

Short Communication

Would Forest Litter Cause a Risk of Increased Copper Solubility and Toxicity in Polluted Soils Remediated via Phytostabilization?

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Received: 30 August 2016

Accepted: 14 September 2016

Abstract

Soil solutions were collected from soils contaminated differently by the emissions from a copper smelter and incubated with beech litter. Five times in 30 days of incubation, soil solutions were acquired with MacroRhizon samplers and examined using the chemometric approach and two ecotoxicological assays: Microtox with *Vibrio fischeri* bacteria and Phytotoxkit with *Sinapis alba* seeds. Copper speciation in soil solutions was modeled in the Minteq program. Application of beech litter to soils resulted in a considerable increase of copper solubility. The toxicity of soil solutions was associated with total copper concentrations in solutions, copper speciation, and pH values. The toxicity measured by the Phytotoxkit test in soil solutions collected from non-amended soils was higher than that measured in Microtox, and was attributed to high concentrations of copper organic complexes in solutions. The application of beech litter, rich in dissolved organic carbon DOC, resulted in a radical increase of soil solution toxicity to both indicating organisms. This effect, particularly well expressed in the case of *V. fischeri*, was apparently caused by a decrease in pH and associated increase of total Cu concentrations in solutions.

Keywords: copper organic complexes, DOC, forest litter, Microtox, Phytotoxkit

Introduction

The most common method used for remediation of soils polluted with heavy metals, such as those surrounding a copper smelter, is immobilization of metals followed by land afforestation. Such a treatment may, however, create hazards associated with an increase of copper solubility in afforested areas caused by the influence of organic matter that derives from forest litter. Copper has a high affinity to organic matter [1-3] and can be complexed by low molecular weight organic acids produced from decomposing litter [4-6]. Moreover, microbiological decomposition of litter leads to acidification of the soil environment, which may directly contribute to enhanced desorption of copper into soil solution. Typically, the higher the concentration of copper in soil solution, the greater its toxicity to soil fauna and flora [7]. However, copper speciation also plays an important role in determining the toxicity of soil solutions. Therefore, simple chemical analyses may be insufficient for the assessment of environmental risk. A more advanced characteristic of contamination that involves speciation studies and recognition of toxicity associated with different forms of pollutants will be necessary for monitoring contaminated areas [8-11].

Several studies carried out in afforested areas contaminated with heavy metals proved that there is a risk of mobilization of copper and other metals from soils by organic acids derived from forest litter [12-13]. It is still unclear, however, if the mobile forms of heavy metals released into soil solutions derived mainly from a mineral matrix of contaminated soils or from metal-enriched forest litter itself.

The aim of this study was to determine the influence of forest litter that may be formed in afforested copper-polluted soils, on possible release of copper from soil solid phase, and to examine the time changes of concentrations, speciation, and toxicity of copper present in soil solutions.

Experimental

Five soils differently contaminated with heavy metals, mainly with copper, were collected from the close vicinity of the copper smelter Legnica – an area that was previously remediated by liming and phytostabilized by afforestation. Soil material was air dried, sieved, homogenized, and mixed with a beech litter (BL) obtained from an uncontaminated area. Basic properties of soils and litter were determined using the methods described by Tan [14] and reported in more detail elsewhere [15].

An incubation experiment was conducted over 30 days. Soil material (control soils) and soil material mixed with beech litter (50 g per 1 kg) were supplied with distilled water and incubated at moisture ca. 80% of field capacity. Soil solutions were obtained repeatedly in 1, 2, 7, 14, and 30 days of incubation using MacroRhizon samplers. The concentrations of copper and other major components of

soil solutions, including dissolved organic carbon DOC as well as pH, were determined by the methods described by Cuske et al. [15]. The toxicity of soil solutions was measured using two tests: a Microtox assay based on inhibition of *Vibrio fischeri* bioluminescence, and a Phytotoxkit assay based on inhibition of seed germination and root and shoot elongation, carried out with *Sinapis alba*. More detailed information on experimental methodology was provided elsewhere [15].

Cu speciation in soil solution was determined by Minteq v. 3.1 program. The SHM (Stockholm Humic Model) model was adopted for modeling the organic components of DOC in soil solutions.

Results and Discussions

Analyzed soils differed significantly in total content of heavy metals (especially copper) and the concentrations of dissolved organic carbon. In addition, their pH values differed considerably (Table 1). The beech litter contained 3.7 g kg⁻¹ of dissolved organic carbon (DOC), and indicated acidic reaction. The concentrations of heavy metals in the litter were negligible (Table 1).

Copper concentrations in soil solutions obtained from non-amended polluted soils, examined in previous studies, turned out to depend primarily on soil pH values, and then on total concentrations of Cu in soils [16-17]. The differences in DOC concentrations in non-amended soils were small, and therefore this factor had no significant impact on diversity of Cu concentrations in soil solutions of untreated soils (Figs 1-3) [17]. However, the concentrations of Cu in soil solutions of BL-amended soils were considerably higher in comparison to soil solutions of control soils (Figs 1-3). This effect may be explained by two mechanisms. First, a decrease in the pH values of soils after the application of acidic forest litter resulted in an intensive desorption of copper into soil solution [9, 18]. Secondly, released copper was bound by low molecular weight organic acids (formed as a result of litter decomposition in biochemical processes) and transformed into the complexed forms [4, 6, 19]. The highest

Table 1. Basic chemical properties of soils and forest litter applied

Material		C _{org.}	DOC	pH	Cu _{tot.}
		%	g kg ⁻¹	in 1M KCl	mg kg ⁻¹
Soils	1	0.90	0.85	7.38	5180
	2	0.76	0.80	6.41	3732
	3	1.30	1.80	5.22	1686
	4	0.66	1.40	7.00	531
	5	0.93	1.60	7.15	301
Beech litter	BL	34.6	3.7	5.5	25

concentrations of copper (15.4 mg L^{-1}) were found in soil solutions collected from soil No. 2 supplied with beech litter (Fig. 2) in the firsts days of incubation. The lowest concentrations of Cu ($0.11\text{-}0.17 \text{ mg L}^{-1}$) were observed in

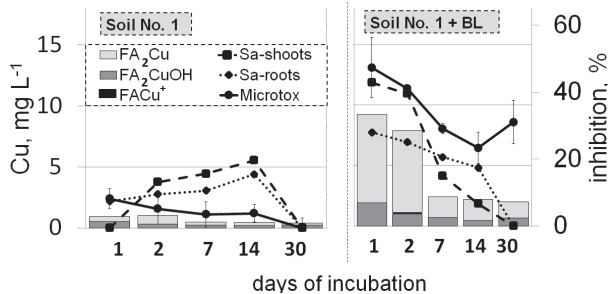


Fig. 1. Concentrations of Cu species and toxicity indices of soil solutions extracted from soil No. 1 – nonamended (left) and treated with a beech litter (right). Luminescence inhibition of *V. fischeri* in Microtox assay (Microtox) and inhibition of elongation measured in Phytotoxkit for shoots (Sa-shoots) and roots (Sa-roots) of *Sinapis alba* (right axis) are shown as related to the concentrations (mg L^{-1}) of predominant Cu species in soil solutions (left axis). FA (abbreviation of “fulvic acids”) stands for organic ligands in soluble Cu complexes (as modeled in Minteq). Error bars indicated for Microtox data represent a standard deviation.

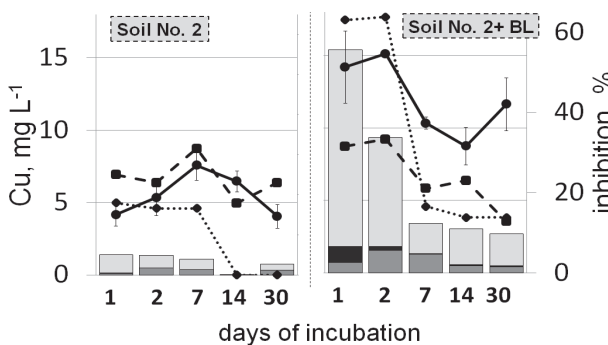


Fig. 2. Concentrations of Cu species and toxicity indices of soil solutions extracted from soil No. 2 – nonamended (left) and treated with a beech litter (right; see Fig. 1 for explanations).

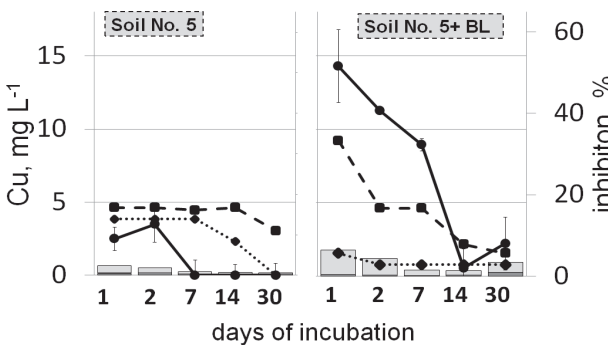


Fig. 3. Concentrations of Cu species and toxicity indices of soil solutions extracted from soil No. 5 – nonamended (left) and treated with a beech litter (right; see Fig. 1 for explanations).

soil solutions obtained from the least polluted, untreated soil Nos. 4-5 (Fig. 3) in the lasts days of incubation.

Almost 100% of copper present in soil solutions occurred in the forms complexed with low molecular organic matter, specified as FA_2Cu , FA_2CuOH , and FACu^+ , where the symbol FA stands for various soluble organic compounds, and represented in the SHM model by fulvic acids, which is confirmed by other studies [20]. The presence of free copper ions Cu^{2+} in soil solutions, in trace concentrations, was only confirmed in the case of soil Nos. 2 (Fig. 2) and 3, in the first and last days of incubation, when the pH values were, due to BL application, the lowest. In those cases, the cationic forms of free Cu^{2+} made up ca. 0.03% of soluble copper.

The concentrations of particular forms of copper complexed organically (FA_2Cu , FA_2CuOH , and FACu^+) in the solutions of control soils were dependent mainly on soil pH values. The predominating forms of copper complexes in soil solutions were non-ionic complexes FA_2Cu , the contribution of which ranged from 59.3 to 93.6% (Figs 1-3). The highest concentrations of these complexes were observed in soil solutions collected from soil Nos. 2-3 (with the lowest pH values; Fig. 2). In those solutions, the contributions of ionic FACu^+ complexes were higher compared to soil solutions collected from the other soils. In turn, the soil solutions of more alkaline soils (Nos 1, 4-5) contained higher contributions of non-ionic hydroxyl-copper complexes FA_2CuOH (Fig. 1). Applying beech litter to soils did not significantly affect the main patterns of copper speciation in soil solutions. However, the contributions of hydroxyl-copper complexes FA_2CuOH were in the case of BL-treated soils Nos. 1 (Fig. 1), 3, and 4 – significantly higher than in control soils. Additionally, in soil solutions of BL-treated soils, Nos. 2-3 (i.e., those with the lowest pH values; Fig. 2), the contributions of ionic copper complexes FACu^+ increased significantly compared to non-treated soils.

Total concentrations of copper in soil solutions collected from untreated soils, as well as from all BL-treated soils, tended generally to decrease with time, which may be explained by a so-called “aging effect” associated with stabilization of contaminants bound to transformed, stable organic matter [21]. In some cases in our experiment, i.e., in soil solutions of control soil No. 2 and BL-treated soil No. 5, an apparent, though statistically insignificant, increase of copper concentrations was observed after 30 days of incubation. Similar observations were also reported by several other authors, but the mechanisms involved in such an effect remain unclear and need further investigation [22-23].

The toxicity of soil solutions differed greatly in the experiment, depending on pH values of soil solutions, the time of incubation, and total concentrations of heavy metals (particularly of copper) in soils (Figs 1-3). The results obtained from two different ecotoxicological tests were different, depending clearly on the type of testing organism.

Soil solutions of untreated soil Nos 4-5 indicated the lowest toxicity, with the inhibition of *V. fischeri*

luminescence oscillating at the level of 10%, and the maximum inhibition of *Sinapis alba* roots and shoots elongation at 14% and 17%, respectively. Relatively low total concentrations of copper in these two soils, as well as their considerably high pH values, explained low concentrations of copper released into soil solutions [24] and their low toxicity. In fact, all soil solutions obtained from non-treated soils indicated toxicity below 20%, and according to a toxicity and hazard system for natural waters elaborated by Persoone et al. [25], would be classified as class I: “no acute hazard”.

The toxicity of soil solutions acquired from BL-treated soils, measured in both Microtox and Phytotoxkit assays, was definitely higher compared to the solutions of control, non-treated soils. Generally, a Phytotoxkit assay indicated higher sensitivity than Microtox in the case of soil solutions collected from control soils. On the contrary, soil solutions of BL-treated soils exhibited higher toxicity if measured in Microtox test compared to Phytotoxkit assay. It was undoubtedly associated with lower pH values of soil solutions (after the application of beech litter), to which *V. fischerii* bacteria are particularly sensitive [26]. Numerous studies have confirmed that the microorganisms used in the Microtox test are highly sensitive to the changes of pH and the presence of ionic forms of copper in solution [27-28]. In contrast, copper species complexed with low molecular organic matter are particularly intensively taken up by plants [15], which results in high sensitivity of Phytotoxkit assay to enhanced concentrations of organically complexed Cu in solution.

Applying soil solutions into incubation plates containing the seeds of *Sinapis alba* generally caused a higher inhibition of root elongation compared to the elongation of shoots. This observation stays in agreement with other studies that proved the particularly high accumulation of contaminants in early stages of plant growth in plant roots and a substantial reduction of root elongation compared to the shoots [29-30]. However, the results obtained in the case of soil solutions acquired in the first two days from BL-treated soil No. 2 were opposite; and the inhibition of shoot elongation was stronger compared to roots. It should be stressed that the toxicity of those solutions (BL-treated, No.2), measured in both tests, was the highest of all the solutions examined, and related pH values were the lowest (pH 6.5-7.1).

The toxicity of soil solutions, both of control soils and BL-treated soils, decreased with time. After the seventh day of incubation, a substantial decrease in the toxicity was observed in all treatments and was proved by all the tests. Such an effect was apparently associated with decreasing concentrations of copper in soil solutions and a slight increase of pH values. Exceptionally, a statistically insignificant increase of toxicity was noted in the case of soil solutions collected from some BL-treated soils in the last days of incubation. This effect was particularly well seen in the Microtox assay results, and should probably be attributed to the slight decrease in pH values over time.

Our previous studies, carried out with the same soils [15-17] showed that the concentrations of lead and zinc,

i.e., two other potentially toxic metals, in the soil solutions of control soils were low (in the ranges 0.01-0.09 mg L⁻¹ and 0.03-1.50 mg L⁻¹, respectively), and had no effect on the toxicity measured in both tests. This statement was confirmed by the results of a PCA analysis presented elsewhere [15].

Conclusions

The formation of forest litter and its decomposition in soils contaminated with heavy metals, in particular with copper, may be a cause of increasing solubility of metals, both in effect of their complexing by low-molecular-weight organic matter and due to a decrease of soil and soil solution pH values generated by microbiological decomposition of the litter.

Regardless of soil pH, the presence of decomposing forest litter may cause a release of copper from heavily contaminated soils into soil solution, which in turn will result in increased toxicity of soil solutions. This observation confirms the need for further research, including field studies, to examine the real environmental risk associated with the release of heavy metals from polluted soils due to their complexing with organic compounds derived from forest litter.

The highest toxicity of soil solutions, measured in the biotests with *Vibrio fischeri* and *Sinapis alba*, was observed in the first two days of incubation. Then, both the concentrations of Cu in soil solutions and their toxicity tended to decrease considerably with time. However, in the case of soils with relatively low pH values, a secondary slight increase of copper concentrations and the toxicity of solutions was observed after 30 days of incubation. Therefore, further studies on long-term changes of copper concentrations in soil solutions and their toxicity should be carried out with forest-litter-affected, highly contaminated soils.

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