The influence of old leather tannery district on chromium contamination of soils, water and plants

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ABSTRACT

The paper demonstrates the actual level of chromium contamination at the site of the old tannery waste lagoon in Serniki community (SE Poland). Despite the fact that since 1994 this area is not exploited, the chromium presence is still detected in soil, water and plants. The current study is the 4th phase (after 1989, 2001 and 2004) of checking and controlling the chromium concentration in this territory. The chromium content was assessed after soil and plant mineralization using FAAS method. Total chromium concentrations in soil samples reached values of 2.25 - 18.25 mg·kg\(^{-1}\) dry mass. Levels of chromium in water samples ranged between 0.008 and 0.017 mg·dm\(^{-3}\) for Wieprz River and the trough, situated in the tannery centre, respectively. Absorption of chromium by plants depended from the plant species. The highest Cr content in the stalks of *Oenothera biennis*, whilst the lowest in *Phragmites australis* were detected. Finally, it was assumed that soil chromium content noted in 2009 was by 42% - 83% lower in relation to the data from 2004. Consequently, nowadays the tannery waste lagoon site is not so threat to the health of local residents and to the surrounding ecosystem.

Keywords: Chromium; Soil Pollution; Tannery Wastes; Chromium Accumulation

1. INTRODUCTION

Chromium is a heavy metal which presence in soil, water and atmosphere can be hazardous to the natural environment [1-3]. Therefore, knowledge of the transport and transformation of this element in the soil environment is important for predicting threats from its presence [4].

Chromium is commonly used in various industries such as metallurgical, refractory, chemical pigments, electroplating and tanning [5,6]. One of the major emerging problems of the tanning industry is the disposal of chromium-contaminated sludge that is produced as by-product of the wastewater treatment [5]. The world production of chromium is in the order of 10\(^7\) tons per year; 60% - 70% is used in alloys, including stainless steel, and 15% is used in chemical industrial processes, mainly leather tanning, pigments and electroplating [7,8].

The migration of chromium in soils and surface- or ground-waters, from its source of pollution, and its toxicity mainly depends on its speciation [6]. Its commonly existing forms are trivalent and hexavalent chromium [2,3]. In oxidized Cr (VI) form, chromium is highly soluble in water, and therefore mobile, whereas the reduced Cr (III) form is almost insoluble in water and thus immobile [4,9,10]. Hexavalent chromium is very toxic, and it can be an inhaled carcinogen, poisonous to humans and other mammals, whereas trivalent chromium is an essential nutrient as mineral supplement [1,3,9,11].

Values of chromium are strongly related to pH and redox potential (Eh) conditions [4]. Most of authors demonstrated that Cr (VI) is stable in the more oxidising environment with Eh level above 600 mV and pH above 6.0 [1,12,13]. Under conditions of pH 3 to 7 and Eh from -200 to 500 mV compounds of Cr (VI) tend to reduce to more thermodynamically stable Cr(OH)\(_3\) [4,14]. So, under appropriate pH and Eh conditions as well as in the presence of reducing or oxidising agents one form of chromium can transform to another [1,3]. The reduction of highly toxic Cr (VI) to the less toxic Cr (III) is likely to be useful for the remediation of contaminated waters and soils [9]. It was also indicated that Cr (III) can be transformed to Cr (VI) mainly inside root cells, and/or in the aerial part of the plant, however, roots accumulate 10 - 100 times more chromium than shoots and other plants tissues [7,8]. As a result of this, there is observed an inhibition of growth, respiration activities and photosynthesis [8,9]. Also progressive stages of chlorosis and necrosis are the symptoms of chromium toxicity in plants [7]. There are however still conflicting results regarding,
which form of chromium is taken up and accumulated by plants [7,8,15]. Some authors considered that both translocation as accumulation of chromium inside the plant depends on the oxidation state of the supply and/or on the plant species [8,15,16].

Yet, among heavy metals, chromium has been paid much less attention compared to, e.g., cadmium copper, zinc, lead etc. One of reasons is that this element is constantly considered as a “local source” contaminant, whilst the second is that the dominant naturally occurring form of chromium, Cr₂O₃, is known essentially immobile and non-toxic in the environment [10].

Thus, based on above presented assumptions we have started a monitoring program aiming to assess the actual chromium accumulation in the soils, waters and plants of a tannery waste lagoon vicinity in SE Poland.

Our interest was also connected with chromium transformations in soils in the pH and Eh conditions, described in the present tannery waste lagoon site. By realized measurements of pH and Eh in the soil samples the estimation of the dominant form of chromium was possible.

2. MATERIALS AND METHODS

2.1. Site Location and Description

In the current study the tannery waste lagoon (6 ha) in Lubartów (SE part of Poland), located in Serniki community (51°26’N 22°40’E) has been studied. The storage yard is situated both in a basin as in the direct neighbourhood of the Wieprz River (approx. 50 m away) and cultivated fields [1]. In the active period (1970-1994) more than 200,000 t of tannery waste, containing 5% - 10% of Cr₂O₃ were produced [6]. High chromium level in wastes deposit results from the application of chromal Cr(OH)SO₄ (chromium (III) hydroxysulfate), used for protein denaturation in the amount of more than ten thousand tons per year. The second problem was the fact that the tannery waste lagoon was not appropriately secured. Its bottom was not sealed, and this still causes problems connected with the possibility of leaching the harmful substances getting away from it. Consequently, even at present (15 years after closing the Lubartów tannery) a migration of chromium from deposited wastes to the soil and ground water has taken place.

Lubartów tannery was declared insolvent on September 1998. During past 10 years the tannery waste lagoon has undergone reclamation twice (in 2001 and 2005) by covering the top of the storage layer by rocks originating from Bogdanka coal mine. Phytoremediation method was also applied by introducing pine plantings.

In 1989 the old tannery waste lagoon site was first examined for chromium contamination. After 10 years the level of chromium contents in soils, waters and plants were re-analysed [17]. Five years later (in 2004) environmental samples (soils, plants and water) were investigated again [6]. So, the current study is the fourth phase of monitoring the chromium concentration at this site.

2.2. Samples Collection

The study material was collected in October 2009 at different distances from the tannery dumping ground (Table 1).

The soil sampling points are numbered from 1 to 7, depending from their location (Figure 1). Soil samples were taken from the surface layer (0 - 20 cm).

Plant samples: leaves of Quercus petraea (DB) and stalks of Oenothera biennis (WD), Juncus sp. (S) and Phragmites australis (TP), representatives for this place were also tested. Water samples were collected from the Wieprz River and the trough situated in the centre of waste deposit (see Figure 1).

2.3. Mineralization of Soil and Plant Samples

Under laboratory conditions the soil samples were

![Figure 1. Aerial view of the tannery waste lagoon with marked sample points (maps.geoportal.gov.pl).](image-url)
dried at 105°C for 24 h and after that sifted and weighed. Both soil as plant samples were mineralized. Samples (0.5 g) were digested using the mixture of concentrated HNO₃ and HF and performed (4 min, temp. 150°C) in open microwave system (Maxidigest MX 4350, Prolabo). After filtration the extracts were diluted with distilled water to the final volume of 50 ml.

2.4. Total Chromium Concentration Measurements

Environmental samples (water, plants and soils) were analysed at λ = 359.3 nm for total chromium concentration by AAS technique, with flame atomisation—FAAS (Z-8200, HITACHI), in three replications. Total chromium concentration was expressed as element content in dry mass of the soil (mg·Cr·kg⁻¹).

2.5. Determination of pH and Eh

The pH and Eh were determined potentiometrically using a multifunctional potential meter pIONneer 65 (Radiometer Analytical S.A., France), equipped with a following electrodes: glass pH electrode—(Cartridge pH E16M340) and combined platinum and Ag/AgCl (reference) electrode (E31M004)—for Eh. Each measurement was replicated three times.

3. RESULTS AND DISCUSSION

3.1. pH and Eh Values in the Soil Samples

Values of pH and Eh in the soil samples taken from different points of the old tannery waste lagoon are presented in Figure 2. Generally, Eh level varied between 266 - 510 mV, reaching the highest values in soil sample 5—taken from neighbouring cultivated field.

The lowest redox potential level (266 - 281.3 mV) in soil sample 6—situated in the central point of the lagoon and sample 7, located nearby Wieprz River was noted.

Soil pH represented acidic conditions (2.7 - 5.4), characteristic for sandy soils with small CEC (cation exchange capacity) and high H⁺ saturation. Sandy soils predominate at investigated tannery waste lagoon area [1]. Moreover, decrease of pH to the value of 2.6 resulted in Eh increase to 510 mV. This trend is compatible with earlier observations [1,4,18,19]. Presented range of Eh is characteristic for good oxygenation conditions with O₂, as a main electron acceptor. Furthermore, these Eh values and acidic pH favoured the dominance of less toxic and immobile Cr (III) form. It was also pointed out that the main oxidant responsible for oxidation of Cr (III) in soil was manganese (IV) oxide [4]. Authors observed that the sorption of Cr (III) on soil decreased in the presence of MnO₂, what is probably caused by comparative reaction, which is oxidation of Cr (III) to Cr (VI) by MnO₂ and also by process of manganese (III) sorption on the soil [4].

3.2. Chromium Contamination in Water Samples

Total chromium concentration in water sample originated from Wieprz River reached 0.008 mg·dm⁻³ and 0.017 mg·dm⁻³ in the sample taken from the trough situated in the tannery centre. Registered in 2009 water contamination was by 98% lower from its level in Wieprz River (0.41 mg·dm⁻³) noted in 2004 [6]. Additionally, in 1989, the chromium concentration in water was estimated at the level of 1.18 mg·dm⁻³ [1]. Decidedly, in comparison with earlier studies a decreasing trend of the chromium content in Wieprz River was found.

3.3. Chromium Contamination in Plants

The highest total chromium concentration was found in the following plant samples (see Figure 3): stalks of Oenothera biennis (WD) and leaves of Quercus petraea (DB), where Cr content was on the level: 11 mg·kg⁻¹ and 5.5 mg·kg⁻¹ for WD and DB, respectively. In the case of
stalks of *Juncus sp.* (S) and *Phragmites australis* (TP) less ability for chromium absorption was noted, as level of Cr concentrations remained at the range of 2.75 - 3.0 mg·kg⁻¹. It was assumed that only a small amount of metal passes through plants [20], however one should remember that the mechanisms of chromium resistance in plant cells have not been properly discovered and described [7]. Suggestion of World Health Organisation (WHO) indicated that the chromium concentration in plants growing in unpolluted soil should not exceed 0.19 mg·kg⁻¹ [21]. Other authors reported [10] that in normal conditions, chromium concentration in plants is less than 1 µg·g⁻¹, but in plants growing on soils treated with tannery waste, concentrations may increase significantly [22]. The study by Isikli *et al.* [20], who attempted to determine the chromium content in the plants taken from a rural area, exposed to cement factory emissions, indicated that an average chromium concentration was c.a. 167 mg·kg⁻¹. It was also observed that in serpentine soils the mean chromium content was 45 mg·kg⁻¹, but plants growing on these soils may contain up to 100 mg·Cr·kg⁻¹ [23]. In that context a comparison of our data with these mentioned above demonstrates that tested plant samples were not as highly contaminated with chromium. It may be an effect of phytoremediation treatments realized in 2005 on investigation site. It is known, that the application of plants to decontaminate soils is an initial stage for detoxification of heavy-metal-polluted areas [7]. Phytoremediation based on the ability of the some plant species to absorb, sequester, an/or transform the soil contaminant [24], can be very efficient, particularly in soils with small CEC.

### 3.4. Chromium Contamination in Soil Samples

Large differences in soil chromium concentration were observed in the site investigated, with a very scattered distribution. Total chromium concentrations in soil samples are presented in Figure 4. The highest chromium content is still present in sandy soil samples 1 (18.25 mg·kg⁻¹) and soil sample 2 (8.0 mg·kg⁻¹), located in the centre of the old tannery waste lagoon (see Figure 1). At the remaining soil sample points (3 - 7) chromium concentration not exceeded the level of 5 mg·kg⁻¹.

Finally, comparison of the total Cr concentration changes (from 2003 to 2009) in a investigated soil samples were done (Figure 5). A decreasing trend of the chromium concentration with time in the monitored soil points was stated. The highest chromium content was noted in 2004, what may be connected with the second step of tannery waste site restoration—realized in 2005. Results from 2009 showed that chromium concentration in soil samples was lower from these in 2004 by 42% - 83% (depending from the soil point location).

In normal conditions, chromium in soils derives from that content in the parent material [10]. Significant chromium enrichment in soils is always of anthropogenic origin. As is reported by WHO [21] the chromium content in soils ranges between 2 and 50 mg·kg⁻¹. The results obtained in the current study have not exceeded the acceptable level of 50 mg·kg⁻¹. Even, the highest chromium concentration stated in the centre of the old tannery waste lagoon (18.25 mg·kg⁻¹), was by 63.5% lower from WHO limit. Decreasing trend of chromium content observed during last 10 years in the investigated site of tannery lagoon is optimistic, taking into account that the residence time of chromium in soils is estimated to be 10³ - 10⁴ years [10,25].

### 4. CONCLUSION

Our investigations demonstrated that total chromium concentration in soil samples from the vicinity of tannery waste lagoon ranged from 2.25 to 18.25 mg·kg⁻¹·dm and not exceeded Polish acceptable level (50 mg·kg⁻¹). Similarly, water contamination in the range from 0.008 - 0.017 mg·dm⁻³ was not an infringement acceptable.
standard \((0.05 \text{ mg} \cdot \text{dm}^{-3})\). Noted in the current experiment the average concentration of chromium in soil landfill \((5.5 \text{ mg} \cdot \text{kg}^{-1})\) was about half lower, in comparison with the concentrations noted in 2003 \((9.7 \text{ mg} \cdot \text{kg}^{-1})\) and 2004 \((13.8 \text{ mg} \cdot \text{kg}^{-1})\). The present study confirmed also previous findings that the chromium absorption from the soil by plants depends on the plant species, as the highest Cr content was detected in the stalks of \textit{Oenothera biennis} \((11 \text{ mg} \cdot \text{kg}^{-1})\), whilst the lowest in \textit{Phragmites australis} \((3 \text{ mg} \cdot \text{kg}^{-1})\). Determined soil redox potential (Eh) values, not exceeding the level of 600 mV and pH 2.7 - 5.4 reflected the ranges characteristic and favourable for Cr (III) dominance. Moreover, it was assumed that applied reclamation methods seemed to be an effective in reduction of Cr contamination, as nowadays the tannery waste lagoon site is not so threat to the health of local residents and to the surrounding ecosystem.

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