limit. The maximum contamination level (MCL) drinking water standard is 50 ppb (Table 1).

Cu is essential nutrient but excesses can be toxic and it is essentially immobile under basic pH conditions. Zn is essentially immobile under basic pH>5 and remains sorbed to clay and free oxides in upper layer of the soil. lead is non-essential and toxic heavy metal (Siegel, 2002).

Table 1. Summary of the heavy metals (HM) abundances in the continental crust (UCCA), maximum contamination levels (MCL) and lower limits of detection (LDL) by ICP-AES* technique (after Siegel, 2002)

<table>
<thead>
<tr>
<th>S. No</th>
<th>HM</th>
<th>UCCA (ppm except Fe (%)</th>
<th>MCL in water (ppb)</th>
<th>LDL for ICP-AES (ppb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Pb</td>
<td>20</td>
<td>1.5</td>
<td>10</td>
</tr>
<tr>
<td>5</td>
<td>Zn</td>
<td>71</td>
<td>5000</td>
<td>4</td>
</tr>
<tr>
<td>6</td>
<td>Cu</td>
<td>25</td>
<td>1000</td>
<td>4</td>
</tr>
<tr>
<td>7</td>
<td>Fe</td>
<td>3.5</td>
<td>200</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Mn</td>
<td>600</td>
<td>50</td>
<td></td>
</tr>
</tbody>
</table>

*ICP-AES: Inductively Coupled Plasma- Atomic Emission Spectrometer.

Several workers have developed a method for the selective dissolution of manganese oxides from soils and sediments. This method uses a solution of 0.1M hydroxylamine (NH₄OH) and hydrochloride in 0.01M HNO₃ at pH=2 (Thompson, 1983). The action of the hydroxylamine hydrochloride solution under the above conditions is specific to manganese oxides (Fig. 1). Crystalline iron oxides such as goethite, hematite and magnetite do not dissolve in this solution (Cardoso and Ferreira, 1998). Selective extraction of Mn oxides and Fe oxides is usually performed by using reagents, hydroxylamine hydrochloride being the popular choice (Hall, 1997).

The aim of this paper is to examine the significance of selective and total extraction techniques in dissolving Mn and Fe mineral phases and assess heavy metals pollution in the mineral phases. These techniques are used to investigate how trace elements are distributed between the phases. Moreover, these extractions have been used in the past for geochemical exploration because it has been perceived that elements associated with mineralization (especially those hosted by sulfides) are more likely to be released during weathering than those in barren rocks (hosted by silicates), more widely dispersed, and more probably held by secondary minerals and hence likely to be preferentially released by extraction solutions. In contrast, some of the new extraction techniques
currently used for geochemical exploration are partial rather than selective digests, i.e. they extract part of a phase or phases, rather than a selected mineral (Gray, 1996).

2. MATERIALS AND METHODS

2.1. Study area

The Study area is located in the South-eastern part of Spain (Fig. 2A). The approximate total catchment area considered is about 18 Km$^2$. Extensive mining workings and ore processing wastes have caused exposure of sulfide rocks to the atmosphere and loading of waste sediments to the stream systems eventually reaching the Sea.

A number of studies have previously been conducted in the study area with aspects ranging from pure exploration geological to environmental pollution studies as well as remote sensing applications (Arribas et al., 1995; Wray, 1998; Ferrier, 1999; Chica-Olmo et al., 2001; Estifanos, 2006). The gold-alunite deposits are located in the central part of the volcanic field within the Rodalquilar caldera complex where four volcanic complexes are identified. The volcanogenic complexes are overlain by younger fossilized calcareous limestone (Arribas et al., 1995) (Fig. 2B).

Figure 2. Showing (A) location and regional geology and (B) local geology (After Arribas et al., 1995).
2.2. Sampling and geochemical analysis

Stream sediment samples were collected in June, 2005, at the junctions of tributaries in the lower and higher orders of the main stream beds of the Rodalquilar area (Fig. 3). At each sample point five to seven central scoops of the dry sediments were collected from the stream bed using a stainless steel and was sieved with a <1mm stainless steel sieve. All sampling equipments were thoroughly cleaned between each sampling spot to avoid sample-to-sample contamination. Blank and standard samples as well as 5 duplicate sample analyses were made to check the quality of the analysis of ICP- AES. The results of these controlling samples indicate that the quality of the analysis was very good.

![Figure 3. Spatial distribution of stream sediment samples in the study area (coordinates in UTM).](image)

The samples were subjected to geochemical analysis in International Institute for Geo-information Science and Earth Observation (ITC) geochemical laboratory where ICP-AES analytical instrument was used. A number of elements were extracted from three solutions: water, hydroxylamine (partial extraction) and aqua regia (total extraction) following the procedures described in table 2.

10gm of each sample was dissolved in water and shaken for some time. ICP-AES could detect cations of Fe, Mn, Pb, Zn, Cu, Ca, Mg and Na being dissolved for concentration above the detection limit. Whereas the major ions in water can be determined directly, the background levels
of most heavy metals, including Cu, Pb, Zn and Cd are too low for direct determination even by ICP (Thompson, 1983).

For partial extraction, 1gm of each sample was digested in 20 ml solution of 0.25M hydroxylamine and 0.1mol hydrochloric acid before Mn, Cu, and Zn, Pb were analyzed by ICP-AES. The partial geochemical extraction was aimed to extract the pollutants that might have been attached to amorphous phases of Mn and Fe minerals.

In total extraction, a solution of aqua regia (3:1 ratio of hydrochloric and nitric acids), is used to dissolve a wide range of mineral phases releasing cations of heavy metals from crystalline phases. Aqua regia has similar extraction characteristics to nitric acid – perchloric acid but the former is more effective in dissolving some sulfides notably pyrite (Thompson, 1983). A 1:4 mixture of perchloric and nitric acid if slowly evaporated to dryness with 0.25 gm samples of rock, soil or sediment, provides powerful decomposition of the greater part of any heavy metals present (Thompson, 1983). The elements to be determined satisfactorily are Cu, Pb, Zn, Cd Mn, Fe, Co, Ni, Mo and P (Thompson, 1983).

Table 2. Geochemical analytical procedure for the three leachates.

<table>
<thead>
<tr>
<th>Water extraction (W)</th>
<th>Partial extraction (HA)</th>
<th>Total extraction (AR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 gm, Adding 25 ml demineralized water</td>
<td>1 gm, 20 ml of 0.25M hydroxylamine and 0.1M HCl solution</td>
<td>1 gm, 5 ml aqua regia and 5 ml demineralized water</td>
</tr>
<tr>
<td>Shaking for 2 hours</td>
<td>Shaking for two hrs in 60°C water</td>
<td>Shaking for 4 hrs in 70°C water</td>
</tr>
<tr>
<td>pH measurement</td>
<td>Decantation</td>
<td>Decantation</td>
</tr>
<tr>
<td>Centrifuging 4000 rpm</td>
<td>Centrifugal 4000 rpm</td>
<td></td>
</tr>
<tr>
<td>Settlement and decantation</td>
<td>Settlement and decantation</td>
<td></td>
</tr>
<tr>
<td>Filtering over a funnel</td>
<td>Filtering over a funnel</td>
<td>Filtering over a funnel</td>
</tr>
<tr>
<td>Storage in a 20 ml Pyrex test tube with stopper</td>
<td>Storage in a 20 ml pyrex test tube with stopper</td>
<td>Diluting for 100 ml de-ionized water</td>
</tr>
<tr>
<td>ICP AES analyses for Fe, Mn, Zn, Pb, Cu</td>
<td>ICP AES analyses for Fe, Mn, Zn, Pb, Cu, Pb</td>
<td>ICP AES analyses for Fe, Mn, Sb, Cu, Pb, Zn</td>
</tr>
</tbody>
</table>
2.3. Statistical analysis

The geochemical analysis dataset from both the total and partial extraction were treated in such a way that the cations concentration from the crystalline and amorphous phases of the sample materials is separated. This is done simply by subtracting the cations concentration in the partial extraction from that of total extraction to separate the cations that were in crystalline phases and considering the difference between partial and water extraction for cations in amorphous phases.

3. RESULTS AND DISCUSSION

3.1. Distribution of Mn, Zn, Pb and Cu in the three phases

The partial geochemical extraction conducted using 20 ml of 0.25M hydroxylamine and 0.1M HCl solution was aimed to extract the contaminants that might have been attached to amorphous phases of minerals. Unlike, sequential extraction which can selectively dissolve particular phase, the partial extraction includes the water soluble mineral phases also. To enhance the heavy metals concentration attached to amorphous and crystalline/primary rocks, the difference between partial and water extractable values (HA - W) as well as difference between total and partial extraction values (AR - HA) were considered respectively.

Among zinc, copper and lead detected in water, partial and total extractions (Table 3), lead exceeds the MCL of 1.5ppb even in the water extractable concentration level (Fig. 4). It is widely accepted that the water extractable metal in sediments is more mobile, highly toxic and the most bio-available fraction (Lalita and Srilert, 2011). Moreover, its average concentration from partial extraction accounts about 50% of the total extraction. The average concentration of the total extraction values of these heavy metals exceed their crustal abundances (Fig. 5).

Table 3. Statistical summary for the geochemical parameters in three extraction datasets.

<table>
<thead>
<tr>
<th>Water extractable</th>
<th>Descriptive Statistics</th>
<th>Partial extraction</th>
<th>Descriptive Statistics</th>
<th>Total extraction</th>
<th>Descriptive Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe ppm</td>
<td>N 50</td>
<td>Min. 0.03</td>
<td>Max. 732</td>
<td>Mean 294.26</td>
<td>Std. Dev. 104.39</td>
</tr>
<tr>
<td>Mn ppm</td>
<td>N 50</td>
<td>Min. 0.01</td>
<td>Max. 0.29</td>
<td>Mean 0.07</td>
<td>Std. Dev. 0.07</td>
</tr>
<tr>
<td>Cu ppm</td>
<td>N 50</td>
<td>Min. 0.07</td>
<td>Max. 1.92</td>
<td>Mean 0.40</td>
<td>Std. Dev. 0.22</td>
</tr>
<tr>
<td>Pb ppm</td>
<td>N 50</td>
<td>Min. 0.01</td>
<td>Max. 0.28</td>
<td>Mean 0.18</td>
<td>Std. Dev. 0.18</td>
</tr>
<tr>
<td>Zn ppm</td>
<td>N 50</td>
<td>Min. 0.01</td>
<td>Max. 1.92</td>
<td>Mean 0.28</td>
<td>Std. Dev. 0.28</td>
</tr>
</tbody>
</table>

Mn and Fe are considered in this paper to verify their scavenging property towards the heavy metals. Mn exceeds the MCL set to be 50ppb in its water extractable phases while MCL of 0.2ppm...
is set for Fe which is claimed to be potentially toxic and thus it is in polluting level in at least the water extractable phase in the study (Table 3).

![Histogram showing the elevated concentration of heavy metals (Pb, Zn and Cu) relative to environmentally acceptable level.](image)

**Figure 4.** Histogram showing the elevated concentration of heavy metals (Pb, Zn and Cu) relative to environmentally acceptable level (*the values are plotted on log scale to enhance the low values of the environmentally accepted limits for relative comparison purpose)*

![Histogram showing comparison of average concentration of pollutants with upper crustal abundance.](image)

**Figure 5.** Histogram showing comparison of average concentration of pollutants with upper crustal abundance

The following pie diagrams illustrate the approximate and relative proportion of the contaminants in the three broad phases (Fig. 6). The pie diagrams show that Fe-Zn-Cu-Pb-Mn cations are attached to amorphous phase (HA-W) in increasing order. The higher proportion of Mn in the amorphous phase indicates the presence of Mn minerals predominantly as either amorphous-minerals or coating material since the partial extraction is known to be efficient in dissolving amorphous Mn-related minerals. A number of studies strengthen that Mn minerals have scavenging affinity towards a number of heavy metals (Hall, 1997; Kuang-Chung et al., 2001; O’Reilly, 2002; Wenfeng et al., 2005).
Manganese (III/IV) oxides/hydroxides are common in soils and sediments as particles and coatings on mineral surfaces (Russell, 1998). Manganese is known to be the most important scavenger for many inorganic cations such as Cu, Pb, Zn, Co and Ni among others. This is because it exists in several oxidation states forming non-stoichometric oxides with different valences. Its higher valence oxides exist in several crystalline forms that co-precipitate and solid solution with Fe oxides. It can have surface areas of several hundred m²/g and larger CEC (cation exchange capacity) than some clay minerals, though they are much less abundant than iron oxides (Hall, 1997; O’Reilly, 2002). However, the metal’s mobilities are expected to be affected also by the prevailing natural pH conditions of the area.

3.2. Spatial Patterns of the Pollutants

The spatial association of the heavy metals with manganese was assessed using plots. The partial extraction values were subtracted from the total extraction concentration values to determine the portion contained in the crystalline phases of each trace element: Mn, Pb, Cu and Zn. In the spatial assessment, the proportion of the amorphous phases relative to the crystalline phases of the trace elements is considered. The plots in figure 7 show the spatial variability of both manganese and the other heavy metals and evaluate the scavenging relationships in the amorphous phases.

Figure 6. Pie diagrams showing relative proportions of Pb, Zn, Cu, Fe and Mn Cations in the three phases (W: water extractable, HA-W: amorphous and AR-HA: crystalline phases).
Figure 7. Plots of the partial extraction values in proportion to total extraction values (A. Pb vs Mn; B. Cu vs Mn and C. Zn vs Mn).

Figure 8. Spatial pattern of pH values in the study area.
In the plots, it is evident that Mn has spatial association with lead (Fig. 7A), with copper (Fig. 7B) and with zinc (Fig. 7C) in decreasing order. The proportion of the trace metals in their amorphous phases is significantly higher in the downstream part and show consistent correlation with Mn amorphous proportions. This indicates the scavenging effect of Mn on the heavy metals whose ore minerals could also have undergone physical and chemical weathering in the course of transport downstream (Nair and Ramachandran, 2002). The inconsistencies of the relationship between Mn and the three heavy metals in some of the samples is attributed to the possible formation of their amorphous phases by virtue of other metal accumulation mechanisms than scavenging by Mn minerals (Xiangdong et al., 2001). Such processes can also include weathering of the ore minerals and/or co-precipitation of the heavy metals in the stream sediment transport (Elza et al., 2012).

The pH of the stream sediments in general is alkaline (7.6) with few acidic extremes in the mine wastes (Fig. 8). Alkalinity in the mill tailing (ground sulfide materials) can possibly be due to the metallurgical methods practiced to concentrate gold involving cyanide solutions in a medium with pH 9 to 11. There is a general trend of pH values to increase down stream which implies the dilution effect of more water accumulation in the higher order of streams which experience occasional flux of water due to the arid climate.

### 3.3. Down Stream Patterns

There are two main streams in the study area that drains towards east and joins together before reaching the Mediterranean Sea. Figure 9A shows the two paths from which path-II is selected for assessing the downstream patterns since it experiences Pb-Cu-Zn loading from mineralization at around sample spots 45 and 28. In the following graphs, the peaks at sample spot 28 are due to Pb-Zn-Cu loading from the tributary towards sample 45 (Fig. 9B-E).

The comparison is between the concentration values of the heavy metals in their amorphous including water extractable phases (HA) and crystalline phases (AR-HA). These two phases explain the relative proportions of the metals in weathered/adsorbed and ore minerals portions of the samples respectively.

All the heavy metals except Cu and Fe showed lower concentration in the partial extraction values including water extractable values (HA) than their extraction values crystalline phases (AR-HA) in the most downstream part of the study area. Fe exhibit by far lower values in its partial extraction implying that the dominant host mineral is crystalline such as hematite which is the most common ore mineral of Iron (Fig. 9C & F). This suggests that Fe has lower scavenging role to the heavy
metals in its amorphous phases. The lower values of copper at down stream can possibly be due to erosion of the amorphous phase to the sea in the form of suspension while the portions in the crystalline phase remain in the stream sediment. This can be supported by the fact that heavy metals are not necessarily attached permanently to stream sediment particles, but they can also be remobilized through chemical, physical and biological processes (Edward et al., 2005; Jelodar et al., 2012).

Copper’s partial extraction value exceeds its counterpart, at only sample spot 28 where Mn showed elevated value in its amorphous phase (HA) implying the possible preferential association of copper and manganese in the vicinity including sample spot 36 (Fig. 9C & E). Zn, Pb and Mn revealed higher concentration values in their amorphous (HA) than crystalline phases (AR-HA) downstream. It is evident that there is a shift in the prevalence of their mineral phases at sample spot 4 with the predominance of crystalline phases upstream and amorphous phases downstream of this sample spot along the path. This phenomenon can be elaborated by the sediment load from path-1 which covers the catchment area of the old gold mining areas and mill tailings in most upstream down to the sea (Fig. 9A).

3.3. Statistical correlations

Four transects were selected to investigate the possible relationships among the three heavy metals (Fig. 10A). Such relationships include co-mineralization (mineral assemblage), adsorption (scavenging) and co-precipitation (weathering). Transect-b is found to be more relevant in the transect-wise correlation of the pollutants (Pb, Zn and Cu) and scavengers (Mn and Fe) followed by transect-c.

Scatter plots reveal the existence of association of Mn with Zn, Cu and Pb in the modes of co-mineralization and absorption. To illustrate these findings, the geochemical analysis values of Mn from the total extraction (AR), the difference between total and partial (AR-HA) as well as values from the partial extraction (HA) are correlated with Zn, Cu and Pb values from the same extraction phases. The relationship among the three heavy metals due to co-mineralization is better explained using the values from (AR-HA) while that due to absorption is better explained using the values from the partial extraction (HA).
Figure 9. A-Drainage map showing the path selected; B-downstream trend of Zn leachates; C-downstream trend of Cu leachates; D-downstream trend of Pb leachates; E-downstream trend of Mn leachates; F-downstream trend of Fe leachates
Figure 10. (A) The drainage map of the study area with four transects; Scatter plots; (B) Zn vs Cu, AR-HA, along transect-b (co-mineralization); (C) Zn vs Cu, HA, along transect-b (co-precipitation); (D) Zn vs Mn, HA, along transect-b (Adsorption); (E) Cu vs Mn, HA, along transect-b (Adsorption); (F) Zn vs Pb, HA, along transect-c (co-precipitation); (G) Cu vs Fe, HA, along transect-b (Adsorption).
Significant correlation between the concentration values of Zn and Cu in their crystalline phases (AR-HA) along transect-b is explained by the assemblage of the metals during mineralization in the upstream and consequent transport downstream (Fig. 10B). Association by weathering processes (co-precipitation) among the heavy metals is strong along transect-b as the correlation of the partial extraction values depicted (Fig. 10C).

The relationships among the heavy metals along the other transects than transect-b was found relatively weak with the possibly co-precipitation between Zn and Pb along transect-c being the highest (Fig. 10F). The main reason for these weaker correlations can be attributed to weaker degree of Pb-Zn-Cu mineralization and other factors such as the tailing materials, milling processes and local geology.

The scavenging effect of Mn and Fe has been clearly indicated by their significant correlations with the heavy metal pollutants along transect-b (Fig. 10D, E & G). In the course of stream sediment transport along transect b, amorphous manganese minerals has played adsorption role on the heavy metals when they get free through weathering and/or water dissolution.

4. CONCLUSION

This present work has further indicated the potential application of partial extraction techniques in assessing the distribution of heavy metals due to absorption. The absorption affinity of Mn to Zn and Cu has been examined using geochemical data from total and partial extraction values as well as their respective difference. Manganese was found to have stronger scavenging effect on Cu than on Zn and Pb as shown by simple statistical correlations. Iron was found to exit predominantly in its crystalline than amorphous phases.

The partial extraction technique has proved to play great role in heavy metal pollution studies by successfully detaching the heavy metals that might have been scavenged by Mn when transported by stream water or leached out in situ by hypogene soil formations. The three extraction results showed that the area is severely polluted with Pb, Cu and Zn in decreasing order. The partial extraction result is found helpful to investigate the distribution of the heavy metals in amorphous phases by enhancing the scavenging, co-precipitation and co-mineralization pattern of heavy metals. The potential application of partial extraction technique in assessing the distribution of heavy metals spatially as well as in mineral phases has been verified.
5. ACKNOWLEDGEMENT

My heartfelt gratitude is due, to Drs. Boudewijn de Smeth for his devoted assistance and guidance in the laboratory works at ITC geochemical laboratory, the Netherlands whereby the dataset was generated and served in this work for further investigation as extension beyond the scope of my M.Sc thesis. I am highly grateful to Dr. K. Bheemalingeswara and the anonymous reviewers for their important feedbacks that helped much for the initial manuscript to reach its current form.

6. REFERENCE


