

Food-chain transfer of zinc from contaminated Urtica dioica and Acer pseudoplatanus L. to Microlophium carnosum and Drepanosiphum platanoidis Schrank

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1 Food-chain transfer of zinc to aphids

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- 5 Number of words:
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- 9 Figure 1: Zn concentration in a) the leaf tissue of A. pseudoplatanus (n=25) and above-
- 10 ground tissue of *U. dioica* (n=15) exposed for 98 and 54 days respectively and b) *D.*
- 11 *platinoidis* (n=23) and *M. carnosum* (n=15) exposed for 14 and 28 days respectively
- 12 compared to the Zn concentration in Hoagland's solution and c) is the Zn
- 13 concentration in *D. platinoidis* (n=23) and *M. carnosum* (n=15) compared with the Zn
- 14 concentration in leaf tissue of *A. pseudoplatanus* and above-ground tissue of *U. dioica.*
- 15 (Where [ZnNettle], [ZnLeaf], [ZnAphid] is the concentration of Zn in the tissue of U.
- 16 dioica (mg/kg), the leaf tissue of A.pseudoplatanus (mg/kg), M. carnosum (mg/kg) and
- 17 the watering solution (mg Zn/l) respectively).
- 18 Figure 2: Zn concentration in the phloem tissue of U. dioica (n=15) and A.
- 19 pseudoplatanus (n=15) exposed for 54 and 98 days respectively to Zn in Hoagland's
- 20 solution.

- 21 Food-chain transfer of zinc from contaminated Urtica dioica and Acer
- 22 pseudoplatanus L. to Microlophium carnosum and Drepanosiphum

23 platanoidis Schrank

- 24
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31 Abstract - This study examines the food-chain transfer of Zn from two plant species, Urtica 32 dioica (stinging nettle) and Acer pseudoplatanus (sycamore maple), into their corresponding 33 aphid species, Microlophium carnosum and Drepanosiphum platanoidis. The plants were 34 grown in a hydroponic system using solutions with increasing concentrations of Zn from 0.017 35 to 42 mg Zn/l, although U. dioica only survived in solution containing up to 18 mg Zn/l. 36 Above-ground tissue total and phloem concentrations in U. dioica and M. carnosum 37 concentrations increased with increasing Zn exposure (p<0.001). When U. dioica were 38 exposed to the 18 mg Zn/l solution the corresponding above-ground plant tissue, phloem and 39 *M. carnosum* concentrations were around 2100, 50 and 131 mg/kg respectively. Although Zn 40 concentrations in *M. carnosum* were lower than total plant concentrations bioaccumulation 41 was taking place as concentrations were greater than those in the phloem which represents 42 the Zn reservoir to which the aphids were exposed. Zn concentrations in A. pseudoplatanus 43 also increased with solution concentration from the control to the 9 mg Zn/l solution, after 44 which concentrations remained constant at around 160 mg/kg. Zn concentrations in both D. 45 platanoidis and the phloem tissue of A. pseudoplatanus were not affected by the Zn 46 concentration in the watering solution with concentrations of 6.2 and 375 mg/kg respectively 47 for exposure to solutions of 18 mg Zn/l and above. It appears that A. pseudoplatanus is able 48 to regulate Zn, whereas U. dioica is not resulting in increasing Zn exposure to the aphids on 49 the latter species. Despite this Zn concentrations in *M. carnosum* were around a third of 50 those in *D. platanoidis*, suggesting that the latter species may have naturally elevated Zn 51 concentrations.

52

53 Keywords – stinging nettle, sycamore maple, common nettle aphid, sycamore aphid,

54 contaminated land

55

INTRODUCTION

56 The importance of the impact of contaminated land on terrestrial ecological receptors is 57 increasingly being recognised in the site investigation, risk assessment and remediation 58 process. Many practitioners commonly use an Ecological Risk Assessment (ERA) to 59 determine the potential for harm that a site may pose to ecological receptors and many 60 countries have produced frameworks and guidance for conducting such investigations [1]. 61 The ERA process often makes use of a combination of field and laboratory analysis and 62 models to determine the risk to either ecological function or the food-chain transfer of 63 pollutants. The majority of the ecotoxicological tests used in ERA are based on ecological 64 function and use endpoints such as mortality, reproduction and growth. In order to estimate 65 the risk to higher organisms from a contaminated site it is often necessary to use models to 66 predict the pollutant concentrations through the food-chain and relate these to published 67 toxicological endpoints for the species of interest. There are a variety of models available to 68 estimate the food-chain transfer of pollutants (e.g. [2,3]). However, the models are often not 69 species specific, may have been based on aquatic organisms (for example in the case of 70 flying insects), or may only be applicable to a certain group of contaminants [2,3]. This has 71 serious implications for those using such models to estimate risk from contaminated land to 72 ecological receptors. At best it may result in significant gaps in the range of species for which 73 such a risk assessment can be conducted, at worst it may result in an over or under-74 estimation of the risk leading to either unnecessarily costly remediation or no remediation 75 taking place where it is needed.

Urtica dioica L. is prevalent in almost all urban ecosystems and is an early coloniser of
contaminated land [4,5]. It is extremely important in urban ecosystems as it provides a habitat
for a wide range of invertebrates [5,6]. In addition, it is also relatively simple to cultivate,
widely available and fast growing [6], and as such, may be a useful species for
ecotoxicological testing. *Acer pseudoplatanus* L. is a tree species that has been introduced to
the UK, but is commonly found in urban areas [7]. It is an early coloniser [8] and tolerant of a
wide range of site conditions [9]. *U. dioica* and *A. pseudoplatanus* both have extremely

prevalent species-specific aphids associated with them; *Microlophium carnosum* Buckton and *Drepanosiphum platanoidis* Schrank respectively.

85 The food-chain transfer of metals to a variety of aphids have been assessed in a number of 86 studies (e.g. [10-12]), although these studies have all concentrated on aphids whose hosts 87 are agricultural plant species. Aphids are an important source of food for a large number of 88 other insects, either indirectly for their honeydew (e.g. ants) or directly (e.g. parasitoids and 89 ladybirds) [7,13]. M. carnosum is a large aphid (3.3 to 3.8 mm) commonly found on U. dioica, 90 primarily on the underside of the leaves and the stem [14], during May to October [5,6]. D. 91 platanoidis is abundant on the underside of leaves of A. pseudoplatanus, during April to 92 October, with population peaks in June and October [8]. Used in conjunction with U. dioica 93 and A. pseudoplatanus, M. carnosum and D. platanoidis have the potential to assess the risk 94 of food-chain transfer of metals in urban ecosystems.

This study aims to assess the transfer of Zn to *M. carnosum* and *D. platanoidis* from *U. dioica* and *A. pseudoplatanus* grown under hydroponic conditions in order to determine the potential for Zn transfer to aphid predators in urban ecosystems. The study was originally carried out with Cd in addition to Zn, however the small masses of aphids combined with the smaller concentrations of Cd in their tissue meant that Cd concentrations in aphids were often below detection limits and therefore Cd data are not reported here due to the patchy nature of the dataset.

102

MATERIALS AND METHODS

103 Transfer of Zn into Microlophium carnosum and Drepanosiphum platanoidis

U. dioica cuttings, taken from Alice Holt Forest, Farnham, UK, and *A. pseudoplatanus* (bare
rooted 1+1 stock; Prees Heath Forest Nurseries, Shropshire, UK) were planted individually in
1 litre containers filled with perlite. Perlite was used as it has no inherent sorption capacity
that could influence Zn availability. Additionally, pores between individual perlite beads
ensure an aerobic environment. Pea shingle was placed on the perlite to a depth of 2 cm to
minimise evaporation.

110 A fully replicated randomised block experiment with five replicates was set up in a 111 glasshouse. Plants were grown under 16 h of artificial light and 8 h darkness per day (PAR = 112 0.37 mmol/m/s). The temperature of the glasshouse was regulated to 20 °C (±5 °C). 113 Each container was watered with one of five solutions: control (1/4 strength Hoagland's 114 solution for A. pseudoplatanus and full strength for U. dioica [15]) or one of four Zn treatments 115 in Hoagland's solution. The Hoagland's formulation provided background micronutrient 116 concentrations of 0.02 or 0.08 mg Zn/l and 0.0008 or 0.0032 mg Cu/l for the ¼ strength or full 117 strength solutions respectively. Zn amendments were added as ZnSO₄.7H₂O to provide concentrations of 0.02 (control), 5, 10, 20 and 50 mg Zn/l. The solution in each container was 118 119 replaced by mass when necessary.

120 M. carnosum were added to the U. dioica pots 26 days after planting whilst D. platanoidis 121 were added to the A. pseudoplatanus pots 84 days after planting. Differences in timing were 122 due to the availability of sufficient aphid populations in the field. Leaves with aphids on them 123 were removed from Alice Holt Forest and placed at the base of each plant. Enough leaves 124 were used so that at least 5 aphids were transplanted to each pot. Each pot was then 125 covered individually with a fine mesh net suspended from the ceiling, this was tied securely 126 around the lip at the top of the pot to prevent the aphids from moving to different plants. At 127 each watering, the netting was loosened round the pot and lifted enough to add the 128 appropriate solution and re-secured.

129 U. dioica and A. pseudoplatanus were harvested 28 and 14 days respectively, after the 130 aphids had been added. D. platanoidis populations appeared to be declining on the A. 131 pseudoplatanus so these were harvested earlier than U. dioica in order to ensure that enough 132 aphid mass was available for analysis. Reproduction rates of D. platanoidis vary during the 133 season, being closely linked to the amino-nitrogen content of the leaves and this decline is 134 likely to have been a result of the leaves reaching maturity [8]. The netting was loosened 135 from around the pot and the stem cut, the netting was then closed at the bottom and detached 136 from the ceiling, the netting along with its contents were then placed in the freezer at -20 °C 137 for 2 hours. The plants were then removed from the freezer and the dead aphids collected 138 with a fine brush. A. pseudoplatanus were split into their stem, shoot and leaf components. 139 The above-ground tissue of U. dioica and leaf and shoot tissues of A. pseudoplatanus were

washed in deionised water to remove the honeydew, weighed and dried at 70 °C for 24 hours
and reweighed. The stem tissues of *A. pseudoplatanus* were discarded as the aphids do not
feed on this woody material. The aphids were weighed, dried at 50 °C for 24 hours and
reweighed. The aphid and plant material samples were then milled and analysed to
determine their Zn concentrations (see below).

145 Determination of phloem Zn concentrations

146 In order to understand the different Zn exposures to the aphids a further experiment was set 147 up to determine the phloem Zn concentrations within U. dioica and A. pseudoplatanu. U. 148 dioica cuttings and A. pseudoplatanus were planted individually in 1 litre containers filled with 149 perlite in the same way as for the aphid exposure experiment. A fully replicated randomised block experiment with five replicates for U. dioica and five replicates for A. pseudoplatanus 150 151 was set up in a glasshouse under the same conditions as the aphid exposure experiment. 152 Each container was watered with one of three solutions: control (1/4 strength Hoaglands 153 solution for A. pseudoplatanus and full strength for U. dioica [15]) or one of two Zn treatments 154 in Hoaglands solution. Zn amendments were added as ZnSO₄.7H₂O to provide 155 concentrations of 0.02 (control), 5 and 20 mg Zn/l. The solution in each container was 156 replaced by mass when necessary.

157 U. dioica and A. pseudoplatanus were harvested after the same duration as the aphid 158 experiment in order to ensure that the plants had been exposed to the Zn solutions for the 159 same time. The method used to determine the concentration of Zn in the phloem tissue was 160 based on that of Thornber and Northcote [16] which extracts the water-soluble material within 161 the phloem. The leaf and shoot tissues of A. pseudoplatanus were removed from the stem 162 tissue and discarded. The bark was carefully removed from the stem tissue using a grafting 163 knife and the phloem tissue was then removed, again with a grafting knife. The phloem tissue was weighed and then boiled at 100 °C in 200 ml of deionised water for 3 hours. Following 164 165 boiling, the samples were centrifuged and the solution removed and filtered through a 0.45 166 µm Whatman filter. It was not possible to separate the phloem tissue of U. dioica from the rest of the stem so the entire above-ground biomass was subjected to boiling under the 167 168 assumption that the water-soluble fraction of the plant material will give an indication of the

phloem concentration. The above-ground tissue of *U. dioica* was removed, weighed and
boiled at 100 °C in 300 ml of deionised water for 3 hours. The solutions were then analysed
to determine their Zn concentrations (see below).

172 Determination of Zn concentration

173 The Zn solutions used for watering and the phloem extracts were analysed using a Spectro

174 Flame Inductively Coupled Plasma – Optical Emission Spectrometer (ICP-OES; Spectro

175 Analytical Instruments, West Midlands, UK). The target Zn concentrations in the solutions

used for watering of 0.02, 5 and 20 mg Zn/l were found to be 0.017, 4.71 and 17.97 mg Zn/l

177 respectively.

178 Plant samples were prepared for analysis by dry-ashing at 450 °C for 18 hours and wet

digestion [17]. Wet digestion was achieved by incubating each sample for 1 hour at 60 °C in

180 0.75 ml concentrated HNO₃, followed by a further 14 hour incubation with 2.25 ml

181 concentrated HCI and heating for 2 hours at 110 °C. After cooling, 0.15 ml of 30 % H₂O₂ was

added to each sample followed by heating for 30 minutes at 110 °C. To ensure complete

183 oxidation of all organic matter the H₂O₂ treatment was performed twice. The digested

184 samples were analysed for Zn using the ICP-OES [18].

Aphid samples were digested in 1 ml concentrated HNO₃ at 180 °C for 1 hour, after which 1

186 ml of deionised water was added and the sample further digested at 180 °C to dryness. A

187 further 0.01 ml of concentrated nitric acid was added and the sample digested at 60 °C for 1

hour. The digested samples were analysed for Zn using the ICP-OES [18].

189 The limit of detection was 0.67 µg/kg for Zn. Bush branches and leaves (NCS DC73349,

190 China National Analysis Centre for Iron and Steel), oriental tobacco leaves (CTA-OTL-1,

191 Commission for Trace Analysis of the Committee for Analytical Chemistry of the Polish

192 Academy of Sciences and Institute of Nuclear Chemistry and Technology, Warsaw, Poland),

193 mussel (CE278, European Commission, Geel, Belgium) and bovine liver (1577b, US

194 Department of Commerce, National Institute of Standards and Technology. Gaithersburg, MD

195 20899, USA) tissues were used as Certified Reference Materials (CRM) with batches of plant

196 and aphid samples as appropriate. Mean recovery from oriental tobacco leaves was 104.9

and 98.6 % from the bush branches and leaves and oriental tobacco respectively. Mean

198	recovery from mussel and bovine liver was 92.9 and 93.6 % respectively for the M. carnosum
199	samples and 111.8 and 102.6 % respectively for the <i>D. platanoidis</i> samples.

200 Statistical analysis

201 The plant and aphid Zn uptake data were subjected to general linear regression analysis to

202 assess the significance of changes in plant and aphid concentrations with increasing Zn

203 concentration in hydroponic solutions and plant material respectively, using Genstat version

8.1 [19]. Mean values are reported with \pm standard errors throughout.

205 Linear and exponential models of Zn uptake into each of the plant and aphid tissue types

206 compared to that of the solution concentration and, in the case of aphids, the leaf

207 concentrations were fitted using Genstat version 8.1 [19]. A comparison of the residual sum

208 of squares of alternative models relative to the smallest residual mean square was used to

209 determine the most appropriate model. This comparison used for nested models and is

210 referred to an F-distribution with 1, n degrees of freedom where n is the residual degrees of

211 freedom from the exponential model.

212

RESULTS

213 Zn concentration in solution had a significant affect on the Zn uptake into the above ground 214 tissue ($F_{1,13}$ =533.63; p<0.001) of *U. dioica*; no plants survived in the 42 mg Zn/l solution 215 treatment (Figure 1). Zn concentration in both solution and nettle tissue had a significant 216 affect on the Zn concentration in *M. carnosum* ($F_{1,13}$ =107.95; p<0.001 and $F_{1,13}$ =77.38; 217 p<0.001 respectively; Figure 1). The concentration of Zn in the phloem extracts from *U.* 218 *dioica* increased significantly with increasing Zn concentration in solution ($F_{1,13}$ =138.89; 219 p<0.001; r²=0.908).

220 Zn concentration in solution did not have a significant effect on the Zn uptake into either the 221 leaf or shoot tissue of *A. pseudoplatanus*. This is because the Zn concentrations in the 222 tissues reached a plateau between the 9 and 18 mg Zn/l solutions; the exponential model 223 was, however, significant for both leaf ($F_{2,22}$ =3.57; p=0.046) and shoot ($F_{2,22}$ =5.43; p=0.012) 224 tissue (Figure 1). There was no significant effect of the concentration of Zn in solution or in the leaf or shoot

tissue of *A. pseudoplatanus* on the concentration in *D. platanoidis* using either the linear or

exponential models (Figure 1). Similarly, the concentration of Zn in the phloem extract was

228 not significantly related to the concentration of Zn in solution.

229 The concentration of Zn in the above-ground tissue of U. dioica were approximately 13 times

that in the A. pseudoplatanus as a result of exposure to the 18 mg Zn/l solution;

231 2153±68.7mg/kg compared with 163±20.6 mg/kg. The phloem extract concentrations at this

18 mg Zn/l exposure were 48.2±2.4 mg/kg in *U. dioica* and 6.1±1.2 mg/kg in *A.*

233 pseudoplatanus (Figure 2). Despite this, the Zn concentration in *M. carnosum* was less than

a third of that in *D. platanoidis*; 131.5±11.0 mg/kg compared with 406±21.2 mg/kg. Phloem

235 concentrations of both species were lower than those in above-ground tissue in U. dioica or in

the leaf and shoot tissues in A. pseudoplatanus, this difference increased with increasing Zn

concentrations; from 3 up to 17 times lower and 11 up to 25 times lower in *U. dioica* and *A.*

238 *pseudoplatanus* respectively.

239

DISCUSSION

240 Zn concentrations in the above-ground tissue of *U. dioica* increased with Zn exposure, reaching a mean of approximately 2100 mg/kg for the 18 mg Zn/L solution. In A. 241 242 pseudoplatanus tissue concentration increased up to the 9 mg Zn/l solution and then 243 remained constant at around 160 mg/kg despite the increasing Zn concentration in solution. 244 Zn concentrations in the above-ground tissue of U. dioica have been reported to range 245 between 42 and 52 mg/kg in uncontaminated soils [20]. Leaf concentrations of between 23 246 and 532 (mean 113 mg/kg) have been reported in U. doica growing on dredged sediments 247 with a Zn concentration of between 149 and 1817 (mean 54 mg/kg) [21]. Zn concentrations in 248 U. dioica around the Avonmouth smelter have been found to be as high as 3000 mg/kg, 249 although this is likely to have occurred from atmospheric deposition as well as soil uptake 250 [22]. The substantial quantities of Zn that nettles appear to be capable of accumulating make 251 this species an important pathway for Zn in the food-chain. Mertens et al. [23] found Zn 252 concentrations with a mean of 74 mg/kg in A. pseudoplatanus grown on dredged sediments 253 with a Zn concentration of 359 mg/kg. The normal range of Zn in plant tissue has been

reported to be 27-150 mg/kg with an upper toxic limit of 100-500 mg/kg [24], which suggests that the concentrations reported here for *A. pseudoplatanus* are unlikely to cause a toxic effect.

257 The Zn concentrations in the tissue of U. dioica and A. pseudoplatanus showed large 258 differences; at the lowest Zn solution concentration the tissue concentration of A. 259 pseudoplatanus is greater than that of U. dioica, but at higher concentrations the reverse is 260 true, increasing from a 3 fold to a 13 fold difference at the highest solution concentration. The 261 relationships between solution and tissue concentration between the species were also 262 different; U. dioica having a steep linear relationship whilst for A. pseudoplatanus the 263 relationship was exponential with the Zn tissue concentrations reaching a plateau at around 264 160 mg/kg. This suggests different responses to Zn between the two species. U dioica is 265 unable to regulate Zn and continues to accumulate this metal until a toxic concentration is 266 reached and the plant can no longer survive, in the present experiment this must have 267 occurred to plants grown in the 42 mg Zn / L solution. In contrast A. psuedoplatanus is able to 268 regulate the Zn concentration in its above-ground tissue and therefore survive in media 269 containing higher concentrations of Zn.

270 Previous studies investigating the transfer of metals into aphids have used wheat grown in 271 sewage sludge amended soils. In these studies the Zn concentrations in the plant tissue 272 were substantially lower (<150 mg/kg) [11,12,25-27] than those found in U. dioica in the 273 current study and more comparable to those in A. pseudoplatanus. Despite this, the 274 concentrations of Zn in *M. carnosum* reported in the current study are similar to those found in 275 these previous studies, which used different aphid species [11,12,25-27], whereas the 276 concentrations in D. platanoidis were generally two to three times greater, even at the lowest 277 Zn solution concentration. All of these studies found that Zn was bioaccumulated in the 278 aphids Rhopalosiphum padi and Sitobian avenae feeding on wheat. In our study, from the 279 total plant concentrations it appeared that *M. carnosum* was not accumulating Zn as the 280 U.dioica bulk tissue concentration from the 18 mg Zn/l solution was around 2100 mg/kg and 281 the aphid concentration was 131 mg/kg. However, the analysis of the phloem tissue of the 282 nettle tissue revealed that *M. carnosum* were accumulating Zn as this concentration was

283 around 50 mg/kg. Zn concentrations in D. platanoidis were greater, at around 375 mg/kg, 284 than both the total plant and the phloem concentrations of 160 and 6.2 mg/kg respectively. 285 It has been reported that Zn is concentrated in the stem tissue as well as the roots [28] and is 286 readily transported in the phloem of A. pseudoplatanus [29] and wheat [28,30]. Aphids feed 287 directly on the phloem sap [8] and are therefore exposed to the Zn within this solution. The 288 chemical form that Zn takes within the phloem is not well understood [31], although is likely to 289 be in a soluble form, bound to chelators, amino acids and/or organic acids, it is also unclear 290 whether the Zn is transported apoplastically or symplastically [31]. Studies on barley have 291 shown that, whilst most of the Zn in the roots is soluble, that in the leaves is primarily located 292 in the mesophyll cells and, to a lesser extent the epidermal cells; where is it present primarily 293 in the cytoplasm, followed by the chloroplasts. The Zn in the apoplatsic solution is mainly (97 294 %) bound to cell walls [31]. The ability of A. pseudoplatanus to regulate Zn in its above-295 ground biomass, and because the transfer of Zn to the phloem is regulated by the 296 requirements of the plant, may explain why, in this species, the concentrations in the phloem 297 are similar regardless of the exposure to the plant or plant tissue concentrations. Water-298 soluble concentrations of Zn in U. dioica are much lower than the total plant concentrations, 299 suggesting that the Zn within this species is also bound within the plant tissue and not readily 300 transported in the phloem. However, the water-soluble concentrations increase with 301 increasing Zn concentration in the watering solution and the plant tissue, further suggesting 302 that U. dioica is not able to regulate Zn transport within the plant. This results in increased 303 exposure to aphids as the concentration of Zn in the plant tissue increases. 304 Although it appears that both aphid species bioaccumulated Zn, the concentrations in M. 305 carnosum were smaller than those for D. platanoidis despite an increased level of exposure. 306 This may be because the duration of exposure of D. platanoidis was double that for M. 307 carnosum. Alternatively, M. carnosum may be able to regulate Zn; Crawford et al. [10] found 308 that Aphis fabae on broad beans (Vicia faba) were able to regulate Cu by excretion in 309 honeydew. Unfortunately, it proved impossible to obtain sufficient quantities of honey dew for 310 analysis in this study. The Zn concentrations in *M. carnosum* were comparable with those 311 found in other studies, whereas those in D. platanoidis were elevated. This, coupled with the

312 fact that the concentrations in *D. platanoidis* were elevated even when *A. pseudoplatanus*

was watered with the control solution suggest that this species may simply have naturally
greater Zn concentrations compared with other aphid species regardless of the concentration
within the plant.

316 The greater Zn concentrations in *D. platanoidis* has important implications, both for the

317 estimation of risk to higher organisms and the modelling of food-chain transfer, particularly

318 given that the Zn tissue concentrations in *A. pseudoplatanus* were substantially lower than

- 319 those in *U. dioica*. When the ladybird *Coccinella septempunctata*, lacewing *Chysoperla*
- 320 carnae and carabid bettle Bembidion lampros were fed aphids with Zn concentrations ranging

between 163-249, 104-188 and 60-116 mg/kg respectively their corresponding tissue

322 concentrations were between 184-217, 105-249 and 99-112 respectively [12,26,27]. This

323 suggests that, although only in the lacewing was Zn accumulated, the tissue concentrations of

324 the predators of aphids are likely to reflect the tissue concentrations of their prey. Therefore

325 species feeding on *D. platanoidis* may be exposed to higher concentrations of Zn in their diet

than those feeding on other species of aphid. This demonstrates the importance of species

327 specificity in modelling food-chain transfer in terrestrial ecosystems.

328

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334

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a)

× D. platinoidis — M. carnosum O M. carnosum



C)

Figure 1: Zn concentration in a) the above-ground tissue of *U. dioica* (n=15) and the leaf tissue *A. pseudoplatanus* (n=25) exposed for 54 and 98 days respectively and b) *M. carnosum* (n=15) and *D. platinoidis* (n=23) exposed for 28 and 14 days respectively compared to the Zn concentration in Hoagland's solution in which the *U. dioica* and *A. pseudoplatanus* were grown and c) *M. carnosum* (n=15) and *D. platinoidis* (n=23) Zn concentrations compared with the Zn concentration in the above-ground tissue of *U. dioica* and the leaf tissue of *A. pseudoplatanus*. (Where [ZnNettle], [ZnLeaf], [ZnAphid], [ZnSoln] is the concentration of Zn in the tissue of *U. dioica* (mg/kg dry weight), the leaf tissue of *A.pseudoplatanus* (mg/kg dry weight), *M. carnosum* (mg/kg dry weight) and the watering solution (mg Zn/l) respectively).



Figure 2: Zn concentration in the phloem tissue of *U. dioica* (n=15; mg/kg wet weight) and *A. pseudoplatanus* (n=15; mg/kg wet weight) exposed for 54 and 98 days respectively to Zn in Hoagland's solution.