Heavy Metal Uptake by Herbs. V. Metal Accumulation and Physiological Effects Induced by Thiuram in *Ocimum basilicum* L.

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Received: 31 March 2017 / Accepted: 3 August 2017 © The Author(s) 2017. This article is an open access publication

Abstract Basil (*Ocimum basilicum* L.) is extensively cultivated as either an important spice and food additive or a source of essential oil crucial for the production of natural phenylpropanoids and terpenoids. It is frequently attacked by fungal diseases. The aim of the study was to estimate the impact of thiuram contact time on the uptake of manganese, cobalt, nickel, copper, zinc, cadmium, and lead by *Ocimum basilicum* L. The relevant plant physiological parameters were also investigated. Two farmland soils typical for the Polish rural environment were used. Studies involved soil analyses, bioavailable, and total forms for all investigated metals, chlorophyll content, and gas exchange. Atomic absorption spectrometry was used to determine concentration of all elements. Analysis of variance proved hypothesis that thiuram treatment of basil significantly influences metal transfer from soil and their concentration in roots and aboveground parts. This effect is mostly visible on the 14th day after the fungicide administration. Thiuram modifies mycoflora in the rhizosphere zone and subsequently affects either metal uptake from the soil environment or their further migration within the basil plant. Notable, those changes are more evident for basil planted in mineral soil as compared to organic soil with higher buffering capacity.

Keywords Thiuram · Basil · Heavy metal bioaccumulation and translocation · Fungicide persistence · Farmland soils

1 Introduction

Medical plants are extensively used and consumed all over the world (Basgel and Erdemoglu 2006). *Ocimum basilicum* L. (sweet basil) is an annual, green-leaved herb (*Lamiaceae* family) that has been widely used as a medicinal and seasoning plant for centuries (Stefan et al. 2013; Beatovic et al. 2015). Basil is extensively cultivated as either an important spice and food additive or a source of essential oil crucial for the production of natural phenylpropanoids and terpenoids (Bazaid et al. 2013). It demonstrates wide antibacterial (Koba et al. 2009; Carovic-Stanko et al. 2010; Moghaddam et al. 2011), antifungal (Adigüzel et al. 2005), immunomodulatory (Tsai et al. 2011), antioxidant (Salles Trevisan et al. 2006; Paduraru et al. 2010; Taie et al. 2010), cardiotonic (Muraildharan and Dhananjayan 2004), antihyperglycemic, hypolipidemic (Zeggwagh et al. 2007), and anticonvulsant (Ismail 2006) activities. Notably, composition and activity of these oils depend strongly...
on agricultural practices and environmental conditions (Kandil et al. 2009; Essetlili et al. 2016). Nowadays, basil is widely present in tropical and subtropical regions (Grayer et al. 1996; Javanmardi et al. 2002; Stefan et al. 2013). In Europe, the cultivation of this plants is concentrated mainly in the Mediterranean area (Golcz and Seidler-Łozykowska 2008), but it is also grown in east and central European countries, with Poland being one of the major suppliers (Nurzyńska-Wierdak 2012; Malinowska and Jankowski 2015). Modern cultivation practices reduced impact of pests on crops quite significantly indeed (Bianchi et al. 2006). On the contrary, the plant exposure to fungi is abundant and fungal-induced plagues pose the serious threat to contemporary agriculture. They are difficult to fight and usually lead to substantial harvest losses (Janisiewicz and Korsten 2002; Snelders et al. 2011; Bruni et al. 2016; Sexton and Howlett 2006). Additionally, food plants used for the drug or spice production should be free of pathogens. Basil is frequently attacked by fungal diseases (Zechni D’Aulerio et al. 1995; Hudaib et al. 2001). They strongly influence phytopathological status of particular plant (Hudaib et al. 2002; Bruni et al. 2005) and, in consequence, the quality of essential oil. Therefore, fungal diseases are becoming the major threat to basil farming in Europe and especially in Poland. Plantations can be efficiently protected by the least toxic contact fungicides which should be applied before the harvest, in the time which secures their complete degradation. In a plethora of organic contact fungicides, dithiocarbamates, and thiuram (tetramethylthiuram disulfide) especially play a vital role in a worldwide control of fungal plant diseases and are widely used to protect herbs, arable crops, vegetables, and decorative plants (Kitagawa et al. 2002). Thiuram mode of action is based on inhibition of the pyruvic dehydrogenase system at the fungal cell level. In particular, it hinders uptake of glucose and oxygen and, in consequence, the formation of carbon dioxide by the fungal spores (Dias et al. 2014). It is well documented that thiuram is an effective ligand for diverse coordination species and has the ability to chelate metal ions in soil environment (Victoriano 2000b, 2000a; Filipe et al. 2013; Adamczyk-Szabela 2015). Nevertheless, to the best of our knowledge, thiuram influence on the heavy metal mobility and uptake by plants have not been widely investigated yet (Adamczyk 2006).

The crop productivity and yield as demonstrated by biomass production and plant growth rates are strongly related to the intensity of photosynthesis processes. On the contrary to the numerous experimental data on the relevant physiological and metabolic effects developed in plants cultivated under control of diverse contact fungicides (Garcia et al. 2003; Xiao et al. 2006; Petit et al. 2008), impact of dithiocarbamates on the photosynthesis has not been extensively examined so far (Sau-Man Po and Ho 1997; Dias et al. 2014).

In this paper, the effect of thiuram on physiological changes and uptake of copper, zinc, manganese, cobalt, nickel, cadmium, and lead by basil plants cultivated in the model pot experiments in mineral and organic soils is reported. This work follows our ongoing investigations on the impact of cultivation conditions on the heavy metal uptake by herbs (Adamczyk 2007; Adamczyk and Jankiewicz 2008).

2 Materials and Methods

2.1 Soil Analysis

Two soil types named hereafter A and B were applied. Soil A samples were collected on farmland (uncultivated for at least 2 years before the beginning of the experiment) located away from excessive traffic, according to the procedure as in PN-ISO 10381-4: 2007 at Słupia municipality (51° 51’N, 19° 58’; 20 km from Skierniewice, Łódź province, Poland) in October 2014. Soil B is commercially available universal garden soil enriched with organic matter produced by the Hollas Ltd.

All samples were subsequently dried in a well-ventilated place, sifted through a 2-mm stainless steel sieve, and finally stored in plastic bags. Soil pH was measured by the potentiometric method in 1 mol/dm³ potassium chloride solution (PN-ISO 10390: 1997). The well-established gravimetric method for the determination of soil organic matter by the mass loss at 550 °C was applied (Nelson and Sommers 1996; ASTM 2000; Schumacher 2002).

The bioavailable forms of metals were determined in either 1 or 0.5 mol L⁻¹ solutions of hydrochloric acid extracts for soils A and B, respectively (PN-ISO 11259: 2001). The total metal content was measured in samples mineralized using the Anton Pair Multiwave 3000 closed system instrument. The mixture of concentrated HNO₃ (6 mL) and HCl (2 mL) was applied (0.500 g of soil). Metal concentrations were measured by the FAAS with the GBC Scientific Equipment 932 plus spectrometer. The soil properties are listed in Table 1.
2.2 Preparation of Plant Material

Basil was cultivated under laboratory conditions by the well-established pot method (Adamczyk-Szabela et al. 2015) from April to July. Carefully weighted 500 g samples of soils A and B were placed in 48 plastic pots (24 pots per one soil type) with a diameter of 14 cm and a height to 20 cm. Seeds of Ocimum basilicum L. (P. H. Legutko Company, Poland) were sown in an amount of 0.1 g (approximately 100 seeds) per pot. All pots were kept in a growth chamber at controlled temperatures 23 ± 2 and 16 ± 2 °C for day and night, respectively. The relative humidity was limited to 70–75% while the photosynthetic active radiation (PAR) during the 16-h photoperiod was restricted to 400 μmol m⁻² s⁻¹. All plants were regularly watered by deionized water.

Main cultivation experiments were performed in two arrangements, each was related to one type of soil. A single batch consisted of four series of cultures, each with six pots giving, 24 samples for one arrangement altogether. The fourth series in a batch was cultivated as a reference without addition of a fungicide. Thiuram was applied (6 mg thiuram kg⁻¹ of soil) to plants, 1 month after they had been sown. Herbs were harvested in periods of 14, 28, and 42 days after administration of fungicide. The aboveground parts of plants were cut while the roots were separated from soil by washing and rinsing with distilled water. The entire harvest was oven-dried at 45 °C to a constant weight, homogenized, and grounded.

2.3 Determination of Metals in Basil

The dried roots and aboveground parts of basil plant (0.5 g sample) were subjected to microwave mineralization in concentrated HNO₃ (6 mL) and HCl (1 mL) acid solutions using the Anton Paar Multiwave 3000 closed system instrument. Metal contents were determined by the FAAS using an air/acetylene flame and GAAS with the Scientific Equipment GBC 932 plus and GBC, SensAA spectrometers, respectively. Respective metal nitrates (Me(NO₃)₂, Merck) were used for calibration curve determinations. The reliability of the analytical procedures was checked using the certified reference material INCT-MPH-2, containing a mixture of selected Polish herbs (Dybczyński et al. 2004).

### Table 1 Results of soil A and B analysis

| Analysis          | Results                      |
|-------------------|------------------------------|
|                   | Soil A                       | Soil B                       |
| Soil pH           | 4.7                          | 5.5                          |
| Organic matter (%)| 2.4                          | 61                           |
| Manganese (μg g⁻¹) | Total forms 256 ± 15         | 107 ± 8                      |
|                   | Bioavailable forms 119 ± 7    | 77.8 ± 5.4                   |
| Cobalt (μg g⁻¹)   | Total forms 2.17 ± 0.42       | 4.43 ± 0.44                  |
|                   | Bioavailable forms 0.90 ± 0.14| 2.61 ± 0.52                  |
| Nickel (μg g⁻¹)   | Total forms 8.02 ± 0.64       | 59.5 ± 1.8                   |
|                   | Bioavailable forms 1.13 ± 0.12| 54.8 ± 1.9                   |
| Copper (μg g⁻¹)   | Total forms 4.37 ± 0.62       | 35.9 ± 2.1                   |
|                   | Bioavailable forms 0.93 ± 0.10| 16.5 ± 1.4                   |
| Zinc (μg g⁻¹)     | Total forms 8.27 ± 0.91       | 204 ± 13                     |
|                   | Bioavailable forms 2.12 ± 0.28| 100 ± 7                      |
| Cadmium (μg g⁻¹)  | Total forms 0.26 ± 0.01       | 0.77 ± 0.09                  |
|                   | Bioavailable forms 0.18 ± 0.02| 0.37 ± 0.06                  |
| Lead (μg g⁻¹)     | Total forms 7.38 ± 0.57       | 42.6 ± 3.0                   |
|                   | Bioavailable forms 4.72 ± 0.48| 39.6 ± 1.1                   |

n = 5
p = 0.95
n number of samples, p confidence level
2.4 Basil Plant Growth and Its Physiological Activity

Plant height was measured from the soil surface up to the highest part of the leaf. Index of chlorophyll content was evaluated using Konica Minolta SPAD-502, Japan, methodology in which the chlorophyll concentration is determined by measuring the leaf absorbance in the red and near-infrared regions. Readings were taken around the midrib of each leaf sample. Gas exchange (activity of net photosynthesis, stomatal conductance, intercellular concentration of carbon dioxide, and transpiration) were determined with the gas analyzer apparatus TPS-2 (Portable Photosynthesis System, USA) (Grzesik and Romanowska-Duda 2015; Piotrowski et al. 2016; Kalaji et al. 2012, 2016). All measurements were made in triplicate on separate plants.

2.5 Statistical Analysis

A one-way analysis of variance (ANOVA) as implemented in the Microsoft Excel 2010 was used to test the impact of thiuram contact time on the heavy metal accumulation by basil plant cultivated in soils A and B.

3 Results

Soil analysis as summarized in Table 1 points out that both soils A and B are acidic. The organic matter content indicates mineral or organic character of soils A and B, subsequently (Dobrzański and Zawadzki 1995; Fotyma and Mercik 2003). Manganese, cobalt, nickel, copper, zinc, cadmium, and lead content clearly shows that, according to the generally accepted international standards (Council Directive 86/278/EEC; IUSS Working Group WRB 2006), both soils are not contaminated by these metals.

Metal content in roots and aboveground parts of the basil plant for soils A and B are summarized in Fig. 1; numerical data are shown in Tables S1 and S2. Analysis of the certified reference material is collected in Table S3. Plants cultivated in the untreated reference mineral soil A accumulate metals mostly in the roots. In organic soil B, manganese and cobalt are concentrated in the aboveground parts, while nickel, copper, zinc, and cadmium accumulate in roots. Our preliminary investigations on Melissa officinalis and Valeriana officinalis (Adamczyk 2006, 2007; Adamczyk and Jankiewicz 2008) showed that thiuram affects either metal uptake from the soil environment or their further migration within the plant body and prompted us to examine this effect in herbs in a more detailed fashion.

The influence of thiuram contact time on heavy metal accumulation in the plant body was evaluated by ANOVA at the 0.95 probability level (Fig. 2). Detailed numerical data are given in Supplementary material (Tables S4 and S5). The null hypothesis was, whether thiuram treatment influences the metal transfer from soil and their content in roots and aboveground parts of the plant for a particular period of cultivation after the fungicide administration (14, 28, and 42 days). These calculations clearly showed that thiuram affects metal concentration in investigated plants. Major exceptions involved accumulation of cobalt and cadmium at specific times after the addition of thiuram.

Plant uptake of metal from soil was evaluated by its transfer coefficient (TC). This is defined as ratio of particular element concentration in roots to its content in the soil environment (Chen et al. 2016; Galal and Shehata 2015; Liu et al. 2015). Metal distribution inside the plant body was assessed by translocation factor (TF) which is the ratio of element concentration in aboveground part of the plant to that in roots (Shi and Cai 2009; Testiati et al. 2013; Xiao et al. 2015). TCs and TFs computed for four series of cultures in both soils A and B are presented in Figs. 3 and 4, respectively.

Metal uptake by plants depends on their health status and should not be discussed without connection to the plant growth. The latter can be conveniently evaluated by the standard photosynthesis indicators, i.e., index of chlorophyll content in leaves, the activity of net photosynthesis, stomatal conductance, transpiration rate and intercellular concentration of CO₂ (Fig. 5). In this research, all those parameters clearly showed that basil plants were in reasonable growth conditions. However, they are quite sensitive to the type of soil and thiuram contact time. Decreased plant growth rate was observed in mineral soil A while the opposite situation was in the organic soil B. Generally, alterations in the height of basil plants were quite well reflected by photosynthesis indicators. All those parameters were increased after the thiuram administration, especially in organic soil B. As expected, the only exception was intercellular CO₂ concentration which is decreasing upon the photosynthesis intensification. It means that photosynthesis acceleration is larger than that of stomatal conductance.
Fig. 1 Metal content (μg g⁻¹) in roots and aboveground parts of the basil plant displayed against the thiuram contact time (days). Soils A and B are treated separately.
4 Discussion

It has been documented that fungicide administration influences heavy metal uptake by plants from soil (Adamczyk 2006). However, the impact of thiuram decomposition time has not been widely investigated so far. Thiuram is characterized by limited solubility in water (30 mg L$^{-1}$, 20 °C) and shows a pronounced tendency to adsorb on soil particles; therefore, it is quite safe to groundwater systems. In aquatic conditions at pH = 7, its half-life time is 6 days (Gupta et al. 2012a). Opposite to the model water solutions, in real environment, thiuram degrades more rapidly in acidic soils rich in organic matter. According to Howard (1989), in a humus sandy soil, at pH 3.5, thiuram fully decomposes after 4 to 5 weeks. Rising pH to 7.0 extends that time above 14 weeks (Wauchope et al. 1992; Sharma et al. 2003; Sherif et al. 2011). Thiuram degradation in soil is a complicated
process governed by various factors of which moisture, organic content, and microbial activity are of main concern. The major metabolites in soil are dithiocarbamates, dimethylamine, and carbon disulfide (Gupta et al. 2012b).

Thiuram may affect either metal uptake from the soil environment or their further migration within the basil plant. In particular, fungicide administration decreased the content of Zn and Cd during cultivation in mineral soil A, while in organic soil B, the decline was observed for Cd only (Tables S1 and S2). TCs calculated for plants cultivated in the reference, untreated mineral soil A are in the order \( \text{Mn} > \text{Zn} > \text{Co} > \text{Cu} > \text{Ni} > \text{Pb} > \text{Mn} \). Fungicide treatment alters that order for all metal except Cd (Table 2).

Migration of metals in the plant body may be conveniently examined with the TFs. These factors calculated for basil cultivated in the reference, untreated mineral soil A are in the order \( \text{Mn} > \text{Zn} > \text{Co} > \text{Cu} > \text{Pb} > \text{Ni} > \text{Cd} \). For basal grown in organic soil B (without thiuram), TFs are in the series \( \text{Mn} > \text{Co} > \text{Pb} > \text{Cu} > \text{Ni} > \text{Zn} > \text{Cd} \). The largest TF decreases were detected for zinc, cobalt, and lead in plants cultivated in mineral soil A 14 days after thiuram administration. The respective TF series was \( \text{Mn} > \text{Zn} > \text{Cu} > \text{Ni} > \text{Cd} > \text{Co} > \text{Pb} \) (Fig. 4).

Plants grown in organic soil B showed TF increase for manganese and zinc 14 days after fungicide administration. The longer contact time (i.e., 28 and 42 days) resulted in TF stabilization. After 42 days, TF values computed for majority of metals were quite close to those reported for untreated soil. Opposite situation was observed in mineral soil A where strong TF increase was identified for manganese, zinc, lead, and nickel after 42-day incubation time. Higher impact of thiuram on heavy metal uptake was found in mineral soil A as compared to organic soil B. This may be related to thiuram persistence in complicated soil matrices. According to Gupta et al. (2012a) high concentration of humic acids (as in soil B) increases the rate of thiuram decay and prompts its lower persistence. Additionally, humic acids shows the well-recognized ability to form stable complexes with metals further reducing mobility and rising their retention in organic soils (Pandey et al. 2000). In organic soil B, thiuram alters heavy metal uptake by the basil plant roots in a diverse way with zinc being the most affected species. Its concentration in roots decreases rapidly in 14 days after fungicide administration. This is associated with zinc migration to the fast-growing green parts of the basil plant. The opposite situation is observed in mineral soil A, where thiuram hampers zinc transport to aboveground parts of herbs and stabilizes its accumulation in roots. In this soil, the highest impact of thiuram is visible in manganese uptake, transport, and accumulation. Its concentration increases in either roots or aboveground parts of basil plant. Interaction of fungicide with rhizosphere microflora often leads to substantial local pH modifications (Mukerji et al. 2006) which are more visible in mineral soil A with buffer capacity.

![Table and Figure](https://example.com/table-and-figure.png)
Fig. 3 Transfer coefficients (TC) determined for basil plants cultivated in mineral soil A (a) and organic soil B (b) in the function of time after the fungicide administration. First plot represents untreated control sample.

Fig. 4 Translocation factor (TF) determined for basil plant cultivated in mineral soil A (a) and in organic soil B (b) in the function of time after the fungicide administration. First plot represents untreated control sample.
lower than that of organic soil B. It is well recognized that in either the soil or the plant cell environment, manganese can exist in a number of chemical forms, namely Mn\(^{2+}\) ions and insoluble manganese oxides (Adamczyk-Szabela et al. 2015, Skiba et al. 2017). In acidic conditions, the former are readily available to plants and further prone to migration within the plant body (Adriano 2001; Watmough et al. 2007). Additionally, Mn transfer may also involve superoxide dismutase (SOD) which neutralizes oxygen reactive for thiuram-treated herbs are given in red. The untreated, control samples are in blue. All parameters were determined repeatedly in 14, 28, and 42 days after the fungicide administration.

Fig. 5 Height of the plant, index of chlorophyll content, net photosynthesis (P\(_n\)), stomatal conductance (G\(_s\)), transpiration (E), and intercellular concentrate CO\(_2\) (C\(_i\)) calculated for basil grown in mineral and organic soils A and B, respectively. Data
species (ROS) produced in the plant metabolism. Conserved Mn is an important cofactor which secures the enzyme activity (Whittaker 2010). Metal uptake by roots from soil is strongly dependent on the rhizosphere environment in which bacteria and fungi play the vital role. Additionally, the influence of humic acids cannot be ruled out (Gupta et al. 2016). It is well recognized that mycorrhizal fungi are responsible for nutrients and metal uptake, with zinc and copper being the mostly prone elements (Tinker
and Gilden 1983; Habte 2000). Thiuram reduces fungi populations in soil, but its impact on rhizobacteria is complicated and has not been fully understood yet. Obviously, it deserves more attention in the future.

5 Conclusions

Heavy metals determined in basil originated from soil environment through the root uptake. They are mobilized in rhizosphere by variety of mechanisms involving roots and microorganism exudates. Obviously, thiuram modifies the mycoflora in the rhizosphere zone and subsequently affects either metal uptake from the soil environment or their further migration within the plant. Notable, those changes are more evident for basil planted in mineral soil A as compared to organic soil B with higher buffering capacity. Additionally, migration of metals may be influenced by the formation of sparingly water-soluble metal-fungicide complexes (Beurskens et al. 1971; Zhao et al. 2003). In particular, the latter are presumable responsible for the high manganese uptake and translocation in plants cultivated on mineral soil A. Thiuram impact on metal migration is highly dependent on its persistence and activity in soil. The highest influence was observed 14 days after fungicide administration. Reduction of thiuram activity and, in consequence, the soil and plant recovery is more visible in organic soil B. This effect may influence heavy metal uptake and their further concentration in plant. It should be taken into the consideration when herbal plantations are to be protected with fungicides.

Acknowledgements This work received support from the statutory funds allocated to the Institute of General and Ecological Chemistry by the Polish Ministry of Science and Higher Education.

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