Phytoremediation of Soil Contaminated by Organochlorine Pesticides and Toxic Trace Elements: Prospects and Limitations of Paulownia tomentosa

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Abstract: Paulownia tomentosa (Thunb.) Steud. is a drought-resistant, low-maintenance and fast-growing energy crop that can withstand a wide range of climatic conditions, provides a high biomass yield (approximately 50 t DM ha$^{-1}$ yr$^{-1}$), and develops successfully in contaminated sites. In Kazakhstan, there are many historically contaminated sites polluted by a mixture of xenobiotics of organic and inorganic origin that need to be revitalised. Pilot-scale research evaluated the potential of P. tomentosa for the phytoremediation of soils historically contaminated with organochlorine pesticides (OCPs) and toxic trace elements (TTEs) to minimise their impact on the environment. Targeted soils from the obsolete pesticide stockpiles located in three villages of Talgar district, Almaty region, Kazakhstan, i.e., Amangeldy (soil A), Beskainar (soil B), and Kyzylkairat (soil K), were subjected to research. Twenty OCPs and eight TTEs (As, Cr, Co, Ni, Cu, Zn, Cd, and Pb) were detected in the soils. The phytoremediation potential of P. tomentosa was investigated for OCPs whose concentrations in the soils were significantly different (aldrin, endosulfans, endrin aldehyde, HCB, heptachlor, hexabromobenzene, keltan, methoxychlor, and γ-HCH) and for TTEs (Cu, Zn, and Cd) whose concentrations exceeded maximum permissible concentrations. Bioconcentration (BCF) and translocation (TLF) factors were used as indicators of the phytoremediation process. It was ensured that the uptake and translocation of contaminants by P. tomentosa was highly variable and depended on their properties and concentrations in soil. Besides the ability to bioconcentrate Cr, Ni, and Cu, P. tomentosa demonstrated very encouraging results in the accumulation of endosulfans, keltan, and methoxychlor and the phytoextraction of γ-HCH (TLFs of 1.9–9.9) and HCB (BCFs of 197–571). The results of the pilot trials support the need to further investigate the potential of P. tomentosa for phytoremediation on a field scale.

Keywords: Paulownia tomentosa; phytoremediation; organochlorine pesticides; toxic trace elements; bioconcentration factor; translocation factor

1. Introduction

Emissions from the oil and gas industry and mining and smelting activities are sources of various hotspots [1–4] that negatively affect the surrounding soils and surface and groundwaters. At such contaminated sites, xenobiotics of organic and inorganic origin occur simultaneously, requiring the development of innovative management strategies to address these complex issues. Only a limited number of studies have proposed solutions for addressing these complex ecological problems [1–3], and phytoremediation is among the proposed approaches [5–7].
Phytoremediation is based on the utilisation of different plants for soils contaminated by toxic trace elements (TTEs), hydrocarbons, pesticides, oil products, and radionuclides [8–13]. The plants used in phytoremediation must grow easily and quickly, and produce enough biomass to eliminate notable amounts of contaminants from the soil. Undoubtedly, the nature of remediation depends directly on the ability of the plant to uptake and accumulate the concerned pollutants. In the case of energy crops proposed as phytoagents [14,15], one of the crucial characteristics is the high content of fibre fractions (i.e., lignin, cellulose, and lignocellulose), which corresponds to the quality of the bioproducts to be produced.

Gołab-Bogacz et al. indicated [16] Miscanthus spp. as the most promising plant suitable for phytoremediation. These species have a high annual biomass yield (up to 36.6 t ha\(^{-1}\)) [17] and the highest heating value of 16.3–20.2 MJ kg\(^{-1}\) [18]. Another prospective energy crop, Arundo donax, has been shown to produce even higher biomass yields than M. × giganteus, as it can be harvested several times a year [19]. However, considering the energy costs, M. × giganteus remains a more promising plant as it requires fewer inputs during cultivation [20,21]. Numerous studies have demonstrated the efficacy of M. × giganteus and A. donax in the phytoremediation of TTE-contaminated soils [12,22–26]; however, when the soil was contaminated with organochlorine pesticides (OCPs), M. × giganteus could not tolerate their high concentrations and survived in the soil, overcoming the maximum permissible concentration (MPC) for Kazakhstan (241 ± 16 µg kg\(^{-1}\)) in two instances only [27], while studies investigating the remediation potential of A. donax with respect to OCPs were not found. Indeed, M. sinensis, another representative of Miscanthus spp., showed good growth in OCP-contaminated soil, with concentrations up to 62 × MPC [27]. However, the cultivation of M. sinensis and A. donax is restricted in some countries due to their invasiveness [28,29], which shortens the list of plants eligible for the phytoremediation of soils contaminated with OCPs and TTEs and necessitates the search for new plants that can be used.

In this context, Paulownia tomentosa (Thunb.) Steud [30] can be considered a good candidate for the phytoremediation of soil contaminated with a mixture of OCPs and TTEs. The plant can produce consistent biomass of 50 t DM ha\(^{-1}\) y\(^{-1}\) under non-optimised growth conditions and up to 330 t DM ha\(^{-1}\) y\(^{-1}\) under optimised ones [30,31], which is around two times higher compared to other popular woody plants: willow (23.1–25.7 t DM ha\(^{-1}\)) and poplar (12.2 ± 0.3 t DM ha\(^{-1}\)) [32,33]. Moreover, P. tomentosa is a drought-resistant, low-maintenance plant able to withstand a wide range of climatic conditions (i.e., from −20 to +40 °C) [34]. The biomass content of P. tomentosa is similar to that of Miscanthus sp., i.e., 22.4% lignin, 37.4% cellulose, 33.3% hemicellulose, and 61.5–70.7% holocellulose [30,35], which makes its biomass promising for processing into various bioproducts. The published research on the application of P. tomentosa is mainly concerned with TTE-contaminated soils [36–40]; there are also a few reports on the use of the plant in PCB- and hydrocarbon-contaminated soils [3], while this plant’s tolerance to OCP-contaminated soils is not presented in the literature, although complex contaminated soils are common in the territories of former obsolete pesticide stockpiles [41], mining [42], and post-military soils [27].

The current study aimed to investigate the phytoremediation efficiency of P. tomentosa toward soil historically contaminated by a mixture of OCPs and TTEs and evaluate the interconnections between the background of the aged soils and the plant phytoremediation potential. The use of historically contaminated soils allowed us to conduct experiments in the conditions of the natural heterogeneity of the compounds’ distribution in the soils.

2. Materials and Methods

2.1. Soil Collection

The research soils were sampled on April 2018 at historically contaminated sites in the vicinity of obsolete pesticide storage facilities located in three villages of Talgar district, Almaty region, Kazakhstan: (a) Amangel’dy (GPS 43°18′01.54″ N, 77°12′33.9″ E), hereafter
denoted as soil A; (b) Beskainar (GPS 43°13′16″ N, 77°6′49″ E), soil B; and (c) Kyzylkairat (GPS 43°17′58.7″ N, 77°11′39.6″ E), soil K. According to the updated Köppen–Geiger classification [43], the climate of the Talgar region belongs to group Dfa: it has cold winters, hot summers, and does not have a dry season.

Soil sampling was done according to the standard procedure (ISO 18400-2018) [44]: five samples were collected from a 5 × 5 m testing square at a depth of 0–0.6 m. After sampling, plant materials and stones were manually removed; soil was further air-dried until constant weight and sieved (d = 3 mm). According to the World Reference Base for Soil Resources Classification, the soils at the three research sites belonged to kastanozems [45].

The agrochemical parameters of the research soils were determined using standard methods. The total humus content (C) was measured by the Tyurin method [46]; the content of phosphorus (P$_2$O$_5$) and potassium (K$_2$O) mobile forms by the Kirsanov method with the modification of the Central Research Institute for Agrochemical Agricultural Services [47]; the absorbed bases of sodium (Na) and potassium (K) by the Antipov-Karataev and Mametov method with Grabarov modification; the absorbed bases of calcium (Ca) and magnesium (Mg) by the Arinushkin method with Grabarov modification; and the soil pH according to GOST 26423-85 [48]. The agrochemical parameters of the research soils are presented in Table 1. These soils were used in the pot experiment performed in greenhouse conditions.

Table 1. Agrochemical profile of the research soils.

| Parameter | Unit | Soil A | Soil B | Soil K |
|-----------|------|--------|--------|--------|
| C         | %    | 4.44 ± 0.11 | 5.27 ± 0.10 | 6.10 ± 0.02 |
| pH (water)|      | 7.48 ± 0.01 | 7.49 ± 0.02 | 7.85 ± 0.02 |
| P$_2$O$_5$| mg kg$^{-1}$ | 353 ± 15 | 71 ± 0 | 400 ± 5.0 |
| K$_2$O    | mg kg$^{-1}$ | 965 ± 15 | 740 ± 0 | 885 ± 25 |
| Ca        | meq/100 g | 16.4 ± 0.50 | 19.2 ± 0 | 20.8 ± 0.70 |
| Mg        | meq/100 g | 7.75 ± 1.40 | 5.05 ± 0.72 | 2.70 ± 1.23 |
| Na        | meq/100 g | 0.16 ± 0 | 0.16 ± 0 | 0.38 ± 0.01 |
| K         | meq/100 g | 1.31 ± 0 | 0.80 ± 0.03 | 1.04 ± 0.03 |

2.2. Experiment Design

The clones of _P. tomentosa_ were first obtained by in vitro propagation and further adapted to the open ground conditions. The adaptation took place for three months (January–March 2019), when the clones were illuminated with fluorescent lamps LB-40-4 of the infrared spectrum at 3000 lux. After the adaptation period, _P. tomentosa_ seedlings were planted in pots, and each pot was filled with 350 g of dried research soil. The surface area of one pot was 88.56 cm$^2$; consequently, the total area was 0.74 m$^2$. Overall, 84 seedlings were planted, 28 plants per research soil. The experiment was started on 18 March 2019 and ended on 15 June 2020. Soil moisture was adjusted to 50% by irrigation with regular cold tap water every 3rd day.

2.3. Biomass Collection at Harvest

Research plants were harvested on 15 June 2020. The plant’s roots and aboveground biomass (AGB) were sampled following GOST 17.4.4.02-84 [49]. Root samples were taken by unearthing the plant together with the soil from the pot. The roots were shaken free of the soil and washed thoroughly under cold running tap water to eliminate tiny soil particles and then oven-dried till constant weight. Plants’ AGB was dried in the same way as roots. The samples were separately collected in labelled plastic zip-lock bags and then stored at room temperature until the chemical analysis.

2.4. Chemical Analysis

Concentrations of OCPs were measured by gas chromatography with an electron capture detector (Gas Chromatography 6890N Agilent Technologies, Santa Clara, CA, USA)
equipped with the autosampler Combi-PAL (CTC Analytics AG, Zwingen, Switzerland). Limit of detection (LOD) values for soil and plant samples were 0.1 and 25 µg kg\(^{-1}\), while limit of quantification (LOQ) values were 4.0 and 5.0 µg kg\(^{-1}\), respectively. Quartz sand and cellulose were used as reference samples according to standards ST RK 2131-2011 [50] and ST RK 2011-2010 [51], used to analyse soil and plant samples, respectively. The TTE concentrations were determined by atomic absorption spectrometry with electrothermal atomisation, using a Varian AA240 Atomic Absorption Spectrometer GTA 120 (Agilent Technologies, Santa Clara, CA, USA). The reference samples were the same as for OCP content, while LOD and LOQ values were 0.1 and 2.0 mg kg\(^{-1}\), respectively. The procedure was described in detail earlier [52]; briefly, analysis of soil samples was provided according to standards ST RK ISO 11047-2008 [53] and GOST 23581.8-79 [54]; analysis of plant samples was performed following ST RK ISO 11047-2008 [53], GOST 23581.8-79 [54], GOST 26930-86 [55], and GOST 30178-96 [56].

2.5. Calculation of Phytoremediation Coefficients

The bioconcentration factor (BCF) is the ratio between the pollutant concentration in the plant tissue and its concentration in the soil. The coefficient was calculated according to Zayed et al. [57]:

\[
BCF = \frac{\text{Contaminant concentration in plant tissues (mg kg}^{-1}\text{) at harvest}}{\text{Initial contaminant concentration in soil (mg kg}^{-1}\text{)}}
\]  

(1)

The transfer of pollutants within the plant was quantified by the translocation factor (TLF), which is the ratio between the pollutant’s concentration in the aboveground biomass (AGB) (leaves and stems) and root system [58]:

\[
TLF = \frac{\text{Contaminant concentration in aboveground biomass (mg kg}^{-1}\text{)}}{\text{Contaminant concentration in roots (mg kg}^{-1}\text{)}}
\]  

(2)

2.6. Statistical Analysis

Data analysis was performed using RStudio software (version 1.3.959, R Studio PBC, 2020). A one-way ANOVA was performed to compare the initial concentrations of contaminants in the soils at three research sites, while a two-way ANOVA was applied to compare the contaminant concentrations in the AGB and roots of the plants grown in the different soils. In the case of TTEs, to attain statistical and biological differences, concentrations of these substances were measured in plant tissues, referring to the soils with the highest and lowest concentrations for each particular element. The comparison of BCF and TLF values was carried out using two-way ANOVA.

Tukey HSD tests were performed for pairwise comparison of means when ANOVA showed a significant effect of the tested factors. Then, treatments were categorised by letter in descending order, and boxplots/graphs were generated. Significance was declared at \( p < 0.05 \); however, tendencies at \( p < 0.10 \) were indicated as well.

3. Results

3.1. Contamination of the Research Soils

The levels of contamination of the research soils A, B, and K by OCPs are presented in Table 2. Altogether, twenty pesticides were detected in the soils, including fourteen insecticides, three metabolites, three fungicides, and one herbicide. The concentrations of TTEs in the research soils are presented in Table 2. Due to the natural heterogeneity of the distribution of organic compounds in the complex soil matrix, the OCP contamination appeared more heterogeneous than the distribution of TTEs. To address this heterogeneity, the phytoremediation process was examined for OCPs whose concentrations in the soils were significantly different (\( p < 0.05 \)), i.e., aldrin, endosulfans, endrin aldehyde, heptachlor,
hexabromobenzene, methoxychlor, and lindane (γ-HCH), or tended to be different (i.e., \( p < 0.1 \)), i.e., HCB and keltan.

Table 2. Concentrations of OCPs and TTEs in research soils.

| Contaminant                     | Pesticide Type a | MPC b,c | Soil A | Soil B | Soil K | \( p \)-Value | Root MSE |
|---------------------------------|------------------|---------|--------|--------|--------|--------------|----------|
|                                 | \( \mu g \text{ kg}^{-1} \) |         |        |        |        |              |          |
| Aldrin                          | I                | 2.5     | 12.2 b | 96.0 b | 345.2 a | <0.01        | 59.7     |
| Chlordane                       | I                | 100     | 30.1   | <LOD   | 72.1   | 0.34         | 47.1     |
| Chlorobenzilate                 | I                | 20      | 277.6  | 5590   | 32,242 | 0.45         | 31,134   |
| DDD                             | I                | 100     | 1153   | 2976   | 25,506 | 0.44         | 24,241   |
| DDE                             | I                | 100     | 9709   | 69,847 | 777,967| 0.40         | 716,310  |
| DDT                             | I                | 100     | 1237   | 6274   | 10,023 | 0.33         | 6613     |
| Dibutyl chlorendate             | H                | -       | 511.1  | 1285   | 2135   | 0.33         | 1208     |
| Dieldrin                        | I                | 0.5     | 42.3   | 291.3  | <LOD   | 0.18         | 185      |
| Endosulfans                     | ml               | 100     | 83.2 b | 124.1 b| 759.2 a| <0.001      | 63.0     |
| Endosulfan sulfate              | ml               | -       | 654.5  | 265.7  | 356.0  | 0.46         | 373      |
| Endrin                          | ml               | 1       | 1289   | 181.3  | 44,085 | 0.33         | 42,462   |
| Endrin aldehyde                 | ml               | -       | 62.4 b | 130.8 a| 1088 a | <0.05       | 394      |
| HCB                             | F                | 500     | 21.3   | 41.6   | 14.0   | 0.07         | 11.7     |
| Heptachlor                      | I                | 50      | <LOD   | 118.4 a| 269.0 a| <0.001      | 17.1     |
| Heptachlorepoxide               | F                | 50      | 190.3  | <LOD   | 3029   | 0.39         | 3580     |
| Hexabromobenzene                | F                | 30      | 39.8 c | 187.6 b| 604.0 a| <0.001      | 54.0     |
| Keltan                          | I                | 100     | 11.9   | 22.1   | 32.9   | 0.10         | 10.7     |
| Methoxychlor                    | I                | 1600    | 11.1 c | 137.2 b| 1307 a | <0.001      | 43.9     |
| γ-HCH                           | I                | 100     | 19.3 b | 20.1 b | 76.4 a | <0.001      | 3.0      |
| HCH isomers                     | ml               | 100     | 162.7  | 258.9  | 600.4  | 0.25         | 299.2    |

| Contaminant                     | Pesticide Type a | MPC b,c | Soil A | Soil B | Soil K | \( p \)-Value | Root MSE |
|---------------------------------|------------------|---------|--------|--------|--------|--------------|----------|
|                                 | \( \mu g \text{ kg}^{-1} \) |         |        |        |        |              |          |
| Cr                              | 6                | 1.53 a  | 1.12 b | 0.98 b | <0.01 | 0.12         |          |
| Co                              | 5                | 1.84 a  | 1.89 a | 1.35 b | <0.001| 0.06         |          |
| Ni                              | 4                | 2.36 a  | 1.85 c | 2.08 b | <0.001| 0.08         |          |
| Cu                              | 3                | 4.93 a  | 4.34 ab| 4.28 b | <0.05 | 0.26         |          |
| Zn                              | 23               | 36.07 a | 7.99 b | 12.07 b| <0.001| 4.15         |          |
| As                              | 2                | 0.32 b  | 0.67 a | 0.27 b | <0.001| 0.06         |          |
| Cd                              | 0.5              | 2.17 a  | 1.17 b | 0.85 b | <0.001| 0.23         |          |
| Pb                              | 32               | 5.01 b  | 2.25 b | 11.1 a | <0.01 | 2.12         |          |

\( a \) F: fungicide; I: insecticide; H: herbicide; ml: a metabolite of insecticide; \( b \) MPC values for OCPs as for the Republic of Kazakhstan [59]; \( c \) MPC values for TTEs, as for the Republic of Kazakhstan [60]; LOD = 0.1 \( \mu g \text{ kg}^{-1} \). The concentrations of Cr, Co, Ni, Cu, and Zn reflect the mobile form, i.e., the fractions available to plants, while the concentrations of As, Cd, and Pb represent the total form. Means in the same line with different letters are significantly different.

Among the three research soils, soil K appeared to be the most contaminated with OCPs, followed by soils B and A, while contamination with TTEs showed the opposite tendency: soil A was the most contaminated, followed by soils B and K. Concentrations of aldrin and hexabromobenzene exceeded the MPC in research soils—specifically, aldrin by 4.9, 38.4, and 138 times, and hexabromobenzene by 1.3, 6.3, and 20.1 times in soils A, B, and K, respectively. Endosulfan concentrations slightly exceeded the MPC in soil B (by 1.2 times), while the exceedance in soil K was around 7.6 times. Heptachlor was detected only in soils B and K at concentrations exceeding the MPC by 2.4 and 5.4 times, respectively. The concentrations of methoxychlor and γ-HCH varied considerably between soils; however, they did not exceed the MPC in any of them. The concentration of HCH isomers exceeded the MPC in all three soils, consequently, by 1.6, 2.6, and 6.0 times in soils A, B, and K, respectively (Table 2). The concentrations of TTEs in the research soils varied essentially (Table 2); however, MPC values were surpassed for Cu, Zn, and Cd only. Specifically, the Zn concentration in soil A was 1.6 times higher than the MPC, and the Cu concentration was higher than the MPC by 1.6, 1.4, and 1.4 times in soils A, B, and K, respectively. The highest exceedance...
was recorded for Cd, which exceeded the MPC by 4.3, 2.3, and 1.7 times for soils A, B and K, respectively.

3.2. Phytoremediation Potential of *P. tomentosa* Utilised in Complex OCP- and TTE-Contaminated Soils

To assess the potential of *P. tomentosa* to uptake the contaminants from the research soils, the phytoremediation coefficients BCF and TLF were calculated (Table 3). The patterns of transfer of two OCPs in the soil–plant system did not allow further analysis: heptachlor was detected only in the AGB (BCF of 2.8), while hexabromobenzene only in the roots (BCF of 2.0) of plants grown in soil B. Statistical analysis of OCP concentrations in plant tissues showed the impact of soil contamination on the uptake of endosulfans and endrin aldehyde. Similarly, the cumulative influence of contaminants’ concentrations in soils and the accumulation organs of plants was detected for HCB, keltan, methoxychlor, and γ-HCH (Table 3).

Table 3. OCPs and TTEs concentrations in AGB (sum of leaves and stems) and roots of *P. tomentosa*. Different letters within one compound indicate a significant difference.

| Contaminant       | Soil A     | Soil B     | Soil K     | SO Effect | p-Value | Cumulative Effect | Root MSE |
|-------------------|------------|------------|------------|-----------|---------|------------------|----------|
|                   | AGB | Roots | AGB | Roots | AGB | Roots |                 |          |
| HCB               | 12.170 a | <LOD     | 13.572 a | 2449 b  | 2760 b | 3395 b | <0.001           | <0.001   |
| Keltan            | 198 a   | 206 a     | 206 a     | 179 a   | 50.5 c | 121 b   | <0.001           | 0.11     |
| Methoxychlor      | 71.7 c  | 94.0 c    | 235 b     | 464 a   | 236 b  | 237 b   | <0.001           | <0.01    |
| γ-HCH             | 151 a   | 16.3 c    | 70.0 b    | 19.0 c  | 29.5 bc | 15.5 c  | <0.001           | <0.001   |
| Cr                | 5.04 a  | 3.52 c    | 3.90 bc   | 3.69 bc | 3.83 bc | 4.45 ab | <0.10            | <0.05    |
| Co                | 1.87 ab | 1.60 b    | 2.10 a    | 1.67 b  | 1.56 b  | 1.57 b  | <0.01            | <0.01    |
| Ni                | 8.17 ab | 5.18 c    | 8.65 a    | 6.08 bc | 5.48 c  | 6.81 abc | <0.10            | <0.01    |
| Cu                | 19.1 bcd| 12.7 d    | 30.1 a    | 22.6 b  | 15.6 ed | 19.4 bc | <0.001           | <0.01    |
| Zn                | 56.0 a  | 21.0 d    | 46.6 b    | 29.7 c  | 55.8 a  | 35.4 c  | <0.001           | <0.001   |

Note: OCPs concentrations are presented in µg kg⁻¹; TTEs concentrations are presented in mg kg⁻¹; LOD = 0.1 µg kg⁻¹; SO—soil origin; PP—plant parts.

Aldrin was detected in AGB and roots when the plant was grown in soil K, the most contaminated with OCPs, and in roots only when the plant was grown in the least contaminated soil A; this substance was not detected in the plant’s organs during growth in soil B (i.e., <LOD) (Tables 2 and 3). A bioconcentration effect was observed only during development in soil A (BCF of 3.2), while in soil K, BCFs for AGB and roots were below 0.3 (Figure 1). Thus, aldrin was mainly accumulated in roots and not translocated to the AGB (Figure 2).

Endosulfans were found in the AGB and roots of plants grown in all research soils. The highest concentrations in plants were observed in soil A, followed by soils B and K, with 955, 664, and 520 µg kg⁻¹ average concentrations within the plant, respectively, and all differences were significant (Table 3). Although no differences were observed between concentrations of endosulfans in AGB and roots, BCF values were inversely correlated with concentrations in soils, ranging from 11.5 (soil A) to 0.7 (soil K) (Figure 1). Despite the absence of a statistically significant difference between endosulfan concentrations in soils A and B (Table 2), the corresponding BCF values differed significantly.
Figure 1. Bioconcentration factors for OCPs: (a) BCFs < 20; (b) BCFs > 50. OCPs highlighted in bold indicate a significant difference between BCFs due to soil origin and plant part effects. Different letters on the boxplots within one compound indicate a significant difference at $p < 0.05$.

Endrin aldehyde was accumulated in the plant’s organs for all research soils; the concentration of this substance in the plant tissues was higher for soil A (354 µg kg$^{-1}$) compared to soil B (256 µg kg$^{-1}$) and soil K (179 µg kg$^{-1}$), and all pairs were significantly different. Similar to endosulfans, concentrations of endrin aldehyde in plant tissues were inversely correlated with concentrations in soils: $BCF$ for AGB decreased from 5.4 (soil A) to 0.2 (soil K), and $BCF$ for roots from 6.0 (soil A) to 0.2 (soil K), while OCP concentrations...
in soils were 62.8 (soil A), 130.8 (soil B), and 1088 µg kg$^{-1}$ (soil K) (Figure 1; Table 2). Based on the TLFs (Figure 2), no actual translocation of endrin aldehyde from roots to AGB was observed, so the root system was the main accumulation organ.

HCB was quite strongly taken up by P. tomentosa during cultivation in all three soils studied (the concentration of this substance in plant tissues varied from 2.5 to 13.6 mg kg$^{-1}$ (Table 3)); however, unexpectedly, HCB was not detected in the roots when the plant was grown in soil A (i.e., <LOD) (Table 3). The highest and lowest concentrations of OCP were detected in the AGB and roots of plants grown in soil B, most contaminated by HCB (41.6 µg kg$^{-1}$) (Table 2). BCF values for HCB ranging from 58.9 to 571 (Figure 1) were much higher compared to other OCPs. Uptake of HCB depended on its concentrations in the soils: the average BCFs calculated for the whole plant in soils K and B were equal to 193 and 220, respectively, being not significantly different. In soil K, with the lowest concentration of HCB (14.0 µg kg$^{-1}$), its accumulation in AGB and roots was not significantly different, with BCFs of 197 and 243, respectively. The HCB migration within the plant, represented by TLFs, was directly correlated with concentrations in the soils: the more OCP was present in the soil, the more enhanced migration into the AGB was observed (TLFs increased tremendously) (Figure 2).

Keltan was detected in both the AGB and roots of P. tomentosa during growth in all research soils. When the plant was developed in soils A and B, with relatively low concentrations of keltan (11.9 and 22.1 µg kg$^{-1}$, consequently), the accumulation of OCP in the AGB and roots did not differ significantly between soils or plant organs (Table 3). In contrast, keltan concentrations in AGB (50.5 µg kg$^{-1}$) and roots (121 µg kg$^{-1}$) were significantly lower when plants grew in soil K and differed within plant organs (Table 3). The highest BCF of 17.0 was observed for soil A, followed by 8.7 and 2.6 for soils B and K, respectively (Figure 1). Accordingly, the uptake of keltan reduced with the increasing concentrations in soils. A similar trend was detected for TLF values (Figure 2).

Methoxychlor was accumulated almost equally in AGB and roots when the plants grew in the most (soil K) and least (soil A) contaminated soils (Table 3). In contrast, the concentrations of OCP in AGB (235 µg kg$^{-1}$) and roots (464 µg kg$^{-1}$) differed significantly when the plant was grown in soil B. BCF values decreased from 6.5 to 0.2 for AGB and from 8.5 to 0.2 for roots in soils A and K, respectively, representing the inverse correlation between the uptake of methoxychlor and its concentrations in the soils (Figure 1). The opposite trend was observed for the migration of OCP to AGB, i.e., the TLFs were 0.5, 0.8, and 1.0 for soils B, A, and K, respectively (Figure 2). Thus, the translocation occurred at relatively high concentrations in the soil (1307 µg kg$^{-1}$).

The uptake of γ-HCH as well as endosulfans, endrin aldehyde, and keltan decreased with their increasing content in the soils studied (Tables 1 and 3). For all soils, γ-HCH accumulation was higher in AGB than in roots; however, it was without a significant difference for soil K. BCFs for roots changed in a rather wide range (0.2–1.0), albeit not significantly different (Figure 1). BCFs for AGB decreased from 7.8 (soil A) to 3.5 (soil B) and 0.4 (soil K) in parallel with the increasing OCP concentrations in the soils. Translocation of γ-HCH was observed in all soils (Figure 2), and even at a sufficiently high concentration in the soil, OCP migration to AGB remained reasonable, with a TLF of 1.9, indicating the phytoextraction potential of P. tomentosa concerning this substance.

In addition to OCPs, research soils contained different TTEs; for some elements, the concentration in the soils exceeded the MPC (Table 2). Although As was detected in the three research soils, this element was not detected in the plant tissues (i.e., <LOD). Other TTEs originally presented in the research soils, i.e., Cr, Co, Ni, Cu, Zn, Cd, and Pb, were detected in P. tomentosa tissues (Table 3) at concentrations that varied significantly between soils (at least $p < 0.05$). The concentrations of Cr, Co, Ni, Cu, and Zn differed between roots and AGB as well (Table 3). The uptake of Cr, Cu, Zn, and Pb into the AGB correlated directly with their concentrations in the soils, while the accumulation in the roots showed the opposite trend. The uptake of Ni and Cd to both parts of P. tomentosa correlated inversely with their concentrations in the soils, while Co concentrations correlated directly (Table 3).
Since concentrations of Cu, Zn, and Cd in the research soils exceeded the MPC (Table 2), the potential of *P. tomentosa* to uptake and accumulate these elements was investigated in detail. The accumulation capacity of *P. tomentosa* concerning Cu ranged from 12.7 to 30.1 mg kg\(^{-1}\), depending on the soil. In soils A and B, Cu concentrations in the AGB were 19.1 and 30.1 mg kg\(^{-1}\), respectively, being significantly higher than in the roots (12.7 and 22.6 mg kg\(^{-1}\), respectively). In soil K, the Cu concentration was slightly higher in roots (19.4 mg kg\(^{-1}\)) than in AGB (15.6 mg kg\(^{-1}\)). The highest BCFs for AGB and roots were found in soil B with a medium Cu concentration (4.34 mg kg\(^{-1}\)) (Figure 3). The TLFs correlated directly with the Cu concentrations in the soils (Figure 4).

The accumulation of Zn varied significantly in a range from 21.0 to 56.0 mg kg\(^{-1}\). Zn uptake to AGB was higher than uptake to roots in all research soils. BCFs for both AGB and roots correlated inversely with the element’s concentration in the soils, while the TLFs correlated directly (Figures 3 and 4). However, in soil K, with an average concentration of Zn equal to 12.07 ± 3.26 mg kg\(^{-1}\), element uptake decreased and TLF remained at the same level. Furthermore, reduced uptake was observed in soil A, with high Zn contamination; however, translocation increased to 2.7.

**Figure 3.** Bioconcentration factors for TTEs in plant parts: (a) AGB; (b) roots; (c) TTE concentrations in the research soils. Different letters on the bar and boxplots within one element indicate a significant difference at \(p < 0.05\).
The accumulation of Zn varied significantly in a range from 21.0 to 56.0 mg kg\(^{-1}\). Zn uptake to AGB was higher than uptake to roots in all research soils. BCFs for both AGB and roots correlated inversely with the element’s concentration in the soils, while the TLFs correlated directly (Figures 3 and 4). However, in soil K, with an average concentration of Zn equal to 12.07 ± 3.26 mg kg\(^{-1}\), element uptake decreased and TLF remained at the same level. Furthermore, reduced uptake was observed in soil A, with high Zn contamination; however, translocation increased to 2.7.

Cd concentrations accumulated by \textit{P. tomentosa} ranged from 0.53 to 1.29 mg kg\(^{-1}\), with almost equal distribution between plant parts. At the same time, plants showed an ability to bioconcentrate this element in soil K, with BCFs above 1 for both AGB and roots.

### 4. Discussion

The uptake and translocation of contaminants by \textit{P. tomentosa} are highly variable and determined by their characteristics and the level of soil contamination. When analysing the phytoremediation process, some peculiarities were observed, such as the detection of heptachlor in the plants’ AGB in the absence of its traces in the roots. This observation can be explained by the rapid transformation of heptachlor in the living organisms into the more persistent and hazardous metabolite heptachlor epoxide [61], detected in the tissues of \textit{P. tomentosa} (data not shown; soil B: 95.5 ± 13.5 µg kg\(^{-1}\) and 102 ± 3.3 µg kg\(^{-1}\) in AGB and roots, respectively). The absence of hexabromobenzene translocation to AGB could be due to its sufficiently high hydrophobicity coefficient (log \(K_{ow} = 5.7\)) and low water solubility (0.16 µg L\(^{-1}\)) [62]. The behaviour of plants concerning aldrin was quite different:
in soil A, this substance was detected only in the roots of *P. tomentosa*; in soil B, it was not detected in any of the plant organs, and in soil K, aldrin was detected in both AGB and roots. Such behaviour could be explained by the rapid metabolisation of aldrin to less hydrophobic (log *K*<sub>ow</sub> = 5.4) dieldrin [63]. This assumption was confirmed by the presence of dieldrin in the AGB and roots of *P. tomentosa* when the plant was grown in soil K, despite this substance not being originally determined in the soil (Table 2).

Literature data on the phytoremediation potential of *P. tomentosa* toward OCPs are quite limited, as most researchers have mainly studied spiked soils with an essentially narrow range of OCPs, unlike the aged soils investigated in the current study. Therefore, the comparison of the phytoremediation potential of *P. tomentosa* was made with other phytoremediation agents when at least one of the above criteria matched. In the earlier research, we studied the phytoremediation potential of *M. sinensis* while growing in soil K [52]. The comparison of results presented in [52] and obtained in the current study revealed that *M. sinensis* had a better ability to accumulate aldrin and γ-HCH compared to *P. tomentosa* (Table 3).

Results showed that *M. sinensis* accumulated aldrin only in the roots at a concentration of 308 µg kg<sup>-1</sup>, while *P. tomentosa* accumulated aldrin in the AGB (22.5 µg kg<sup>-1</sup>) and roots (57.5 µg kg<sup>-1</sup>) (Table 3). The accumulation of γ-HCH showed a reversed trend: TLF calculated for *M. sinensis* was 4.5 and thus 2.4 times higher than that of *P. tomentosa* (Figure 2). Rissato et al. [9] observed the uptake capacity of *Ricinus communis* L. in soil spiked with OCPs, most of which were present in the current research soils (A, B, and K). Although the uptake potential of *P. tomentosa* for aldrin, heptachlor, methoxychlor, and γ-HCH was higher than that of *R. communis*, the translocation of aldrin to AGB was lower. Sojinu et al. [64] studied the residues of 25 OCPs in 22 native plants, including energy crops (*Citrullus colocynthis*, *Manihot esculenta*, *Zea mays*, and *Pennisetum purpureum*). The phytoextraction ability of *P. tomentosa* related to aldrin, endosulfans, endrin aldehyde, heptachlor, methoxychlor, and γ-HCH when the plant was grown in soil A was compared with the phytoextraction ability of indigenous plants studied by Sojinu et al. [64]. The comparison showed that OCP concentrations were significantly higher in *P. tomentosa* AGB compared to the concentrations of the same substances in various plants investigated by Sojinu et al. [64].

In contrast to results obtained by Sojinu et al. [64], in the current study, aldrin and heptachlor were not detected in the AGB of *P. tomentosa*. The BCFs for AGB calculated from data presented by Sojinu et al. [64] showed that four energy crops were not able to bioconcentrate aldrin (BCFs ranged from 0.02 to 0.12). *P. tomentosa* showed more substantial phytoextraction potential for endosulfans, endrin aldehyde, methoxychlor, and γ-HCH, with BCF values of 10.8, 5.4, 6.5, and 7.8, respectively (Table 4).

| Pollutant      | Current Data of *P. tomentosa* | Data of Sojinu et al. [64] | BCF       |
|----------------|-------------------------------|----------------------------|-----------|
|                | *C. colocynthis*               | *M. esculenta*             | *Z. mays* | *P. purpureum* |
| Aldrin         | ND                            | 0.05                       | 0.02      | 0.12         |
| Endosulfans    | 10.84                         | 0.38                       | 0.99      | 0.57         | 3.46       |
| Endrin aldehyde| 5.38                          | 0.51                       | ND        | 0.77         | 0.53       |
| Heptachlor     | ND                            | 0.33                       | 4.07      | 0.74         | 19.95      |
| Methoxychlor   | 6.46                          | ND                         | 0.70      | 0.51         | 0.38       |
| γ-HCH          | 7.82                          | 0.59                       | 1.10      | 0.55         | 0.58       |

There are different ways to characterise the ability of plants to uptake elements. Peterson [65] proposed two types of accumulation: the accumulation of an element to concentrations higher than in the growth medium (generally soil) or the possession of more significant quantities of an element than usual for such organisms (needing a reference concentration for the same plant). This concept is mainly valid for TTEs accumulation; however, it can also be extended to the accumulation of organic compounds.
Baker et al. [66] defined hyperaccumulation as when the concentrations of TTEs in the plant tissues surpass 0.01% for Cd; 0.1% for Co, Cu, Ni, and Pb; or 1% in the case of Mn and Zn. In the current research, the hyperaccumulation effect was not reached for any of the elements as their concentrations in plant tissues never exceeded 30 mg kg\(^{-1}\). The ability of \(P.\) tomentosa to bioconcentrate TTEs was variable: no real bioconcentration (i.e., BCFs around 1 or less) was noted for Co and very little for Cd and Pb (BCFs up to 2 only for slightly contaminated soils B and K). Significant bioconcentration was observed for Ni, Cr, and Zn (BCFs up to 4.7, 4.6, and 3.9, respectively), which became essential for Cu (BCFs up to 7.0). BCFs of several elements (Cr, Ni, Cd, and Pb) decreased more or less distinctly with their increasing concentrations in the soil, which seems to reflect a protection mechanism of \(P.\) tomentosa, which would limit the plant’s phytoremediation ability.

Translocation of TTEs from the roots to AGB of \(P.\) tomentosa was generally low (i.e., TLFs between 0.8 and 1.5). The observed high translocation of Zn in soil A could be linked to the higher concentration of the element in soil and its generally higher mobility [67]. This would lead to the preferable accumulation of Zn in plants’ AGB, especially when TTEs essential for plants are abundant in soil [67]. Indeed, Bahri et al. [39] observed the same peculiarity, with the increased translocation of Zn to the AGB of \(P.\) tomentosa grown in soil with higher Zn content.

Summarising the data presented in Figures 1–4 and Table 3 it can be stated that \(P.\) tomentosa showed strong potential to accumulate certain soil-bound OCPs and TTEs, and can be proposed as an eligible species for phytoremediation programs for soils historically contaminated by a mixture of OCPs and TTEs.

5. Conclusions

The results of the current study indicate the successful cultivation of \(P.\) tomentosa during one growing season in soils historically contaminated with twenty OCPs and eight TTEs, collected in the vicinity of obsolete pesticide stockpiles in Talgar district, Almaty region, Kazakhstan. The phytoremediation potential of the plant was investigated in detail for OCPs, whose concentrations in the soils were significantly different (aldrin, endosulfans, endrin aldehyde, HCB, heptachlor, hexabromobenzene, keltan, methoxychlor, and \(\gamma\)-HCH), and for TTEs, whose concentrations were above the MPCs (Cu, Zn, and Cd). It was revealed that the potential of \(P.\) tomentosa to uptake OCPs and TTEs varied greatly depending on the type of contaminant and their concentration in the soils. Along with the ability to bioconcentrate Cr, Ni, and Cu, the phytoremediation potential of \(P.\) tomentosa to accumulate endosulfans, keltan, and methoxychlor provided very encouraging results. Moreover, the phytoextraction effect was found in the case of \(\gamma\)-HCH (TLFs of 1.9–9.9) and HCB (BCFs of 197–571). The TLF values for TTEs ranged from 0.8 to 1.5, indicating the low translocation of elements from the roots of \(P.\) tomentosa to AGB. Nevertheless, the observed high accumulation of Zn in the plant AGB can possibly be linked to the high mobility and concentration of this element in the soils studied.

The results indicate the strong potential of the use of \(P.\) tomentosa in phytoremediation programs applied to soils contaminated with a mixture of organic and inorganic contaminants (OCPs and TTEs) with the simultaneous production of valuable biomass. A more extensive study is necessary to investigate the phytoremediation efficiency of \(P.\) tomentosa during multiyear vegetation at field scale, as the complexity of field conditions may modify the results that we obtained under greenhouse conditions.

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