Review Article

Metal (Loid)s in Farmland Soils and Strategies to Reduce Bioavailability

Abstract

High concentrations of heavy metal (loid)s in farmland soils reduces crop yield and contaminates the food chain. Exposure to HMs in the diet results in several adverse health effects such as cancer, reproductive health problems and cardiovascular diseases. Understanding the origin and fractionation of these toxic substances will provide direction for reducing their bioavailability in contaminated farmland soils. HMs are added to farmland soils through activities such as irrigation, organic and inorganic fertilization, pest control, and mining. Weathering of parent material and atmospheric deposition can also increase the levels of HMs in the soil. Fractionation of HM contaminated soils provides information on availability of HMs such as Pb, Cd, As, Cr and Cu to soil biota and plants. Several studies have reported that Pb is mostly associated with Fe and manganese oxides (reducible fraction) while Cd is mostly associated with the most mobile fraction (exchangeable fraction). The application of organic and inorganic soil amendments such as vermiculite, zeolite, composts and crop residue to contaminated farmland soils converts HMs from the plant available fractions to the less mobile fractions. HM resistant microbes can change HMs to a less mobile fraction or less mobile oxidation state. The combination of HM resistant microbes, HM tolerant plants, and soil amendments can be used to reduce mobility of HMs in contaminated farmlands.

Capsule: The mobility of heavy metal (loid)s in polluted farmland soils depends strongly on its association with soil components.

Introduction

Total soil concentrations of heavy metal (loid)s (HMs) are increasing in agricultural soils globally due to both natural and anthropogenic sources in the environment. Several studies have reported HM concentrations above acceptable limits in farmland soils [1-4]. However, total soil concentrations do not provide enough information about the availability or toxicity of HMs to soil biota and growing plants.

HMs can be bound to solid constituents of the soil such as clay, organic matter, carbonates, sulfides, iron (Fe) and manganese (Mn) oxides, which reduces its availability for plant uptake. The amount of HM available to soil biota and plants is referred to as bioavailability. Bioavailability is a better estimate of the environmental impact of contaminated soils. The bioavailability of HMs depends strongly on the sources of contamination, speciation and fractionation of HMs in the soil.

Plant uptake of HMs in contaminated soils is dependent on the fractionation and mobility of HMs in the soil. However, depending on tolerance levels, plant species may either exclude or accumulate toxic HMs from contaminated soils. Food crops growing on HM contaminated soils show a reduction in growth, performance, and yield [5]. Several studies have reported a decline in crop yield and quality in metal polluted farmland soils [6-8]. Declining crop yield due to metal pollution of agricultural soils is a threat to global food security.

HM contamination of farmland soils may also contaminate the food chain and cause health problems in nearby communities [9]. HM contamination of farmland soils has led to high concentrations of HM in food crops such as rice grains, wheat and vegetables [10-12]. These food crops are staples in many countries worldwide indicating that food chain contamination exposes a large population of consumers to the adverse effects of HMs.

HMs disrupts metabolic activities which affects virtually every organ and system in the body. Exposure to arsenic (As) has been associated with health problems such as cancer, cardiovascular and respiratory diseases, hearing impairment, reproductive health problems in pregnant women and it affects the unborn [13]. Lead (Pb) is especially dangerous to infants and young children because it can damage their developing brain while adults can suffer from reproductive
health problems, high blood pressure, digestive problems, neurological disorders, memory and concentration problems, muscle and joint pain, and kidney dysfunction [14]. Cadmium (Cd) is toxic to the kidney while chromium (Cr) exposure can lead to problems in the respiratory system, gastrointestinal system and skin [15,16].

It is important to reduce the bioavailability of HMs in contaminated agricultural soils in order to prevent the health hazards of HM and reduce economic losses to farmers due to reduction in crop yield. Sustaining high crop yields is very important to ensure adequate food supply for the increasing population in the world. There are a variety of strategies and techniques that can be used to control and reduce mobility and bioavailability of HMs in farmland soils.

While some strategies can be used to remove contaminants permanently, some others can only be used to reduce the migration of contaminants in the soil. Physical methods that severely alter soil structure such as capping of soil with cement are not suitable for farmland soils. The selection of the right technique will depend on site characteristics, operation costs, source of contamination and contaminant chemistry.

In order to effectively remove, control and reduce mobility of HMs in contaminated soils, it is imperative to identify the sources of contamination, understand fractionation of HMs and the factors that enhance their mobility in contaminated soils. Hence this paper provides a review of sources of contamination, fractionation and mobility of HMs in farmland soils as well as strategies for reducing their bioavailability.

**Heavy metal (loid)s in farmland soils**

**Sources**

Anthropogenic sources: Anthropogenic sources (Figure 1) of HMs include agricultural activities and industrial activities which contribute to elevated concentrations of HMs in the environment [17]. Sources depend on location, farm management practices and environmental conditions. Major sources of HMs reported for England and Wales include industrial activities, sewage sludge, animal manures and composts, inorganic fertilizers and lime, agrochemicals, and irrigation water [18].

The quality of water used for irrigation is important because it has been associated with metal pollution in farmland soils. Sewage irrigated soil contained much higher concentrations of Cd, copper (Cu), Pb, zinc (Zn), nickel (Ni), Cr, as and mercury (Hg) compared to clean water irrigated soil [19]. Scarcity of high quality water sources for irrigation has forced many farmers to use industrial wastewater or sewage water/sludge with high metal concentrations. Consequently, farmlands irrigated with industrial wastewater for 30 years became heavily polluted with HMs [2].

Wastewater irrigation of vegetable gardens in Zimbabwe also increased concentrations of HMs in the soil [1]. The use of wastewater also contaminates the crops growing on the irrigated farmland soils. Green vegetables irrigated with wastewater had higher concentrations of HMs than vegetables irrigated with well water in Pakistan [20]. However, in regions with arsenic contaminated groundwater, the use of well water for crop irrigation has also led to soil and crop contamination [10,12, 21].

Surface waters are also another source of irrigation water in farmlands. However, these surface water sources may be polluted due to discharge of urban and industrial waste into rivers. Irrigation of alfalfa fields with river water has been reported to account for up to 31 % of HMs in surface soils of Texas and Mexico [22]. Long term irrigation with river water polluted by mining operations in Jiangxi Province of China resulted in heavy metal (loid)s pollution of the soil with the concentrations of the two primary pollutants in the soils varying from 0.52–2.55 mg kg$^{-1}$ for Cd and 27.87–426.15 mg kg$^{-1}$ for Cu [23]. Higher concentrations of HMs were found in the agricultural soils closer to mines and metal smelters than in other soils.

The mining of HMs often lead to contamination of nearby agricultural soils [24]. For example, mining activities in Tunisia has led to high concentrations of Pb (17,229 mg kg$^{-1}$), Zn (4,813 mg kg$^{-1}$) and Cd (42 mg kg$^{-1}$) in nearby agricultural soils [3]. Elevated concentrations of HMs were also reported in wheat grown in these contaminated soils. Pb concentrations were 1824 and 54 mg kg$^{-1}$ and Zn concentrations were 1546 and 636 mg kg$^{-1}$ in the roots and shoot respectively. These concentrations are far above acceptable limits and indicate potential health risks for consumers.

Even when concentrations of HMs in irrigation water were below permissible limits, high levels of HMs were found in vegetable cultivated in farmlands of Northern Ethiopia [25].

![Figure 1: Sources of contamination in farmland soils.](Image 1.png)

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This suggests that the established acceptable limits were not effective to prevent contamination of the food chain. Similarly, the use of treated sewage water in Saudi Arabia led to an increase in HM levels in soils and test crop (okra) [4]. This suggests that pretreatment of sewage water was not able to remove a large portion of heavy metals contained therein.

The use of organic manures such as biogas slurry in vegetable farmlands of the Taihu Basin, China led to high concentrations of Ni, Zn, Cd and Pb in vegetables and soils above acceptable limits [11]. Livestock manures and sewage sludge were found to be responsible for 37–40 and 8–17% of total Zn and Cu inputs, respectively, in farmland soils of England and Wales [17].

Long term use of agrochemicals such as fertilizers and pesticides (insecticides, herbicides and fungicides) to increase crop yield often contribute to HM pollution because of their composition and soil characteristics. In a past study, the high concentrations of As, Pb, and Cd in agricultural soils were attributed to the application of agrochemicals [26]. A previous study has also shown that copper based fungicides have the potential to increase concentration of Cu in the soil to phytotoxic levels [27]. P fertilizer treatments increased the Zn solution concentration in 83% and 53% of treatments while it increased the concentration of Cd by 87% and 80% in field and laboratory experiments respectively [28]. Agricultural activities that may increase HM concentration in the soil include use of contaminated irrigation water, mostly from industrial or domestic wastewater, sewage sludge or bio solids; use of farmyard manure with high HM concentration; use of agrochemicals such as fertilizers and pesticides.

Natural sources: Natural sources of heavy metal (loid)s (HMs) include parent material and atmospheric deposition from volcanic activity and forest fires. Soils are formed as a result of a long period of physical and chemical weathering of the underlying parent rock. Physical weathering breaks the soil into smaller sizes while chemical weathering releases minerals present in the parent rocks which forms part of the soil. These minerals are sources of essential plant nutrients and heavy metal (loid)s.

Parent material was associated with the high concentrations of Cu, Cr, Ni, Pb and As of farmland soils of the urban–rural transition zone in [29]. The soil parent material and point sources of pollution significantly contributed to Cr, Ni, Cu, Zn and Cd levels in agricultural soils [30]. The extent of contamination has been found to depend greatly on the mobility of the metal (loid)s.

It has been estimated that atmospheric deposition was responsible for about 25–85% of total inputs of HMs in agricultural soils in England and Wales [18]. HM contamination in topsoil from remote farmland in China was attributed to long range transport and atmospheric deposition of HMs since no pesticides or fertilizers have been used in the remote farmland stations studied and no industries were situated nearby [31].

Sometimes, human activities such as urbanization, industrialization and mining also contribute to atmospheric deposition. Increased levels of metals in the soil have been attributed partly to atmospheric fallout from a local ore smelter in alfalfa fields in Texas and Mexico. Metal concentrations in unwashed alfalfa forage tissue were at least five times less than those in soil and showed no consistent association with soil concentrations [22].

Industries are responsible for atmospheric deposition of mercury (Hg) in agricultural soils of a rapidly industrializing area [30]. Ferroalloy industries in Brescia province, Italy produced particulate emissions enriched in Mn, Pb, Zn, Cu, Cd, Cr, Fe, and aluminum (Al). This was responsible for the HM (Al, Cd, Fe, Mn, Pb, and Zn) contamination of home gardens close to the industries. Unwashed spinach showed higher metal concentrations than washed spinach indicating atmospheric deposition [32].

Sometimes, the presence of HMs in the soil may be due to multiple sources. The results of a multivariate geostatistical method showed that Cd, Cr, and Cu in reclaimed farmland soils was associated with chemical fertilizers, Pb and Zn with atmospheric deposition and Ni was associated with the electroplating factories’ wastewater discharge [33]. Atmospheric deposition of Pb and Zn may be from vehicular emissions from nearby roads and highways and industrial emissions.

In a representative agricultural suburb in Beijing, China, multivariate and geostatistical analyses indicated that soil contamination of Cd, Cu and Zn was mainly derived from agricultural practices. As and Pb were due mainly to soil parent materials, while Hg was caused by the atmospheric deposition of particulate matter [34]. In Taiyuan City, China, HMs (Cd, Cu, Hg, Pb, Zn, and Cr) contamination of agricultural soils was attributed to farming practices while as and Ni were associated with parent material [35]. Contamination of soils with as is commonly associated with parent material because it can be mobilized naturally through evaporative concentration, leaching from sulfides by carbonate, and microbial mobilization [36].

Bioavailability of heavy metal(loid)s in farmland soils

Bioavailability provides information on the amount of contaminants available for chemical reactions or plant uptake. Total HM concentrations are insufficient to determine its environmental impact in contaminated soils because HMs are often bound by different soil constituents such as organic matter, Fe/Mn oxides and clay minerals [37]. Sequential extractions with different extractants are often used to partition metals into operationally defined chemical associations in the soil.

Though there are several methods of sequential extractions for HMs, the most common one is the five stage extraction method (Figure 2) which separates HMs into five different fractions; water soluble and exchangeable fraction; bound to carbonate; bound to iron and manganese oxides (reducible oxides/oxidic fraction); bound to organic compounds and sulfides; and residual fraction [38].
Figure 2: Fractionation of heavy metal (loid)s in the soil.

Another method was developed by the European Community Bureau of Reference, called the BCR method [39,40]. The BCR method is a three-step procedure that extracts the exchangeable and carbonate-bound fractions in a single step versus the two steps used in the Tessier procedure [41]. The fractions extracted by the BCR method are: exchangeable (extraction step 1), reducible (extraction step 2), oxidizable (extraction step 3) [42]. The fractionation method for metals may not be appropriate for metalloids such as vanadium (V), As, and Se. The method for phosphorus (P) fractionation is used for As contaminated soils because of similarities with As and P. Sequential extraction of As is a four-step procedure separating into four fractions; water-soluble and exchangeable fraction, Fe-bound fraction, and Ca-bound fraction [43].

Fractionation of heavy metal (loid)s in Agricultural soils; Fractionation of HMs into operationally defined fractions is important because it shows the distribution between different soil components and provides information on potential mobility of HMs in the soil. Fractionation of HMs in farmland soils depends on the soil properties, sources, cropping systems, farm management practices and plant species.

Pb was mainly associated with the reducible fraction (bound to Fe and Mn oxides) while Cu, Zn and Cd were dominated by the residual fraction in agricultural and nonagricultural soils [44]. Similarly, V and Pb were associated with the reducible fraction, Hg and Cd with the exchangeable fraction, Co with the carbonate fraction, while Cr, Cu, Zn, Ni were mostly associated with the oxidizable fraction in agricultural soils contaminated by wastewater irrigation in Ethiopia. There was a higher concentration of HMs in the non-residual fractions than the residual fraction suggesting high mobility of HMs in the soil [45].

Some other studies have also shown that Pb was strongly associated with the reducible fraction. Fractionation of agricultural soils close to mines and smelters showed that Pb was in the reducible fraction; Cd in the exchangeable and carbonate fraction while Cu and Zn were mainly in the residual fraction [23]. The reducible fraction was the largest fraction for Cu and Pb in agricultural soils contaminated from improper disposal of tailings [24].

On the contrary, Pb and Cu seemed to be more tightly bound to the organic fraction while cadmium was mostly bound to the Fe/Mn oxides fraction in contaminated soils from Sicily; though a significant amount of all HMs (Cd, Pb, Cu, Co, Cr, Fe, Mn, Ni, Zn) were bound to the organic fraction [46]. The oxides fraction usually has more affinity for Pb than Cd, so this may be due to high concentration and strong negative charges on organic fraction of the soil.

The partitioning of agricultural soils contaminated with wastewater or smelter dust demonstrates how organic matter content may influence fractionation of HMs. HMs in the wastewater polluted soil (with high organic matter) were mostly bound to the organic fraction in the wastewater while they were mainly associated with Fe/Mn oxides in the smelter dust polluted soil. Cadmium was mostly in the exchangeable fraction in both soils while lead in the smelter dust polluted soil was dominated by the reducible fraction [47].

Cd in soil profiles was predominantly associated with the exchangeable fraction while Pb was closely associated with reducible fraction in a BCR extraction of contaminated vegetable soils from the Pb/Zn mining and smelting areas of China [48]. The mobile fractions (exchangeable, reducible, and oxidizable) of Mn, Cd and Pb were higher than the immobile fractions (residual) in a modified BCR extraction of contaminated agricultural fields of Çanakkale, Turkey [49].

There was also a strong correlation between concentrations of HMs in corn grain and the more available fractions (exchangeable and reducible) [49]. Cu Cr, Ni and Zn were mostly in the residual fraction; Cd was mainly associated with the exchangeable/carbonate and reducible fractions, while Pb was mainly associated with the reducible fraction in a BCR extraction of compost amended and unamended field plots [50].

Several studies have reported that Pb is mostly associated with the reducible fraction (bound to Fe/Mn oxides) while Cd is mostly associated with the most mobile fraction (exchangeable fraction) [24,44,45,48]. This may be due to higher affinity of oxidic surfaces in the soil for Pb than for Cd. This may explain why Cd had the highest transfer factor in vegetables grown in farmlands amended with biogas slurry [11]. Cd was probably very mobile in the amendment and treated soil.

Guillén et al. [17], reported that Cd poses a serious threat to human health and the environment due to the high percentage of mobility. The high mobility of Cd might be responsible for the high accumulation of Cd in wheat grains reported in China [51]. Additionally, a study on metal accumulation in farmland...
soils across China reported that Cd had the highest pollution index [52]. Furthermore, Cd has been reported as a major contaminant in rice fields of East and South Asian countries [53]. All these suggest that Cd is a threat to food security in Asian countries due to its high mobility.

It has been shown that metals with anthropogenic sources were mainly associated with residual, oxidizable and reducible phases which are relatively immobile and only potentially bioavailable to plants. However, micronutrients such as Zn and Cu showed a tendency to become more readily mobile and bioavailable as their total content in soil increased and were more easily transported to the edible parts of lettuce than other pollutants [54].

Fractionation of HMs in contaminated agricultural soils also varies with soil classification. A greenhouse experiment with rice grown in two paddy soils showed that Pb was more mobile in the Ultisol (Plinthudic Aquults) than in the Inceptisol (Fluventic Umbrepts) because Pb in Ultisol was mostly associated with the residual and exchangeable fractions, while Pb was mainly associated with the residual and Fe/Mn oxides fractions in the Inceptisol [55]. This suggests that Pb will be more available to plants in the inceptisol while it will be less available in the ultisol.

**Phytoavailability of heavy metal (loid)s in agricultural soils:** Phytoavailability is a measure of the amount of HMs available in the soil solution for plant uptake. Phytoavailability of trace elements especially HMs depends on their speciation rather than on total soil concentrations [56]. Speciation of HMs determines their fractionation in the soil and availability for plant uptake. Phytoavailability of HMs can be measured either by single extractions or directly by plant uptake in contaminated soils.

Four single extractants commonly used include: calcium chloride (CaCl₂), diethylene triamine pentaaetic acid (DTPA), ethanoic acid (CH₃COOH), and water [57]. Some other single extractants include Mehlich 3, ammonium nitrate (NH₄NO₃), sodium nitrate (NaNO₃), ethylene diamine tetraacetic acid (EDTA) and nitric acid (HNO₃) [58].

Some studies have reported significant correlation between concentrations of HMs in single extractants and plant uptake [57–60]. Lavado et al. [59], reported significant correlations between EDTA extractable copper in soils and maize grown in biosolids amended soils. Wang et al. [57], also reported that the phytoavailability of HMs to Chinese cabbage was significantly correlated with concentrations of metal extracted by CaCl₂. However, phytoavailability was influenced by soil properties such as soil pH, organic matter and cation exchange capacity (CEC).

Fractions of HMs in sequential extractions have also been correlated with plant uptake. The Cr concentration of rice grain had the highest correlation with water soluble and exchangeable Cr while the straw Cr best correlated with the Fe and Mn oxide fraction [61]. The sum of the metal percentages extracted from the first two fractions of Tessier’s procedure was found to be suitable in predicting the phytoavailability (to Lactuca sativa and Spinacia oleracea) of most of the HMs present in the soil of a contaminated site in Piedmont, Italy [62].

On the contrary, NH₄NO₃ extractant was ineffective in determining the availability of HMs (Cd, Cr, Cu, Ni, Pb and Zn) for sugarcane [63]. Even though DTPA extraction indicated higher metal availability in sludge amended soils, it did not increase HM concentration in plant leaves, suggesting that determination of HM availability by extraction may not accurately predict phytoavailability [64].

A series of empirical equations involving extractable metals, solid–phase fractions and soil properties have been used to significantly predict the phytoavailability of Mn and Zn for barley [65]. The importance of solid–phase Cd fractions, EDTA fraction and/or HNO₃ fraction, in predicting phytoavailability of Cd in soils amended with manure compost have also been demonstrated [66].

Leafy vegetable *Amaranthus hybridus* had 1.2–8.2 folds enrichment of Cr, Co, Ni, Cd, Cu, and Pb than *Abelmoschus esculentus* in amended soils irrigated with wastewater suggesting that phyto–accumulation of HMs is plant–specific and dependent on physiological set-up [67]. Metal accumulation in vegetables grown in urban gardens varied by crop type further confirming that phytoavailability may vary with crop type [68].

This may explain why phytoavailability of Cd to rice grain was strongly correlated with Mehlich–3–extractable Cd while the same Mehlich–3 extractant was not efficient to evaluate Ni phytoavailability to maize plants [69,70]. This suggests that soil extractions may not be the best method to determine phytoavailability of Cd. It has been shown that the soil type, Cd source, loading rate, soil pH, and plant tissue are important factors in evaluation of Cd phytoavailability [66,70].

**Factors influencing the mobility of heavy metal(loid)s in agricultural soils**

**Soil properties:** Soil properties such as pH, soil texture, organic matter content, Fe/Al oxides and moisture content control metal mobility (Figure 3) in soils [71]. Amongst all, pH is a major factor controlling the mobility of HMs in contaminated soils. Soils with low pH have higher concentrations of HMs in the most mobile/soluble phase (exchangeable fraction) [72]. HM concentrations significantly increased while pH decreased down the soil profile to a depth of 30cm [73]. HMs are more available at a low pH probably because there is a higher concentration of protons displacing the metals from exchange sites on the solid phase of the soil.

Recently, it was shown that increasing the soil pH to 10 can decrease the zinc content of *Valeriana officinalis* significantly [74]. Decrease in heavy metal mobility with increasing soil pH has been attributed to precipitation of hydroxides, carbonates or formation of insoluble organic complexes [75]. On the contrary, increasing pH increased Cu and Mn in plant tissue of *V. officinalis* probably due to formation of soluble organo–metal complexes at high pH [74].
A past study showed that Zn concentrations in the soil increased with increasing soil acidity while it decreased with increasing clay content of the soil [76]. This indicates that soil texture also plays an important role in metal mobility and retention in the soil. Generally, there is a higher mobility of HMs in sandy soils than clayey soils [77]. Adsorption of the HMs on clay surfaces retards the movement of HMs and reduces leaching and run off.

Soil organic matter (SOM) is also an important adsorbent for HMs in agricultural soils because of the presence of charges which reduces their mobility and availability in the soil. The presence of SOM decreased Zn binding energy, increased immobilization of Zn and decreased the proportion of nonspecific adsorbed Zn [78]. Metal release due to solubilisation of organic matter and formation of organo metallic complexes has also been reported in agricultural soils [79,80].

Fe and Mn–oxides and hydroxides also play an important role in heavy metal sorption due to their high specific surface area and reactivity [81]. The strong affinity between acidic surfaces (of Fe, Al and Mn) in the soil and HMs makes them effective adsorbents of HMs. Hence, increasing concentrations of Fe/Al/Mn oxides in the soil reduces the mobility and phytoavailability of HMs in the soil [77].

The mobility of HMs is determined by their concentration in the soil solution which depends on amount of soil moisture. It was found that soil moisture regime significantly affected the transformation rate from the more mobile fractions (exchangeable) to the less labile ones (Fe-Mn oxide and organic matter-bound fractions). In general, the paddy soil under flooding regime had higher metal reactivity compared with 75% field capacity and wetting-drying cycle regimes, resulting in the more complete movement of metals toward stable fractions [82].

It is well known that metal availability decreases as the soil ages [83-87]. During aging, as moved from the more available (non-specifically and specifically sorbed) to less available (amorphous and crystallized Fe/Al) fractions while Pb moved from the first three fractions (exchangeable, carbonate and Fe/Mn hydroxide) to organic fraction [88]).

**Environmental conditions:** Environmental conditions such as rainfall, temperature and wind greatly impact the fate and transport of metals [89]. Rainfall events increase soil and water runoff which also increases the mobility of HMs in agricultural soils [90]. Less than 4% of Cu and more than 58.3% of Zn were transferred from composts to groundwater under simulated rainfall conditions in a soil column experiment [91]. This suggests that the effect of rainfall may depend on the chemistry of HM being investigated and their interaction with the soil. Rainfall can also increase the area impacted by metal contamination by increasing the distance travelled by HMs applied in composts, sewage sludge, farmyard manure and bio solids.

Chemical composition of rainwater is responsible for its effect on the mobility of HMs. It is well known that gaseous air is a mixture of gases such as oxygen, nitrogen, carbon dioxide and sulfur dioxide. Normal rain is slightly acidic because the carbon dioxide in the air dissolves in rainwater to form carbonic acid, a weak acid. Acid rain occurs when there are high concentrations of sulfur dioxide and nitrogen oxides in the air. Dissolution of sulfur dioxide in rainwater may produce sulfuric acid while nitrogen oxides may form nitric acid. These are both strong acids and their presence in rainwater acidifies the soil.

Acid rain lowers the soil pH and increase metal mobility in the soil [92]. Acid rain is caused by volcanic activity, fossil fuel emissions, wildfires and lightning and has been reported in several countries including USA and South Korea [93]. This acidic input may further exacerbate metal pollution and cause leaching in agricultural soils. A study on the effect of simulated acid rain on metal mobility found that Cu and Zn were released by the soil in larger amounts than Pb and Cd [94].

Temperature is also another environmental condition that may impact metal mobility. High temperature will increase rate of organic matter decomposition (Figure 3) which may lead to formation of organic acids that acidifies the soil and increase metal dissolution and mobility. High temperature may also lead to higher microbial activity which may either decrease or increase metal mobility. A past study has shown that microbes reduced as and metal concentrations in the absence of organic matter but increased metal concentrations in the presence of organic matter [95].

Higher microbial activity may lead to lower metal mobility due to microbial reduction of metals to a lower redox state that is less mobile [96]. Bacteria reduced metal mobility (reduced water-soluble Pb) due to adsorption by bacterial cell walls and possible sedimentation reactions with phosphate or other anions produced through bacterial metabolism [97]. Metal mobilization by soil microorganisms can arise from complication and methylation while immobilization can result from sorption to biomass or exopolymers, transport
and intracellular sequestration or precipitation as organic and inorganic compounds [98].

**Plant characteristics:** The presence of plants in the soil can increase metal mobility by increasing acidity via processes like root exudation, acidifying effects of CO$_2$ produced in root respiration and H$^+$ released due to cationic nutrient intake [99]. Plant roots produce low molecular weight organic acids (LMWOA) which have the potential to enhance metal mobility in the soil by reducing soil pH and forming complexes with HMs [100].

On the other hand, some studies have reported that root exudates reduced metal availability due to their strong metal binding ability [101,102]. The immobilization of metals by root exudates may be a coping mechanism for exclusion (reduced uptake) of toxic HMs in nonmetal hyperaccumulators [103,104]. Metals present in the rhizosphere of hyperaccumulators are more bioavailable than those in the rhizosphere of non hyperaccumulators [105–107]. However, root exudates produced by the Cd and Zn hyperaccumulator *Thlaspi caerulescens* did not significantly increase metal mobility [101,108].

Root exudates are an abundant source of energy and nutrients for microorganisms which exist in associations with plant roots in the soil [109]. Plants usually form symbiotic associations with microorganisms such as rhizobacteria and arbuscular mycorrhizal (AM) fungi. AM fungi in symbiotic associations with a range of plant species are able to enhance uptake of nutrient elements and water by host plants through their extraradical mycelial networks [6].

Some of these microorganisms are able to tolerate high concentrations of HMs and have evolved resistance strategies [110]. Arsenic resistant bacteria have been isolated in the rhizosphere and plant tissues of an as hyperaccumulator [111,112]. Arsenic resistant rhizobacteria and AM fungi have been shown to increase as uptake and translocation in the as hyperaccumulator *Pteris viitata* [113–115]. Similarly, a heavy metal resistant bacteria strain, isolated from the rhizosphere of *Elscholtzia splendens* increased the uptake and translocation of copper (Cu) in *E. splendens* [116].

The presence of a bacterium and fungus in agricultural soil increased plant biomass and HM uptake in corn (*Zea mays*) [117]. However, it is also possible for AM fungi and rhizobacteria to decrease metal uptake of the host plant and protect them against heavy metal toxicity. Copper resistant bacteria increased root and shoot length, plant dry weight and leaf area, reduced plant uptake and protected lentil plants from copper toxicity. The plant growth promoting bacteria was also able to improve anti-oxidative defense mechanism of the plants by increasing activities of anti-oxidant enzymes such as catalase and superoxide dismutase [118].

**Farm management practices:** Management practices such as cropping systems, land management, tillage methods, fertilization and water management influence HM mobility in agricultural soils. A study has shown that cultivation can decrease the humified carbon content and metals bound to the humic fraction of organic matter suggesting that cultivation may potentially increase the mobility of heavy metals [119].

A study investigated the metal contents in open field cultured soils versus soils cultured under shelters. Results showed that the respective average concentrations of Al, Fe and Pb in the soils were 2.66, 1.41 and 2.46 times higher in open fields than soils under shelters, whereas the respective average soil concentrations of Cu and Zn were 1.48 and 1.41 times lower [121]. This was attributed to exposure of open fields to soil erosion from runoff and leaching of Cu and Zn within soil profile [121].

Change in land use or conversions also affect metal mobility. The conversion of farmland from wetland resulted in an increase in the concentration of Pb and Cr in the soil while the concentrations of Zn, Cu, Ni, and Cd decreased when wetland was converted into farmland. The levels of Pb increased by approximately 28.6 % when dry land was converted to wetland and by 24.7 % when paddy land was converted [122].

Land application of organic manure may lead to HM contamination of the soil at the surface layer. Tillage practices in farmland soils may affect mobility of HMs by redistributing HMs. In a previous study, no-tillage simulations showed consistent HM accumulations in the surface soil. Even though soil tillage reduced HM concentrations over the top 20 cm of the soil profile, it increased their transfer to deeper layers [123].

The effect of tillage methods, crop rotation and fertilization on Cd availability in tuber and cereal grains was determined in a study in Europe. Results show a higher plant uptake of Cd in wheat grain in continuous wheat under direct drilling than under reduced till or conventional cultivation [124]. Cd concentration in grain was highest in wheat grown after a legume and lowest in wheat grown after a cereal. Cd in wheat grain and potato tubers can increase with increasing rates of nitrogen regardless of the crop rotation [124].

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A recent study has reported that fertilization generally increased more available fractions (exchangeable and carbonate-bound Mn) in the soil [125]. Similarly, mineral fertilizers increased the mobility of Ni and Cu in a study in Poland [126]. Long-term fertilizer use has been reported to increase soil metal content, soil organic carbon, cation exchange capacity, and decrease soil pH [127]. The acidification of the soil by inorganic fertilizers may be responsible for the increase in metal mobility.

### Strategies for reducing bioavailability of metal(loid)s in contaminated farmlands

**Stabilization:** This method involves the use of soil amendments to limit contaminant migration in contaminated soils. Highly solid materials like cement that changes soil structure cannot be used on farmlands. Organic amendments such as crop residues, farmyard manure, composts, and biochar have been used to reduce mobility of HMs in agricultural soils. Similarly, inorganic amendments such as lime, vermiculite and phosphates have been used to immobilize HMs in contaminated farmland soils.

The incorporation of crop residue into the soil may be an economical method of reducing metal mobility because they are readily available on the farm (Table 2). It also takes care of waste disposal on the farm. The addition of rapeseed residue decreased the mobile fraction of Cd by 5–14% and Pb by 30–39% through their transformation into less available fractions, and reduced metal availability to rice plant in a contaminated paddy soil [128].

Addition of rice straw ash (RSA) to rice paddy soils has been shown to retard the movement of Cu by suppressing the release of Cu into the soil solution and transforming Cu into less soluble forms [129]. Rice straw (69.5% reduction) was more effective at reducing Cd uptake in shoots of maize compared with biochar (50.9% reduction) and wheat straw (66.9% reduction) [130].

Biochar, produced from carbonization of organic wastes, has been shown to reduce exchangeable fractions of metals in the soil and decrease plant uptake of HMs in crops such as maize, rice and lettuce [131–133]. Even though biochar has demonstrated potentials of reducing metal mobility in contaminated soils, identifying heavy metal contamination in biochar is crucial before it is used as a soil amendment [134]. Addition of metal rich biochar to lightly contaminated soils may increase metal accumulation in the soil [134].

Even though organic manure may also be a source of HMs, farmyard manure (36–45% reduction) was more efficient than rice husk (23% reduction) and straw dust (14% reduction) in reducing phytoavailability of Cd to *Amaranthus caudatus* [135]. Past study has also shown that increasing rates of Cd applied through chicken or pig manure compost increased its precipitated and residual fractions [66].

Farmyard manure such as pig slurry has been shown to reduce the mobility of Cu in contaminated soils [136]. Although, the application of cow manure and compost to a calcareous contaminated soil increased the solubility of Cu in an incubation experiment, it reduced the mobility of Zn and Pb in the same soil [137]. This suggests that factors such as type of organic amendment, soil type and metal chemistry may affect metal mobility.

Addition of municipal solid waste and manure composts to soil reduced bioavailability and plant translocation of Cu and Zn [138]. It has also been shown that animal manure compost had lower Cd plant uptake (spinach) than soils treated with chemical fertilizers [139]. Furthermore, the addition of 2.5% (dry w/w) municipal compost reduced the Cd concentration in onions, spinach and lettuce by up to 60% in garden soils [140].

Use of fly ash to stabilize sewage sludge has been shown to reduce metal availability to corn by converting metals to less available fractions in the soil [141]. Decontaminated sludge has also been used to reduce bioaccumulation of metals in maize cultivated farmland soils [142]. However, lime was more effective in reducing metal extractability than sewage sludge mixed with green waste compost [143].

A combined application of limestone and red mud reduced Ca(NO3)2 extractable As, Cd, Pb, and Zn by 58%, 98%, 98%, and 99%, respectively which led to reduction in plant uptake along with an increase in microbial activity [144]. Combined soil amendments limestone+sepiolite and hydroxyhistidine+zeolite reduced exchangeable fraction of metals (Pb, Cd, Cu, and Zn) and inhibited uptake and accumulation of Pb, Cd, Cu, and Zn in rice plants [145]. Inorganic amendments such as vermiculite have also been shown to reduce plant uptake of HMs from contaminated soils [62].

Lime based materials increase pH which reduces metal mobility while other inorganic amendments such as vermiculite and zeolite provide surfaces for adsorption of HMs. A recent study has shown that pH change-induced soil immobilization (dolomite, steel slag, and lime) was more effective than sorption agents (zeolite and compost) in reducing plant uptake of Cd.
and Pb in rice grains grown in a contaminated paddy soil [146]. Another study confirmed that pH control was the management option most likely to reduce phytoavailable Cd and Pb, since organic matter content and oxalate extractable sesquioxides only partially influenced HM retardation ($K_r$) [147].

Although, phosphates have been identified as a source of HMs in agricultural soils, they are widely used to immobilize metals due to their strong affinity for metals [148-150]. Phosphate was able to reduce plant uptake of metals by maize and reduce plant uptake of Pb, Cd and Zn by vegetables [151-153]. Recently, it has been shown that the application of phosphate fertilizer and lime was more effective than sawdust in reducing metal uptake by lettuce, especially for Cu and Pb [8].

Crop–Hyper accumulating plant rotation: Hyper accumulating plants are plants that are able to tolerate and accumulate HMs above standard concentrations in their above ground biomass. They can be used to reduce contaminant levels in contaminated farmlands by planting them ahead of crops to be grown on the farm. Hyper accumulating plants may be planted for about one or two years before the land is cultivated with food crops again. Selection of hyper accumulating plant to be used in phytoremediation will depend on the primary contaminant in the soil. Plants that are tolerant to multiple metal (loid)s will be particularly useful in soils contaminated with more than one contaminant.

About 500 plant species have been reported to accumulate HMs [154]. One of them, the arsenic hyper accumulator, Pteris vittata, has been used to reduce contaminant levels in arsenic contaminated soils [155-157]. The fern can also be used in multi-metal contaminated soils because it is tolerant to a wide variety of metals such as Pb, Cd, Zn, Se, Si, Sb, B and Hg [158-161].

There was a 3.5–11.4% reduction in the total soil as when the fern was grown in five contaminated paddy soils over a nine month period [162]. Results showed that As concentrations in the straw and grain of rice grown following P. vittata decreased by 17–82% and 22–58% respectively, suggesting that the fern reduced bioavailable As in contaminated paddy soils probably due to high uptake of As by the fern [162].

The Cd/Zn hyper accumulator Thlaspi caerulescens can be used in Cd/Zn contaminated soils though it’s sensitive to Cu [163]. Hence, it cannot be used in soils with high levels of Cu. Multi-metal hyper accumulating plant such as Ipomea spp can accumulate Cd, Cu, Mn and Zn showing potentials to be used in multi-metal contaminated soils [163]. Helianthus annuus and Hemidesmus indicus have both been shown to be Pb hyper accumulating plant species and can be used in Pb contaminated soils [164].

Plants that are tolerant but accumulate HMs only in the roots can also be used for stabilization of HMs in contaminated soils. Eight plant species, Alternanthera philoxeroides, Artemisia princeps, Bidens frondosa, Bidens pilosa, Cynodon dactylon, Digitaria sanguinalis, Erigeron canadensis, and Setaria plicata were recently identified as good candidates for stabilization of Mn, Cd, Pb and Zn contaminated sites [165]. Plants suitable for stabilization should have an extensive root system and keep the translocation of metal (loid)s from roots to shoots as low as possible in soils with high HM concentrations [166].

Plant varieties with low translocation of HMs from root to shoot can also be selected for cultivation in metal contaminated soils. Leafy vegetables namely, spinach, amaranthus and mustard seem to be unsafe and not suitable for cultivation on heavy metal contaminated soil while fruit type vegetables could be suggested for cultivation on Cd contained soil [167]. Alfalfa, field pumpkin, and barley had the highest Cu concentration in their roots suggesting they could be good candidates for stabilization of copper contaminated soils. But alfalfa and field pumpkin also accumulated Zn in the leaves indicating they may not be good for stabilization of Zn contaminated soils [168].

The efficiency of hyper accumulating plants can be increased by the addition or stimulation of soil microorganisms in metal contaminated soil. Soil microorganisms support the establishment and growth of plants on HM contaminated soils by producing plant growth hormones, inducing siderophores, solubilizing phosphorus; they increase the bioavailability of metals for plant uptake by hyper accumulating plants and reduce the level of growth inhibiting stress hormone in plants growing in metal contaminated soils [169].

Heavy metal (loid) resistant microbes: HM resistant soil microorganisms can be used to convert both organic and inorganic contaminants to less toxic forms. The method is more widely used to break down toxic complex organic compounds to nontoxic simpler forms. HMs cannot be broken down, so the method has its limitations. HM resistant microbes reduce the toxicity of HMs by converting them from a toxic oxidation state to nontoxic oxidation state or converting them from the more plant available fractions to the less available fractions. These transformations involve redox reactions, precipitation, and sorption and desorption reactions.

The reduction of Cr (VI) to Cr (III) is a detoxification pathway because Cr (VI) is more toxic than Cr (III). In addition, Cr (VI) is more bioavailable than Cr (III) at high pH; hence, the reduction of Cr (VI) to Cr (III) reduces its mobility in contaminated soils [170]. An indigenous actinomycete, Streptomyces sp. MC1 was able to reduce up to 94% of the Cr (VI) after 7 days of incubation showing almost complete Cr (VI) removal from soil without any previous treatment [171].

The addition of bacteria to chromium contaminated soils led to a drastic decrease in water soluble, exchangeable and carbonate bound Cr (VI) while there was an increase in carbonate–bonded Cr(III), Fe and Mn oxides–bonded Cr(III) and organic matter–bonded Cr(III) [Li et al., 2013] [172]. This shows that the more available fractions of the more mobile Cr (VI) were converted to the less mobile fractions by soil microorganisms.

Sulfate reducing bacteria reduced the concentrations of the exchangeable fraction of Cd by 70% while the Fe–Mn oxides
and carbonate fractions increased in Cd contaminated soils [173]. *Rhodobacter sphaeroides* reduced Pb phytoavailability to wheat seedlings by decreasing the exchangeable fraction and increasing the residual fraction. The plant available fractions were converted to less mobile fractions [174]. The bioavailability of Pb in contaminated soil was reduced by incubating with *Phanerochaete chrysosporium* and straw. The concentration of exchangeable Pb in the soil was reduced thereby reducing its toxicity [175].

Indigenous soil microflora was used for in-situ immobilization of HMs in a heavily contaminated soil in two stages. The first stage involved the solubilization and leaching of metals from the top horizon to the sub horizon due to the joint action of the soil microorganisms and leach solutions used to irrigate the soils. The dissolved contaminants were then precipitated as the insoluble forms as a result of the activity of the sulfate-reducing bacteria inhabiting the sub horizon [176].

Microbes are also used to enhance removal of HMs from the soil by leaching. Iron reducing microorganism *Desulfuromonas palmitatis* was used to induce the reductive dissolution of iron oxides and release of retained As. EDTA was also used to remove metal contaminants, such as Pb and Zn, through the formation of soluble metal chelates. Bacterial activity was found to have accelerated the removal of arsenic, which increased from the value of 35% obtained during the pure chemical treatment up to 90% in the presence of *D. palmitatis* [177].

A mercury resistant bacteria strain, SA2, was able to volatilize mercury by producing mercuric reductase enzyme. About 79% mercury in a mercury supplemented culture suspension was volatilized in 6 h showing its potential to be used in mercury contaminated soils [178]. This process reduces contaminant levels in the soil while other methods involving redox and fraction transformations limit mobility of contaminants.

**Soil washing**: The process of removing soil particles enriched with contaminants or using aqueous solutions to mobilize and leach the contaminants out is called soil washing [179]. Soil washing is able to permanently remove contaminants from contaminated soil and contaminants can be recycled from washing solution [180]. However, the physiochemical properties of soil may change after physical separation and chemical extraction processes, which may eventually lead to a decline in soil fertility [181].

Chemical washing of Cd contaminated paddy soils with calcium chloride solution removed 55% of Cd from the exchangeable fraction and 15% of Cd from the acid-soluble fraction. Even though soil washing changed the soil fertility, it could be corrected and had no effect on rice growth, but reduced the average Cd concentration in rice grains by about two-thirds compared to a control plot [182].

Ferric chloride solution has also been used for soil washing of Cd contaminated paddy soils [183,184]. Washing with FeCl₃ led to the formation of Cd–chloride complexes which enhances Cd extraction from the soils. Although, soil fertility was affected by soil washing with 45 mM FeCl₃, it had no negative effect on rice yield and lowered the Cd concentration of rice grain and rice straw [183]. In a similar study, soil washing with calcium chloride solution had no effect on growth of soybean but the seed Cd concentration decreased significantly [185].

Soil washing has been used to effectively reduce toxicity of metal polluted garden soils by removing the plant available fractions. HM contaminated garden soil was washed with EDTA solution in a pilot-scale remediation plant. Soil washing with EDTA removed 75%, 26% and 66% of Pb, Zn and Cd, respectively, from the garden soil, while 71% of EDTA was recycled and no waste water was generated [186]. In another study, soil washing of a metal contaminated vegetable garden soil with EDTA reduced the metal burden by 80, 28 and 72% for Pb, Zn and Cd respectively. However, the treatment significantly decreased the potential soil enzyme activity and the washed soil did not recover its biological properties [187].

The major limitations of soil washing are its effects on soil health and fertility. Although the amount of readily available As in soils were reduced after soil washing with different solutions, the bioassay tests showed that the washed soils had lower seed germination, shoot growth, and enzyme activities which was attributed to the acidic pH and/or excessive nutrient contents of the washed soils [188].

The choice of soil washing solutions and their washing order are critical to ensure high removal efficiencies of metal fractions and sustain soil fertility in contaminated soils [189]. Two-step washing of Cu contaminated soil with 0.34 M citric acid and then 1500 mg L⁻¹ dissolved organic matter (DOM) solution (pH 8.5) was found to be most favorable for the soil than the conventional EDTA extraction. The citric acid and DOM treatment removed 91% Cu from the topsoil while the organic matter, cation exchange capacity, plant-available nitrogen, and available phosphate content increased by 28.1%, 103%, 17.7%, and 422%, respectively [190].

The use of a degradable chelator, N, N-bis (carboxymethyl)-L-glutamic acid (GLDA), as washing solution removed 71.34%, 81.02%, and 50.95% of Cd, Pb, and Zn respectively from a polluted farmland soil. GLDA washing solution was as effective as the conventional EDTA washing, and was able to retain most of the soil nutrients [125].

**Conclusion**

The sources of HMs in agricultural soils include irrigation with wastewater, applications of agrochemicals, organic and inorganic amendments. Natural sources include parent material and atmospheric deposition. Proximity to mines and smelters also increases concentrations of HMs in farmlands. Partitioning of HMs in agricultural soils depends on source, soil properties and farm management practices. The availability of HMs for plant uptake depends on their fractionation within the soil. Soil properties such as pH, organic matter content, soil texture, and Fe/Mn oxide content influence the mobility of HMs in contaminated soils. pH is the major factor determining metal mobility in the soil. Acid rain, high temperature, root exudation...
and microbial activities in the soil may increase the mobility of HMs in the soil by increasing acidity. Farm management practices such as cropping systems, tillage methods, and land use change also influences the mobility of HMs in the soil. Strategies for reducing HM mobility in contaminated farmlands include stabilization with organic and inorganic amendments, crop-hyper accumulating plant rotations, HM resistant microbes and soil washing. These techniques reduce bioavailability by removing the plant available fractions, converting mobile fractions to less available fractions and by changing contaminants to less mobile oxidation states.

**References**


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