Different strategies of cadmium detoxification in the submerged macrophyte Ceratophyllum demersum L.

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The heavy metal cadmium (Cd) is highly toxic to plants. To understand the mechanisms of tolerance and resistance to Cd, we treated the rootless, submerged macrophyte Ceratophyllum demersum L. with sub-micromolar concentrations of Cd under environmentally relevant conditions. X-ray fluorescence measurements revealed changing distribution patterns of Cd and Zn at non-toxic (0.2 nM, 2 nM), moderately toxic (20 nM) and highly toxic (200 nM) levels of Cd. Increasing Cd concentrations led to enhanced sequestration of Cd into non-photosynthetic tissues like epidermis and vein. At toxic Cd concentrations, Zn was redistributed and mainly found in the vein. Cd treatment induced the synthesis of phytochelatins (PCs) in the plants, with a threshold of induction already at 20 nM Cd for PCs. In comparison, in plants treated with Cu, elevated PC levels were detected only at the highest concentrations (100–200 nM Cu). Our results show that also non-accumulators like C. demersum store toxic metals in tissues where the heavy metal interferes least with metabolic pathways, but remaining toxicity interferes with micronutrient distribution. Furthermore, we found that the induction of phytochelatins is not proportional to metal concentration, but has a distinct threshold, specific for each PC species. Finally we could show that 20 nM Cd, which was previously regarded as non-toxic to most plants, already induces detoxifying mechanisms.

Introduction

Cadmium is an important environmental pollutant and toxic to most organisms. It is highly water soluble, rather immobile in soils but nevertheless can easily accumulate in plants, thus entering the human food chain. Cd concentrations in the environment range from 0.2 to 0.4 nM in unpolluted areas (e.g. Lake Constance) to 5 nM in different slightly contaminated rivers in Germany. Cd release from car tires increased Cd concentrations in contaminated stream in North Central Nigeria, Cd concentrations of 17 nM in New Zealand. And in a heavily contaminated stream in North Central Nigeria, Cd concentrations of 17 nM, 195 nM and 1334 nM were measured. Although there are some indications that Cd may have a positive effect at very low concentrations (e.g. in Zn-limited diatoms), there is overwhelming evidence for Cd-induced toxicity in plants. Due to chemical similarity of Cd with Zn, many toxic effects of Cd are correlated with Zn limitation or replacement, starting with uptake into the plants by transporters with similar affinities for Cd and Zn. However, both synergistic and antagonistic effects of Cd on Zn accumulation have been found in roots and shoots, depending on cultivar and the genotype of wheat and tomato seedlings.

As plants cannot avoid unfavorable conditions like soils with high heavy metal concentrations, they developed several detoxification methods, including immobilization, exclusion, chelation and compartmentalization. One group of detoxifying substances are the enzymatically synthesized phytochelatins (PCs), which after their discovery have been found in higher plants, algae, yeast, some fungi (see review by Clemens, 2006) and also in the nematode C. elegans. Phytochelatins have the general structure (γGlu-Cys)n-Gly (with n = 2 to 11) and are generated by the constitutive enzyme phytochelatin synthase (PCS), which is post-translationally activated only after blockage of thiols by a broad range of heavy metal(loid)s, most efficiently by Cd and As, but also by Ag, Pb, Cu, Hg, Zn, Sn, and Au. Metal–PC complexes are most likely...
transported into and stored in the vacuole where the heavy metal(loid)s cannot interfere with photosynthetic actions.

The storage of heavy metal(loid)s in compartments or tissues where they cannot damage metabolic pathways is a general tolerance mechanism for reducing the amount of heavy metal(loid)s in the cytosol. Furthermore, the few enzymes found in the vacuole\textsuperscript{19} have never been found to be sensitive to heavy metal stress. This protective mechanism is best studied in hyperaccumulator plants. These plants actively accumulate >100 ppm Cd, >1000 ppm Cu or Ni, or >10000 ppm Zn in their above ground tissues.\textsuperscript{20} It should be noted that it was suggested to change the definition of “hyperaccumulator” to >500 ppm Cd in shoots under environmentally relevant conditions with less than 20% growth reduction until the stage of maturity and a bioaccumulation coefficient >5 at a Cd concentration leading to >1000 ppm in shoots.\textsuperscript{21} By tolerating heavy metals, these plants gain advantage over related non-accumulator species as they can colonize soils with elevated heavy metal concentrations. Due to the specialization, many metallophytes become endemic to metalliferous soils.\textsuperscript{22} Compared to equally metal-tolerant non-accumulators, hyperaccumulators have the additional benefit of being protected against herbivores. This was originally shown in 1994,\textsuperscript{23,24} and since then has been shown for many further metals and plant species including the Cd/Zn hyperaccumulator (formally \textit{Thlaspi}) \textit{Noccaea caerulescens}: the Cd-accumulation deterred thrips (\textit{Frankliniella occidentalis}) from feeding on \textit{N. caerulescens} leaves.\textsuperscript{25} It was shown that especially the vacuole of large epidermal cells is used for the storage of the heavy metals.\textsuperscript{26-28} Küpper \textit{et al.} found that in \textit{N. caerulescens} the relative Zn concentration in epidermal cells was positively correlated with cell length in young as well as in mature leaves.\textsuperscript{29} In the related species \textit{N. praecox} collected from a heavy-metal polluted area in Slovenia, Cd was localized in the epidermis, the vascular bundle, but also the mesophyll while Zn was preferentially accumulated in the epidermis cells.\textsuperscript{29} Also in the non-accumulator \textit{Anthyllis vulneraria}, cadmium depositions were found in the central vein, in epidermis cells and in the (epidermal) trichomes.\textsuperscript{30}

There are different techniques to investigate metal accumulation in plants. If one is interested in the cellular or sub-cellular distribution, however, it is important to analyze intact tissues as fragmentation will disrupt the different organelles: metals that were bound to weak ligands in the vacuole may bind stronger ligands present in the cytosol.\textsuperscript{31} For the same reason any fixation techniques besides rapid freezing should be avoided. In this study, we used μ-X-ray fluorescence to determine the localization of Cd, Zn and Cu in frozen hydrated leaves of the rootless, submerged macrophyte \textit{Ceratophyllum demersum}. It is sensitive to heavy metal stress and as it does not possess roots, no root-specific inhibition or detoxification mechanisms are possible.

\section*{Results}

\subsection*{Toxicity determination}

Cadmium treatment led to a reduction in the maximum quantum efficiency of PSII photochemistry, measured as \(F_v/F_m = (F_m - F_0)/F_m\) (Fig. 1). While the values were constant over the treatment duration for the control (0.2 nM) and the low Cd concentration (2 nM), a slight decrease occurred for the plants treated with 20 nM Cd towards the end of the treatment (\(P = 0.051\)). The plants treated with the highest Cd concentration showed a clear reduction of \(F_v/F_m\) from the 2nd week onwards. The Cd-induced inhibition of the photosynthetic apparatus is a great threat to photosynthetic organisms as it was shown to be an important inhibition site.\textsuperscript{9} And although the decrease in \(F_v/F_m\) is often part of a very complicated phenotype, it was established as a stress monitor.\textsuperscript{32}

\subsection*{Phytochelatin determination – method evaluation}

After extraction of phytochelatins with a cooled aqueous solution of 1% formic acid, their high-resolution separation by UHPLC and their detection using ICP-MS and ESI-Q-TOF-MS in parallel was successfully applied within 10 min for PC\textsubscript{2} to PC\textsubscript{6}. The ESI-Q-TOF-MS allows the detection of accurate masses with an m/z error less than 3 ppm. With these accurate masses, proposals for ion formulas were created and used for the identification of reduced and oxidized PCs in the plant extracts. In the present case the parallely occurring ICP-MS detection was applied to quantify the uncomplexed Cd ions in the extracts because the acidic mode of extraction and separation does not allow the analysis of initial Cd–PC-complexes. Despite the dissociation of the Cd–PC-complexes, an excellent chromatographic separation of the free reduced and oxidized PCs can be performed using

\begin{figure}
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Fluorescence quantum yield of PS II measured as \(F_v/F_m = (F_m - F_0)/F_m\) before treatment start and at different times during the treatment duration for 0.2, 2, 20 and 200 nM of Cd. \(F_0 = \) minimal fluorescence quantum yield in dark-adapted tissue; all PSII reaction centers open, \(F_m = \) maximal fluorescence quantum yield in dark adapted tissues; all PSII reaction centers closed. \(F_v = \) variable fluorescence. Values are from 2 experiments and 2 different leaves each. Error bars show error of the mean.}
\end{figure}
A 0.1% formic acid combined with an acetonitrile gradient as an eluent. The dissociation of the complexes is not a drawback for the aim of our work, because grinding tissues anyhow leads to mixing of all soluble constituents of all cells and cellular compartments so that originally weakly bound metals from the vacuole would be mixed with strong ligands, e.g. PCs, in the cytoplasm. Thus immediate binding of previously weakly bound Cd to the PCs would occur once the ground plant material melts. Therefore, we concentrated on identifying and quantifying the different phytochelatin species. Fig. 2 and Table 1 present qualitative values for the identified reduced and oxidized PCs under investigation. In Fig. 2 the ESI-Q-TOF-MS signals (total ion current (TIC) and extracted ion current (EIC)) are presented and used for identification and quantification of PCs in the plant extracts. Additionally to the reduced PCs, also the oxidized PC isomers were considered for quantification. The y-axis scales for each EIC were adjusted for an optimal view of the peaks. In Fig. 2 the shoulder on the PC3 peak could be identified as (PC2)_2. In this case two PC2s are linked together via an inter disulphide bridge. In Fig. 3 the peak at 3.08 min has an accurate mass of m/z 595.1433 that did not correspond to known canonic and iso-PCs. The peak at 5.485 min could be identified as an isomer of oxPC3.

**Phytochelatin induction**

Cadmium treatment induced phytochelatin synthesis in the plants (Fig. 4; P < 0.001, 2-way ANOVA). The induction of phytochelatins was not proportional to metal concentration, but had a distinct threshold, specific for each PC species: while PC2 was the only PC-species detected in both tissue ages of all extracts, PC6 was present mainly in extracts from the higher Cd treatments. In the control treatment (no Cd added; 0.2 nM, background concentration), only trace amounts of PC2 and PC3 were detected. PC3 levels in the young tissue significantly differed only between the lowest (control and 1 nM Cd) and the highest Cd concentrations (P = 0.001). In the old tissue, Cd concentrations from 0.2 nM up to 10 nM led to different PC2 inductions from both 100 nM and 200 nM Cd. Altogether, this means that induction of PC2 by Cd stress was rather weak. PC3, in contrast, had the most prominent induction in response to the metal treatments and yielded the highest amounts. The PC3 content was not different within the group of low Cd (0.2–10 nM) or the group of high Cd (20–200 nM) in young tissue, but between these two groups (P ≤ 0.002), emphasizing the switch-like response of PC3 synthesis. Regarding the old tissue, only the two highest Cd concentrations induced PC3 in amounts different from the other Cd concentrations. PC4 was...
present in most extracts, but not detected in the control plants, in young tissue of the plants treated with 1 nM and 2 nM, and in old tissue of the plants at 0.5 nM Cd. Up to 5 nM Cd for the young and 10 nM for the old tissues, the amounts of PC4 were very low. Moderately elevated (maximum like PC2) levels of PC4 occurred for the young tissue in the three highest Cd treatments, for the old tissue only in the 100 nM treatment. Compared to the shorter PCs, PC5 was detected only in trace amounts, mostly from 10 nM (young tissue) or 20 nM Cd (old tissue) onwards. PC6 was detected only in the plants treated with the highest Cd concentrations (50, 100, 200 nM Cd for PC6-induction in young tissue; 100, 200 nM Cd for PC6-induction in old tissue), with the exception of a trace amount being detected in the old tissue of the plant treated with 1 nM Cd. Generally, the respective amounts of PC6 were much higher in the young than the old tissues ($P < 0.008$); in young tissues PC6 reached double the maximum concentration compared to PC5. Phytochelatins were also extracted from Cu-treated plants, in order to compare the induction threshold because Cu is an essential metal and not a main inducer of PCs (Fig. 4). Here, only PC2 and PC3 were detected in all samples. The respective contents were only significant in the old tissue of plants treated with 200 nM Cu. PC4 was present, only in trace amounts, solely in the young tissue of some plants (0.2, 1, 2, 20, 100, 200 nM Cu) and in the old tissue in plants treated with 50 nM or more Cu. In all copper-treated plants, the amounts of the respective PCs were much lower than in the plants treated with cadmium.

Elemental accumulation and distribution

The higher the applied Cd concentration, the higher was the Cd accumulation in the leaf as revealed by $\mu$-XRF (Fig. 5, left panel). Also when looking at the whole-plant level as analyzed by GF-AAS in acid-digests of tissues, the Cd accumulation within the young and old tissue after 6 weeks of treatment was found to increase with the applied concentration (Table 2).
The treatments with more than 2 nM Cd led to a significant increase ($P \leq 0.013$) of up to more than 2500 ppm. Solely the plants treated with 20 nM Cd showed differences in Cd accumulation depending on the age of the tissue. No Cd was detected by μ-XRF in the leaves of the control treatment. More importantly, Cd distribution changed depending on the Cd treatment concentration and time. With 2 nM of Cd, the distribution was homogeneous over the whole area of the leaf after 3 weeks of treatment, the maximal accumulation being 10.5 μM. After 6 weeks, a stronger accumulation of Cd within the mesophyll and the epidermis was observed and the maximal Cd concentration nearly tripled (28 μM). There was no enhanced accumulation within the vein. A more defined distribution occurred after 3 and 6 weeks of 20 nM Cd treatment and the Cd concentration in the leaves was more than 20-fold increased (640 μM). Cd accumulation was detected mainly in the epidermis and the vein, but also in the mesophyll. After 6 weeks, Cd-accumulation in the epidermis was less pronounced compared to 3 weeks of 20 nM of Cd. An enhanced accumulation was also observed in the thorn (epidermal structure) of the plant treated with 20 nM Cd for 6 weeks. When 200 nM were applied, Cd was mainly observed in the vein. The maximum Cd accumulation in the second replicate was even twice as high as in the shown one.

On the whole-plant level, Cu was in the range of 17–30 ppm and decreased at higher Cd concentrations (Table 2). μ-XRF revealed that Cu was always accumulated solely in the vein, no change in its distribution or concentration was observed in response to Cd toxicity (not shown).

On the whole-plant level, zinc accumulation ranged between 400 and 600 ppm and was only slightly decreased at 20 nM of Cd treatment. However, the decreases were not significant (Table 2). In contrast, the tissue-level distribution pattern of Zn strongly dependent on what Cd concentration was applied (Fig. 5). While in the lower Cd concentrations (0.2 and 2 nM) Zn was mainly found in the epidermis of the leaf, at higher Cd concentrations (20 nM) Zn was more homogeneously distributed. Zn concentration was highest after 3 weeks of treatment with 20 nM Cd (4.5, 4.5 and 3.2 mM). At the highest applied Cd concentration, Zn accumulation was reduced and mainly detected in the vein and the mesophyll, but hardly in the epidermis anymore, which is particularly surprising in view of the almost unchanged Zn concentration at the whole-plant level.

### Discussion

The synthesis of phytochelatins is one major strategy of heavy metal(loid) detoxification. In many previous studies Cd–PC-complexes were isolated from plants, e.g. transgenic and wild type *Arabidopsis thaliana* plants treated with 10 μM of Cd. To break the cell walls and to isolate the substances, plants were lyophilized and ground to a homogeneous mixture. During these processes all cell components get mixed and it cannot be assured that Cd was indeed bound to the PCs before disrupting the cells. Therefore we examined the different PC species rather than Cd–PC-complexes. In the present study we show that not only the concentration of the single PCs but also the occurrence of different PC species (PC$_2$ to PC$_6$) in *Ceratophyllum demersum* is dependent on the applied Cd concentration, and that this occurs at several orders of magnitude lower Cd concentrations than previously known. A previous study on *C. demersum* treated with 5000 and 10 000 nM of Cd for a short period (2 or 4 days) revealed the induction of PC$_2$ and PC$_3$. After 4 days of treatment with 10 000 nM Cd, the amount of PCs decreased while the accumulation of Cd within the tissue continued to increase. The overall content of NP-SHs (non-protein thiols; including PCs) was significantly increased in the plants treated with 5000 nM Cd, and highest in those plants treated for 4 days. In the present study, induction of phytochelatins started already in the low nanomolar range, and the strongest rise in their induction occurred at 20 nM Cd, i.e. 250 times lower than the lowest Cd in the previous study. In the present study, also PC$_4$–PC$_6$ were detected in plants treated with the highest Cd concentration (200 nM), which was still 25 times less than the lowest Cd concentration in the study of Mishra and colleagues. However, they used the Hoagland solution, which contains Fe-EDTA and may have led to chelation of Cd ions. In different macrophytes (*Pistia stratiotes*; water lettuce) treated for 21 days with environmentally relevant concentrations of Cd (10 nM to 1 μM applied Cd, but these authors also used EDTA), enhanced phytochelatin concentrations were detected from 20 nM on, both in leaves and roots. The respective concentration was always higher in the shoots, while GSH was higher in roots. The involvement of PCs in Cd detoxification especially in non-hyperaccumulator plants is well documented and non-controversial. In the present study, however, we could show the dose-dependency for the different PC species under naturally occurring Cd concentrations and

### Table 2

Cd, Zn and Cu accumulation in young and old tissues after 6 weeks of treatment with 0.2, 2, 20 or 200 nM Cd. Values are averages from 2 experiments and technical triplicates within AAS measurement. Error = standard error of the mean. Different letters in the Cd column indicate significant differences ($P < 0.05$) according to the Holm–Sidak method (2-way-ANOVA with comparison of age and applied Cd concentration). Accumulation of Cu and Zn did not differ depending on the Cd treatment ($P > 0.4$).

<table>
<thead>
<tr>
<th>Cd treatment (nM)</th>
<th>Cd [ppm]</th>
<th>Cu [ppm]</th>
<th>Zn [ppm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Young tissue</td>
<td>Old tissue</td>
<td>Young tissue</td>
<td>Old tissue</td>
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<tr>
<td>0.2</td>
<td>9.1 ± 4.3</td>
<td>a</td>
<td>129 ± 6.8</td>
</tr>
<tr>
<td>2</td>
<td>41.3 ± 15.8</td>
<td>b</td>
<td>24.3 ± 8.3</td>
</tr>
<tr>
<td>20</td>
<td>791 ± 17.4</td>
<td>c</td>
<td>453 ± 48.4</td>
</tr>
<tr>
<td>200</td>
<td>2461 ± 702</td>
<td>d</td>
<td>3676 ± 157</td>
</tr>
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</table>

Mishra and colleagues. However, they used the Hoagland treatment and the Cd concentration in the leaves was more than 20-fold increased (640 μM). Cd accumulation was detected mainly in the epidermis and the vein, but also in the mesophyll. After 6 weeks, Cd-accumulation in the epidermis was less pronounced compared to 3 weeks of 20 nM of Cd. An enhanced accumulation was also observed in the thorn (epidermal structure) of the plant treated with 20 nM Cd for 6 weeks. When 200 nM were applied, Cd was mainly observed in the vein. The maximum Cd accumulation in the second replicate was even twice as high as in the shown one.

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culture conditions. This revealed a switch-like induction, with different Cd concentration thresholds for different PC species. Thus an increase in Cd toxicity stress does not just lead to a further overall increase in PC synthesis, but to a change in PC composition. This may suggest that different PCs have different tasks (e.g. homeostasis vs. detoxification) or that plants only invest the synthesis effort into producing long-chain PCs once it becomes critical for survival, while for alleviating moderate toxicity only PCs are formed.

Unlike Cd, Cu is essential for plant growth. It is mainly required in at least six locations in a plant cell including the cytosol, the endoplasmic reticulum (ER), the inner membrane of the mitochondria, the stroma of the chloroplast, the thylakoid lumen and the apoplast. Cu can be found in the active center of various enzymes, e.g. of plastocyanin, a small protein involved in the electron transfer from cytochrome b$_6$f to the PSI reaction centre; the Cu/Zn superoxide-dismutase (Cu/Zn SOD) and also in the cytochrome c oxidase. 10 nM Cu were used for the stock cultures and within the Cd treatment as this had been found to be the optimal concentration for C. demersum (personal communication, G.T. & H.K.). At Cu concentrations over the optimum value, the plants nevertheless cannot detoxify all Cu but need to retain the essential Cu for plastocyanin and other Cu containing enzymes. Metallothioneins (MTs; gene encoded and expressed metal-binding proteins) and PCs as well as high molecular weight ligands seem to be involved in the detoxification process. In this context, the non-induction of PCs by low Cu (Fig. 4) was expected, even though there have been studies indicating an increase in MTs activating the Cu-chelate-reductase in pea plants at Cu deficiency. Apparently, only the highest of the applied Cu concentrations in this study induced enhanced PC production (Fig. 3). Around 400 ppm of Cu was accumulated by the plant at this concentration while the lower concentrations were below 100 ppm which clearly indicates the PC enhancement at the higher concentrations.

The technique of $\mu$-XRF of frozen hydrated samples allows us to determine the distribution of heavy metal(loid)s in plants without chemical fixation and potential alteration and should be favored over techniques that require dehydration, resin embedding or fractionation. In combination with silencing or deleting genes that encode for metal transporters, $\mu$-XRF can also reveal information about metal homeostasis. Studies in which $\mu$-XRF was applied to detect Cd were mainly done with hyperaccumulator plants and the results may be very different in non-hyperaccumulators, especially because Cd will be bound to different ligands. Cd distribution patterns were examined by $\mu$-XRF in two ecotypes of Sedum alfredii, one of which is a Cd-hyperaccumulator and one not. In the hyperaccumulator ecotype (HE), Cd was distributed very homogeneously across the stem cross section within the parenchymal tissue, with the exception of the vascular bundle where Zn was located. In the non-hyperaccumulator ecotype (NHE) Cd and Zn were distributed mainly in the cells surrounding the vascular bundle.

Laser-ablation ICP-MS of whole unprepared leaf samples within the same study revealed an even distribution of Cd over the whole area (except the leaf tips) and the vein, and in the palisade mesophyll and the vascular bundle regarding the leaf cross section. In contrast, Zn accumulation was mainly found in the epidermal layers of the HE, but also in the vascular bundle. In this study, only leaf samples of Ceratophyllum demersum were subjected to $\mu$-XRF analyses. The distribution of Cd and Zn was mostly similar, but changing at higher Cd concentrations: in the plants subjected to control conditions or low non-toxic Cd concentrations, Zn was localized mainly in the epidermis cells (Fig. 5), whereas Cd was either not detected (control treatment), or found mostly all over the leaf (2 nM). With increasing Cd concentrations, a strong and increasing sequestration of Cd and also Zn into the vascular bundle and the thorn was observed. Metal sequestration into the epidermis is a phenomenon well-known for hyperaccumulator plants; it protects the more sensitive mesophyll from metal toxicity (e.g. Zn; Ni). In Arabidopsis halleri, a Cd-hyperaccumulator, Cd and Zn accumulation was found to behave only somewhat similar, as sequestration was stronger in the mesophyll, while in the epidermis it was found specifically within the trichomes, unicellular leaf hairs on the surface of the leaves, which were located around the main vein. A changing distribution over time was observed in A. halleri by Huguet et al.: after 3 weeks of treatment, Cd accumulation occurred in the vein and petiole. After 9 weeks, more Cd was found within the leaves, towards the edges. Furthermore, the distribution of Zn was different from that of Cd at the leaf scale. The non-hyperaccumulator Arabidopsis thaliana also showed enhanced Cd accumulation in the trichomes. The sequestration into the leaf hairs and epidermis seems to be part not only of a detoxification mechanism, but also a defense strategy. Herbivores like snails will indispensably ingest the leaf hairs and other epidermal cells by moving over them and will be deterred from further feeding. A similar situation is also possible for the thorns in C. demersum.

Despite the induction of PCs at high Cd concentrations and their expected storage in vacuoles, highest amounts of Cd were observed in the vein at 200 nM Cd. This may suggest that the applied Cd concentration could not be exported out of the vein into the mesophyll. Already from 20 nM (3 weeks) onward, the distribution of Cd became less differentiated, suggesting limited export due to inhibited transporters. Cd shares chemical properties with Zn and Cd enters plants most likely via Ca channels or Zn transporters. Due to chemical similarity, Cd can replace Zn in the active center of enzymes and inhibit its uptake by competing for transporters or by interfering with the transporter’s gene expression. Cd can inhibit transporters and limit the uptake of other nutrients, and in the same way transporters localized in the vein can be inhibited. This could explain why high Cd had an effect on the distribution of Zn (Fig. 5). The high amount of Cd therefore most likely inhibited Zn distribution at the highest Cd concentration (200 nM, 3 weeks), and after a longer treatment duration with moderately toxic Cd (20 nM, 6 weeks). In both samples, Zn was relocated from the epidermis to the vein (and the thorn). The export of Zn from the vein into the mesophyll will also be inhibited by Cd, when the same transporters are used due to chemical similarity. This Cd-induced inhibition of Zn distribution, which will lead to
Zn starvation of tissues outside the vein and possibly added Zn toxicity inside the vein, could be an important factor in overall Cd toxicity in the plants. Interestingly, this inhibited Zn export from the vein was not linked to inhibition of Zn uptake into the plant as measured at the whole-plant level. There was an increased Zn accumulation in the plant treated with 20 nM for 3 weeks, probably showing a defense mechanism against moderately toxic Cd. An increased uptake of Zn could ensure sufficient Zn within the tissue and has been observed in previous studies. However, Zn content in the young and old tissue of C. demersum did not alter due to Cd treatment (Table 2).

Conclusion

C. demersum can tolerate low or moderately toxic Cd concentrations due to upregulation of phytochelatin synthesis and sequestration into non-photosynthetic tissues. Toxicity of Cd inhibited export of Zn from the vein, and various phytochelatins were switch-like induced at individual Cd threshold concentrations. These aspects of toxicity and detoxification were induced already at very low and environmentally relevant Cd concentrations. The switch-like response of PC synthesis could be used as a toxicity marker in natural habitats.

Experimental

The submerged, rootless macrophyte Ceratophyllum demersum was used for the stress experiments. The strain was obtained from an aquaria shop and continuously cultivated since 2005 in hydroponic solution under 12 h day/12 h night light conditions provided by two Osram FLUORAlight fluorescent and two warm white fluorescent tubes (Osram, München, Germany) and a temperature cycle from 18 °C at 6 a.m., over 20 °C at 9 a.m., to a maximum of 22 °C at 3 p.m., back to over 20 °C at 9 p.m. to 18 °C again at 6 a.m. The nutrient solution (Table 3) was optimized for growth of submerged macrophytes (personal communication from H.K., unpublished) and resembles the situation of typical oligotrophic waters, in particular soft waters. The pH was adjusted to 7.8 with KOH. All experiments were carried out under simulations of natural light and temperature conditions: a 12 h sinusoidal light cycle with maximal irradiances at 750 μmol photons m⁻² s⁻¹ (supplied by Dulux L 55 W/12-954, OSRAM München, Germany) and 12 h night; 19 °C at 6 a.m., 21.5 °C at 9 a.m., 24 °C at 3 p.m., 23 °C at 7 p.m., 19 °C at 6 a.m. For each treatment around 1.5 g (fresh weight) of plants were used. For the two experiments of PC-induction each aquarium contained 2 or 3 plants. The experiments used for metal accumulation and μ-XRF were carried out with 2 or 4 plants. The number of individual plants was consistent for each concentration within the same experiment. Differences occurred due to weight and size of the plants at treatment start. Each aquarium contained 2 L of continuously aerated medium to secure a low biomass to water volume ratio. The nutrient solution was continuously exchanged (flow rate 0.5 L day⁻¹) to ensure that the metal uptake into the plants was limited only by the concentration, but not by the amount of nutrient solution available. After 1 week of acclimation to the high light conditions, cadmium was applied as CdSO₄ and copper as CuSO₄ to the medium. The concentrations were “0” (background Cd/Cu ~ 0.2 nM), 0.5, 1, 2, 5, 10, 20, 50, 100, and 200 nM. Epiphytic algae and cyanobacteria were removed weekly from the macrophytes by gentle brushing and the aquaria cleaned. During the treatment duration, stress symptoms were determined weekly by measuring chlorophyll fluorescence of individual leaves as described previously.⁵⁸,⁵⁹

For phytochelatin analyses, plants were harvested after 2 weeks of treatment, separated into young (4 cm from apex, 2 cm from branch) and old (remaining) parts, the respective weight was determined and the plants frozen in liquid nitrogen until further analyses. Plants from one Cu treatment were harvested after 4 weeks to ensure Cu limitation inside the plants. For μ-XRF analyses, leaves from Cd treated plants after 3 and 6 weeks of treatment were prepared as described below.

Phytochelatin extraction & determination

The harvested plant material was ground in a pre-cooled mortar in liquid nitrogen. The ground material was halved and transferred into two 2 ml cups and covered with 1% formic acid in a concentration of 2 mL g⁻¹ fresh weight. The material was thawed, and phytochelatins were extracted for 1.5 hours at 4 °C. The cups were vortexed three times in between. The material was then centrifuged for 15 min at 2500 × g at 5 °C using a swinging bucket rotor. The purple supernatant was transferred into new cups, rapidly frozen in liquid nitrogen and kept at −80 °C until analysis.

The used synthetic PCs with a purity of >95% were purchased from MoBiTec (Göttingen, Germany). A mixture of PCs with a concentration of 0.05 mM each (PC₂–PC₆) was prepared in 1% formic acid solution as external standard stock solution. Standard mixtures of PC₂ to PC₆ were used as calibrants. On the basis of the peak areas of the “Extracted Ion Chromatogram” (EIC) of the reduced and oxidized forms of the PCs the PC-specific calibrations were performed and the concentrations of the PCs in the extracts calculated. The limit of detection of PC₃ was calculated to be 0.4 μM. The reproducibility (relative standard deviation) differs for the PC under investigations in the following order: PC₂ and PC₄ < 5%, PC₃ < 7.5%, PC₅ < 9% and PC₆ < 15%.

<table>
<thead>
<tr>
<th>Cations</th>
<th>Anions</th>
<th>Concentration in μM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca²⁺</td>
<td>BO₃⁻</td>
<td>80</td>
</tr>
<tr>
<td>Co²⁺</td>
<td>Cl⁻</td>
<td>0.01</td>
</tr>
<tr>
<td>Cu²⁺</td>
<td>CO₂⁻/HCO₃⁻</td>
<td>0.01</td>
</tr>
<tr>
<td>Fe⁺⁺(aq.)</td>
<td>CrO₂⁻</td>
<td>0.2</td>
</tr>
<tr>
<td>K⁺</td>
<td>H₂PO₄⁻/HPO₄⁻</td>
<td>650</td>
</tr>
<tr>
<td>Mg²⁺</td>
<td>I⁻</td>
<td>150</td>
</tr>
<tr>
<td>Mn²⁺</td>
<td>MoO₄⁻</td>
<td>0.4</td>
</tr>
<tr>
<td>Na⁺</td>
<td>NO₃⁻</td>
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</tr>
<tr>
<td>NH₄⁺</td>
<td>SO₄²⁻</td>
<td>10</td>
</tr>
<tr>
<td>Ni²⁺</td>
<td></td>
<td>0.01</td>
</tr>
<tr>
<td>Zn²⁺</td>
<td></td>
<td>0.05</td>
</tr>
<tr>
<td>Fe-EDDHA</td>
<td>HEPES</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Table 3: Concentrations of the ions in the nutrient solution in μM
Element-selective ICP-MS detection at the synchrotron DORIS at the Deutsches Elektronen-Synchrotron (DESY, Hamburg, Germany). X-rays created in a bending magnet were monochromatized using a multilayer monochromator at 30.8 keV with a bandpass of approximately 2.3%. Focusing was achieved using a single-bounce capillary to approximately 10 μm spotsize in both horizontal and vertical dimensions. μ-XRF tomography was done approx. 3 mm above the branching point of the leaf with a step size of 5 μm and a dwell time of 0.8 s per step. Ninety one linescans were measured, with the sample being rotated by 2°, yielding a 180° tomogram. Two fluorescence detectors (Vortex-60 EX/Vortex-90 EX; SII Nano-technology USA Inc., Northridge, California, USA) were used under 90° and 270° with respect to the incident beam to maximize detected fluorescence counts from the sample while minimizing background caused e.g. by elastic scattering due to the polarized nature of the synchrotron radiation. The detected μ-XRF spectra were fitted using PyMca. The fluorescence line areas were then normalized to 100 mA DORIS current, the resulting sonograms were tomographically reconstructed with XRDUA using the maximum likelihood expectation maximization (“MLEM”) algorithm. Absolute concentrations were obtained using the tomographic reconstruction of the multielement-standard measured in the identical geometry.

### Determination of accumulated elements

After 6 weeks of Cd-treatment, plants were harvested, separated into young and old parts and lyophilized for 48 hours. 5–10 mg of the material was put into acid-washed (5% HNO₃) glass tubes and digested in 500 μL (85:15%) nitric-perchloric acid for 30 min at room temperature and then gradually heated up to a maximum of 195 °C until all liquid was vaporized. The remaining ashes were re-dissolved in 0.5 mL 5% HCl, and gradually heated to 80 °C. After cooling, the volume was filled to 1.5 mL with ddH₂O and used for analyzing the components using a graphite furnace atomic absorption spectrometer (GBC 932 AA with GF 3000, Braeside, VIC, Australia). Standard solutions for Cd, Zn and Cu were diluted from AAS Standards (TraceCERT, Sigma-Aldrich, St. Louis, MO, USA). Digested plant samples were appropriately diluted to the optimal detection range with ultrapure 1.66% HCl (ROTIPURAN ultra, Roth, Karlsruhe, Germany).
Acknowledgements

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Notes and references

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