1	Impacts of the mycotoxin zearalenone on growth and photosynthetic responses in
2	laboratory populations of freshwater macrophytes (Lemna minor) and microalgae
3	(Pseudokirchneriella subcapitata).
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Highlights • Zearalenone (ZON) is a commonly occurring mycotoxin in freshwater samples • ZON aquatic toxicity data are limited hence algae and macrophytes were studied • 72 h algae EC₅₀ = 0.92; NOEC = 0.1; LOEC = 0.23 mg ZON/L, 7 d macrophyte EC₅₀ = 8.8; NOEC = 3.4; LOEC = 11.4 mg ZON/L • PSII efficiency unaffected by ZON in algae and macrophyte exposures

38 Abstract

Mycotoxins are an important class of chemicals of emerging concern, recently detected in 39 aquatic environments, potentially reflecting the influence of fungicide resistance and climatic 40 factors on fungal diseases in agricultural crops. Zearalenone (ZON) is a mycotoxin formed by 41 *Fusarium* spp. and is known for its biological activity in animal tissues; both *in vitro* and *in vivo*. 42 ZON has been reported in US and Polish surface waters at 0.7 - 96 ng/L, with agricultural run-43 44 off and wastewater treatment plants being the likely sources of mycotoxins. As some mycotoxins can induce phytotoxicity, laboratory studies were conducted to evaluate the 45 46 toxicity of ZON (as measured concentrations) to freshwater algae (Pseudokirchneriella subcapitata) and macrophytes (Lemna minor) following OECD test guidelines 201 and 221, 47 respectively. Zinc sulphate was used as a positive control. In the OECD 201 algal static study 48 (72 h at 24 \pm 1°C), exposure to ZON gave average specific growth rate (cell density) EC₅₀ and 49 50 yield (cell density) EC₅₀ values of > 3.1 and 0.92 (0.74 - 1.8) mg/L, respectively. ZON was less toxic in the OECD 221 static study and after 7 d at 24 ± 1°C. L. minor growth was significantly 51 52 reduced based on frond number and frond area at 11.4 mg ZON/L, showing a higher tolerance than reported for other mycotoxins with Lemna spp. Chlorophyll fluorescence parameters 53 were used as biomarkers of impacts on photosystem II efficiency, with no effect seen in algae 54 55 but, with responses being observed in L. minor between 5.2 - 14.4 mg ZON/L. ZON toxicity 56 seen here is not of immediate concern in context with environmental levels, but this study highlights that other freshwater organisms including algae are more sensitive to mycotoxins 57 than *Lemna* sp., the only current source of toxicity data for freshwater plants. 58

60 Graphical abstract



74 **1. Introduction**

The effect of naturally occurring chemicals can sometimes be overshadowed by the growing development of synthetic chemicals, however many naturally produced chemicals have been shown to possess significant endocrine disrupting potential. For example, Cyanobacteria have able to produce retinoid like compounds, potentially causing significant developmental impacts in amphibians and fish (Wu et al., 2012; Smutná et al., 2017). Feminisation of fish due to oestrogenic compounds is widely recognized, with mycoestrogens and phytoestrogens as likely contributors in highly contaminated ecosystems (Jarošová et al., 2015).

Mycotoxins are produced as secondary metabolites by many fungal species. Mycotoxins are 82 commonly associated with cereal crops, but they are also found in other crops such as nuts, 83 84 fruit and coffee. Animals feeding on mycotoxin-contaminated feed have shown toxic effects such as protein synthesis inhibition, immunosuppression and carcinogenicity (Zain, 2011). 85 86 Hence, due to their potential risk to human and animal health, the levels of mycotoxins in foodstuff are regulated by European Union legislation (EU, 2006). Of the mycotoxins produced 87 by Fusarium sp., Zearalenone (ZON) is a known mycoestrogen. Therefore, ZON is associated 88 89 with potential reproductive effects and can cause hypoestrogenism (Cano-Sancho et al., 2012; 90 Rashedi et al., 2012). The metabolites of ZON, α -zearalanol and β -zearalanol, are also oestrogenic; with α -zearalanol licensed as a growth promoter for cattle in some non-EU 91 92 countries (Le Guevel & Pakdel, 2001; Bartelt-Hunt et al., 2012).

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Studies in the US and Poland have found low levels (0.7 - 96 ng/L) of ZON in streams and rivers
with the main sources being agricultural runoff and wastewater treatment plant effluent
(Gromadzka et al., 2009; Kolpin et al., 2014). However, few studies have considered the levels

97 at which mycotoxins can have toxic effects on freshwater species. For ZON toxicity to zebrafish embryos, Bakos et al. (2013) found a 5 d development effect concentration 50 % (EC₅₀) of 50 98 μg/L and lethal concentration 50 % (LC₅₀) of 893 μg/L. Schwartz et al. (2010) reported a 21 d 99 development LOEC and mortality LOEC of > 3.2 μ g/l, 1 μ g/l for vitellogenin production LOEC 100 101 and 0.1 μ g/l for fecundity LOEC. In a longer life cycle (140 d) test with zebrafish a sex ratio 102 LOEC of 0.32 µg/L was seen (Schwartz et al., 2013). In contrast, there is a lack of ZON 103 phytotoxicity data which is needed in order to develop an environmental risk assessment of 104 this widespread mycotoxin. This is important given that other mycotoxins have been shown to cause phytotoxicity in Lemna spp. (eg growth inhibition of 40 % at 3.2 mg nivalenol/L, 56 % 105 at 3.2 mg deoxynivalenol/L and 72 % at 5.6 mg T-2 toxin/L (Abbas et al., 2013)). 106

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Lemna sp. are popular choice in chemical toxicity monitoring for freshwater primary 108 109 producers, due to their small size, rapid growth and ease of culturing. The microalga 110 Pseudokirchneriella subcapitata, previously known as Selanastrium capricornutum and Rhapidocelis subcapitata, similarly is a well-studied organism with its rapid growth rate 111 112 allowing multiple generations to be studied in a brief time frame. Standardised testing guidelines have been developed for both species (OECD, 2006; OECD, 2011), outlining 113 methods which can be used under laboratory conditions to contribute to the hazard 114 115 assessment of chemicals, through analysing the adverse outcome (AO) at the level of the 116 individual and population. To develop knowledge of the specific mode of action (MoA) of a chemical and link this to the AO, these guidelines can be supplemented with physiological and 117 118 biochemical data. Allowing a flow of events from the molecular changes at the target site to

the eventual population effect to be pieced together via a suggested Adverse outcomepathway (AOP).

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122 The aim of this study was to investigate the phytotoxicity of ZON as a chemical of emerging concern, employing the standardised OECD test guidelines for *L. minor* and *P. subcapitata*. 123 This was achieved with a 7 d or 72 h growth inhibition study for each species respectively. 124 125 Following this, physiological measures of photosynthetic performance and biochemical 126 analysis of lipid peroxidation and catalase activity were performed. These were included as a preliminary investigation into MoA measures which can easily be added to the existing 127 guideline framework and analysed for indication of pathways to be pursued to develop AOP's 128 for the test chemical. For quality control purposes, zinc was used as a reference toxicant as 129 per the UK Direct Toxicity Assessment approach (EA, 2007). Zinc is required as a micronutrient 130 131 in plant growth but in excess causes phytotoxicity and reduces growth, previous literature is available for zinc toxicity in both species used in this study (Paixão et al., 2008; Lahive et al., 132 2011). All toxicity data for ZON and Zn reported from the microalgae and macrophyte 133 134 experiments are expressed as measured concentrations unless stated otherwise.

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136 **2. Materials and methods**

137 2.1. Cultures and exposure

A culture of *P. subcapitata* (type strain 278/4) was obtained from the Culture Collection of Algae and Protozoa and maintained in BG11 media made by diluting a sterile stock solution (Sigma Aldrich, Dorset, UK). Prior to experiments, a sub-culture was prepared and held under testing conditions of constant illumination (105-125 μ mol m⁻²s⁻¹) and placed on an orbital shaker set at 120 rpm with temperature in the media maintained at $24 \pm 1^{\circ}$ C. Static exposures were carried out over a 72 h period in accordance with OECD guideline 201 (OECD, 2011). A healthy exponentially growing culture (monitored by increase in cell density) was used to inoculate 25 ml of growth media in polystyrene 50 ml capacity cell culture flasks with filter caps (Greiner, Gloucestershire, UK, C6481) at a density of 5×10^3 cells/ml. Three replicates per test solution were used. Test vessels were placed randomly on an orbital shaker and rearranged daily.

149 A culture of L. minor (UTCC #490) was maintained in Swedish standard (SIS) media, with the pH adjusted to 6.5 \pm 0.2, under a 16:8 h light dark cycle (white fluorescent light) at 24 \pm 1°C. 150 151 Cultures were sterilised prior to testing and visibly free from algae at the beginning of 152 exposures. A static exposure was carried out over a 7 d period in accordance with OECD test 153 guideline 221 (OECD, 2006). Healthy colonies consisting of three fronds were placed into individual wells of six-well clear polystyrene microplates (Thermo Fisher Scientific, 154 Massachusetts, USA, product code 130184) with 8 ml test solution. For each concentration 12 155 replicate wells (i.e. 2 duplicate 6 well microplates for each concentration) were used and held 156 157 under the same conditions as during culturing, position of plates in the incubator was 158 randomised throughout the test.

159

160 **2.2. Growth rate**

Based on pilot studies for microalgae, ZON test solutions of nominal concentrations were zero (< 0.18), 0.032 (< 0.18), 0.1 (< 0.18), 0.32 (0.23), 1.0 (0.83) and 3.2 (3.1) mg/L were tested (mean measured concentrations in brackets with a limit of detection (LOD) of 0.18 mg ZON/L). A zinc positive control of 0.2 mg/L (made with zinc sulphate heptahydrate, CAS number 7446-20-0; Sigma Aldrich batch number: 31665; purity \geq 99.5%) was used. The pH of test solutions

was measured at the beginning and end of the study (pH 6.9 - 7.5) with each replicate meeting
the test criteria for pH (OECD 2011). Growth was measured at 24 h intervals by removing 5 µl
from each test vessel and manually calculating cell density using a Neubauer chamber.
Average specific growth rate (ASGR) and yield, inhibition of ASGR and inhibition of yield were
calculated according to the test guideline.:

171
$$\mu_{i-j} = (\ln(N_j) - \ln(N_i))/t$$

where, μ_{i-j} is the ASGR for the time period (*t*) i to *j*, N_i and N_j is the measurement variable (cell density) at the time *i* and *j* respectively, and *t* is the time period from *i* to *j*. Percentage inhibition of ASGR (% I_r) for each test solution, compared to the dilution water control, was calculated using:

176 %
$$I_r = ((\mu_c - \mu_T)/\mu_c) \times 100$$

177 where, μ_c is the mean ASGR in the dilution water control and μ_T is the mean ASGR in each 178 test solution.

179 Yield was determined, by the change in biomass (cell density) over 7 d in each test replicate.

180 Mean inhibition of yield for each treatment was calculated by:

181 %
$$I_y = ((\underline{b_c} - \underline{b_T})/b_c) \times 100$$

182 where % I_y is percentage reduction in yield, b_c is change in biomass for the dilution water 183 control group and b_T is the change in biomass for the treatment.

For *L. minor* ZON test solutions were prepared for nominal concentrations of zero (< 0.18), 0.1 (< 0.18), 0.32 (0.36), 1.0 (1.1), 3.2 (3.4) and 10.0 (11.4) mg/L (mean measured concentration in brackets with an LOD of 0.18 mg ZON/L), plus a reference chemical measured exposure of 1.4 mg Zn/L positive control. Physio-chemical parameters were measured at the beginning and end of the study (dissolved oxygen 8.1 - 9.9 mg/L; temperature 23.8 - 24.0 °C; and pH ranged between 6.4 - 7.5, within the recommended variation of less than 1.5 units). Growth measurements of frond number and frond area (using WinDias 1.5 software with Hitachi KP-D40 digital camera) were taken at t = 0, 2, 5 and 7 d, average specific growth rate (μ) and yield were calculated as described previously for algae, with frond number and frond area used in place of cell density.

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195 2.3. Chlorophyll fluorescence

196 Chlorophyll fluorescence parameters for *P. subcapitata* were measured using a portable 197 fluorimeter (ToxY-PAM, Hansatech Instruments Ltd., King's Lynn, Norfolk, UK). After the 198 exposure, all replicates were dark adapted for 20 mins at room temperature and 2 ml removed 199 for analysis. To measure F_V / F_M (variable fluorescence / maximum fluorescence) samples were 200 exposed to a saturating light pulse of 2000 µmol of photons m/s over 1 s.

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202 The chlorophyll fluorescence parameters for L. minor were measured using a portable fluorimeter (Pocket PEA, Hansatech Instruments Ltd., King's Lynn, Norfolk, UK) with a light 203 204 pulse of 3000 µmol of photons m/s over 1 s. A single colony was taken from six wells in each 205 treatment and dark adapted in a leaf clip for at least 20 mins at room temperature before being measurements were taken. Measurements were taken at t = 7 d of a second exposure 206 207 with concentrations of measured ZON concentrations of 4.8 (5.2), 8.1 (7.9) and 15.0 (14.4) 208 mg/L (mean measured concentration in brackets) and reference chemical mean measured 209 exposure of 1.8 mg Zn/L. Physio-chemical parameters were measured at the beginning and 210 end of the study (dissolved oxygen 8.1 - 10.0 mg/L; pH 6.4 - 7.1; temperature 23.8 - 24.0 °C). 211

212 The chlorophyll fluorescence parameters are based upon the alterations to shape of the 213 fluorescence rise seen in all photosynthetic materials, which can be separated into a sequence termed the OJIP transient and analysed using the JIP test (Appenroth et al., 2001; Yusuf et al., 214 215 2010) to generate expressions including: (1) measures of efficiency and performance such as 216 F_V / F_M (variable fluorescence / maximum fluorescence) the maximal quantum efficiency of 217 PSII, PI_{ABS} and PI_{Total} (performance indices representing energy conservation for reduction of 218 intersystem electron acceptors and PSI terminal acceptors respectively); (2) parameters 219 calculated based on F_0 (minimal fluorescence) and F_M such as TF_M (time to reach maximum chlorophyll fluorescence (F_M)) and area (proportional to the pool size of the electron acceptors 220 Qa on the reducing side of Photosystem II (the area above fluorescence curve between F₀ and 221 F_{M}); along with Fv/F_0 (quantum yield of the photochemical and non-photochemical 222 223 processes); (3) specific energy fluxes per reaction centre such as ABS/RC (absorption of light energy per reaction centre), DIo/RC (energy dissipation per reaction centre), TRo/RC (the 224 energy trapping rate per reaction centre), ETo/RC (the photosynthetic electron transport rate 225 226 per reaction centre) and REo/RC (reduction of acceptors in PSI per reaction centre); (4) 227 Quantum efficiencies or flux ratios such as $\varphi(Po)$ maximum quantum yield of primary 228 photochemistry, $\Psi(Eo)$ probability of a trapped exciton moving an electron past $Q_{A^{-}}$ to the 229 electron transport chain, $\varphi(Eo)$ quantum yield of electron transport from Q_{A^2} , $\delta(Ro)$ probability 230 an electron from the intersystem reduces PSI terminal electron acceptors and $\varphi(Ro)$ quantum yield of reduction of PSI terminal electron acceptors. (Misra et al., 2001; Strasser et al., 2000; 231 232 Yusuf et al., 2010)

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234 2.4. TBARS assay and catalase enzyme activity

The biomass generated during the 72 h microalgae study was too low to perform biochemical analysis of these measures, with Soto et al. (2011) reporting an extended exposure period of 15 days to generate a sufficient biomass of *P. subcapitata* for analysis of TBARS assay and catalase activity.

239 To measure the catalase activity in the Lemna plant material, three replicates from each treatment, ZON concentrations 4.8 (5.2) , 8.1 (7.9) and 15.0 (14.4) mg/L (mean measured 240 241 concentration in brackets) and the reference chemical mean measured exposure of 1.8 mg Zn/L, were weighed individually and manually crushed with a mortar and pestle (due to the 242 243 low weight of L. minor in 14.4 mg ZON/L and 1.8 mg Zn/L treatments, two wells were combined for each replicate) in 100 mM phosphate buffer (pH 7) at a ratio of 1 mg (wet 244 weight): 19 μ l of buffer. Homogenates were centrifuged (10 000 g for 10 mins) and the 245 246 supernatants collected for the catalase assay (method adapted from Beers & Sizer 1952; Aebi 247 1984). A kinetic method was used, where 200 μ l of 10mM H₂O₂ was added to 50 μ l of 248 supernatant in a microplate and the decrease in absorbance (correlating to a decrease in H_2O_2) 249 read at 3 s intervals for 3 mins at 240 nm. Five replicates were measured per sample.

250 The thiobarbituric acid reactive substances (TBARS) method was used as a general measure of 251 oxidative stress in the tissue (method adapted from Esterbauer & Cheeseman (1990); Marnett 1999). Three replicates from each treatment were weighed and homogenised individually 252 253 (due to the low weight of *L. minor* in 14.4 mg ZON/L and 1.8 mg Zn/L two wells were combined 254 for each replicate) in 100 mM phosphate buffer (pH 7.5) at a ratio of 1 mg: 9 µl. Homogenates were centrifuged (10 000 g for 10 mins). Sixty (60) μ L of the supernatant along with 10 μ l of 255 10 mM butylated hydroxytoluene, 150 μ l of 100 mM phosphate buffer, 50 μ l of 10 % ($^{w}/_{v}$) 256 trichloroacetic acid and 75 μ l of 1.3 % (w/v) thiobarbituric acid were mixed and incubated at 257

258 90°C for 60 mins. The absorbance was measured at 530 nm and calibrated against malondialdehyde standards. Protein content of the homogenates used for catalase and 259 260 TBARS assays was determined with the Peirce BCA Protein Assay Kit (Thermo Fisher Scientific, 261 Massachusetts, USA). Briefly, the working reagent was prepared by mixing bicinchoninic acid 262 (BCA) reagent 1 and 2 in a 50:1 ratio, then 10 µl of homogenate was added to 200 µl working 263 reagent and incubated at 37°C for 30 mins, absorbance was read at 562 nm. Data for catalase 264 and TBARS are expressed as absorbance change min/mg homogenate protein and nmol/mg 265 homogenate protein, respectively.

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Analytical chemistry of ZON and use of zinc positive controls. Nominal exposure 267 2.5. concentrations of ZON (CAS number 17924-92-4; Sigma Aldrich, Dorset, UK, batch number 268 269 043M4106V) in all phytotoxicity experiments were verified by test solution analysis using UV-270 Vis spectrometry (SpectraMax 190 microplate reader, Molecular Devices, USA). The LOD for 271 this method was 0.18 mg ZON/L, hence in experiments where some concentrations were 272 below the limit of detection values for both nominal and measured concentrations are provided. Briefly, samples were taken at the beginning and the end of studies and mean 273 274 concentrations for the exposure period were calculated. Samples from the end of studies were 275 centrifuged at 5000 g for 10 min and the supernatant used to avoid any interference by algal 276 growth. The absorbance of 300 μ l of each sample was measured in a UV-STAR 96 well microplate (Greiner, product code 655801) at 270 nm. 277

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For quality control purposes, zinc sulphate heptahydrate was used as a positive control in the *Lemna* spp. studies. Test solutions were collected at the beginning and end of exposures and mean measured concentrations of zinc were determined using Inductively Coupled Plasma-

Optical Emission Spectrometry (ICP-OES, iCAP, Thermo Scientific,) with a limit of detection of 0.001 mg Zn/L. Due to unforeseen technical problems it was not possible to evaluate the measured concentration for the microalgae study, but the exposure performed in line with expectations from previous zinc range finding studies.

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287 2.6. Statistical Analyses of Algal and Macrophyte Data

Statistical analyses were performed using Minitab (Minitab Ltd., Coventry, UK) and GraphPad Prism (GraphPad Software, Inc, California, USA). Biological effects data (based on measured concentrations of Zn or ZON) were tested for significance (P < 0.05) using one-way analysis of variance with Dunnett's post-test or Kruskal Wallace with Dunn's post-test where appropriate, for normal with homogenous variances and non-normal distributions respectively. EC₂₀ and EC₅₀ values (with 95 % confidence intervals) were determined using non-linear regression and then by fitting sigmoidal curves to the data sets.

295 **3. Results**

296 **3.1. Growth inhibition**

297 The controls of *P. subcapitata* in the control media showed an average overall growth rate of 1.38 (SD 0.01), confirming the healthy status of the organism. Furthermore, all experimental 298 299 treatments continued to increase in cell density until the end of the study (Figure 1.). However 300 during the first 24 h, the cell density of the control and two lowest treatments (0.032 and 0.1) 301 increased more than the higher treatments with both 0.83 and 3.1 mg ZON/L visibly not 302 recovering from this by 72 h. Based on the calculated endpoints at the end of the experiment, 303 there was a significant decrease in growth at 0.23 mg/L for 72 h Yield and at 0.83 mg/L for the 72 h Average Specific Growth Rate (ASGR) (P < 0.05) (Table 1). The EC₅₀ values from this study 304

showed as expected yield to be a more sensitive measure ($EC_{50} = 0.92$) than ASGR ($EC_{50} = >$ 306 3.2) as is the nature of these secondary measures according to the OECD guideline.

307 The growth of *L. minor* was assessed throughout the 7 d study; controls had a doubling time 308 of 2.4 d and no significant variation to exponential growth throughout the test period. Values for 7 d measurements are seen in Table 2, along with the % growth inhibition values, 309 310 recommended to be used in analysis by the testing guidelines. The only concentration to show 311 significant difference in growth in comparison to the control was 11.4 mg ZON/L, with 312 inhibition of 38 % for both ASGR (frond number) and ASGR (frond area) and 60 % Yield (frond 313 number) and 67 % Yield (frond area). Since only the highest exposure in the range finder showed significant inhibition at 7 d, the concentrations for the following photosynthetic and 314 315 biochemical measures were adapted to exceed the growth no effect concentration (NOEC) 316 values (3.9 mg ZON/L for all growth variables).

317

318 **3.2.** Chlorophyll fluorescence

319 During the growth inhibition test measures of Fv/Fm for *P. subcapitata* showed no significant 320 differences (Table 3.), with a control mean of 0.49 and exposure means of 0.48 - 0.50 (SD < 0.012). The value for the control mean is lower than that reported in other studies of greater 321 322 than 0.6 (Choi et al., 2012; Vannini et al., 2011), but consistent with historical control means 323 at this laboratory therefore considered to be due to inter laboratory variation. Measures of 324 chlorophyll fluorescence for the dark adapted *L. minor* were carried out in a second test and are shown in Figure 2. The maximum efficiency calculated by Fv/Fm was not affected by ZON 325 326 exposure but to understand the tolerance of plants it is important to observe other chlorophyll parameters. Tfm, Area, ETo/RC, REo/RC, Ψ (Eo), φ (Eo), δ (Ro), φ (Ro), PI_{ABS} and PI_{Total} decreased 327

significantly in all ZON treatments. While ABS/RC and TRo/RC were significantly increased at the highest treatment of 14.4 mg ZON/L. Fv/F₀ and φ (Po) did not alter significantly in any treatment.

331

332 **3.3.** TBARS assay and catalase enzyme activity

To assess potential oxidative stress as a result of photoinhibition TBARS and catalase activity was monitored at 7 d in *L. minor* (Figure 3.). ZON lowered mean TBARS content, with the decrease (54 %) at 14.4 mg ZON/L being significant (P < 0.05). The 1.8 mg Zn/L reference chemical treatment did not lead to significant changes in TBARS content. Catalase rates showed no significant deviation from the control values for any treatment of ZON or Zn.

338

339 4. Discussion

The main finding of this study was both the algae and the aquatic macrophyte show growth inhibition in the presence of ZON, with the algal species being approximately 10 times more sensitive based the most sensitive EC₅₀ values. There was also evidence of interference with photosynthesis only in *L. minor*, but at high ZON concentrations, although this effect was probably not mediated by overt oxidative stress (no change in catalase and TBARS decreasing slightly).

346

4.1. Acute toxicity

348 The phytotoxicity seen in *P. subcapitata* exhibited a concentration dependant response, with 349 no effect on the two lowest concentrations. Recovery was seen at 0.23 mg ZON/L between the 24 and 72 h observations, with only yield significantly inhibited at 72 h, and significant 350 inhibition in the both of the higher exposures of 0.83 and 3.1 mg ZON/L. Whereas in L. minor 351 352 there was no constant change with concentration but a significant growth response at the 353 highest concentration. The only published data for ZON toxicity to L. minor found was an 354 exposure at a single concentration of 1 mg ZON/L, which showed no effect on growth at this 355 concentration (Vanhoutte et al., 2017). This supports our findings and considering the L. minor growth inhibition values of 38 - 67% seen at 11.4 mg ZON/L in this study, ZON appears to be 356 less toxic to Lemna sp. than mycotoxins tested by Abbas et al. (2002; 2013). Where reported 357 358 growth inhibition due to deoxynivalenol, nivalenol, T-2 toxin and verrucarin A, was 38 - 72% 359 at concentrations in the range of 0.5 - 4.6 mg/L, resembling more the EC₂₀ values generated in this study of 3.0 – 6.5 mg ZON/L. The only mycotoxin reported as less toxic to Lemna sp. 360 than ZON is butanolide with 62 % inhibition at 66.7 mg/L (Vesonder et al., 1992). No previous 361 studies for mycotoxin toxicity to microalgae were found for comparison, but our findings 362 363 demonstrate the value of expanding phytotoxicity data to include algae such as P. subcapitata 364 when considering the potential risk of mycotoxins to freshwater ecosystems.

365

366 4.2. Sub lethal effects

Further to measuring the adverse outcome in terms of growth as a result of ZON exposure, we investigated potential MoA leading to the observed phytotoxicity; measures of chlorophyll fluorescence in a dark-adapted state and biochemical indicators of oxidative stress. Of the photosynthetic parameters measured using chlorophyll fluorescence, Fv/Fm is commonly

371 used as an indication of inhibition of photosynthesis, representing maximum efficiency of Photosystem II via the reduction of Q_A; the electron acceptor in PSII. This was the only measure 372 possible with the instrument used for P. subcapitata. Fv/Fm was unaffected in all P. 373 subcapitata and L. minor ZON exposures and the reference zinc controls. For the additional 374 375 parameters in *L. minor*, all mycotoxin (5.2 - 14.4 mg ZON/L) and zinc (1.8 mg Zn/L) treatments 376 showed a significantly reduced time to reach maximum fluorescence (TFm) and indicated 377 some stress may be occurring due to the inhibition of electron transfer; measured by the area 378 between Fo and Fm. Both values decreased with increasing concentration of ZON or zinc (Figure 2.). The visual health of the fronds was not affected with no signs of chlorosis or 379 bleaching of the leaves, suggesting that the chlorophyll content of the fronds was not 380 appreciably depleted. Overall, these data suggest only modest effects of mycotoxin on 381 382 photosynthetic ability under these experimental conditions (5.2 - 14.4 mg ZON/L) and appear not to explain the key mechanisms of mycotoxin phytotoxicity in Lemna spp. (as yield) with 7d 383 EC₂₀ and EC₅₀ values of 3.0 and 8.8 mg ZON/L, respectively (Table 2). 384

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The specific energy fluxes ABS/RC and TRo/RC significantly increased in the highest ZON 386 387 treatment, this could represent alteration to the composition of light harvesting complexes to absorb and trap higher energies in a shorter time period. Measuring pigment content to 388 389 assess heterogeneity would determine whether this was the cause of the increase (Mirkovic 390 et al., 2017). Efficiency in terms of PI_{ABS} and PI_{Total} significantly decreased suggesting that with the increase in absorbance and trapping there is an imbalance in light absorption and 391 392 utilization of energy as these parameters are associated with the energy flow in the electron transport chain (Farias et al., 2016; Zhang et al., 2016). Combining this with the reduction in 393

ETO/RC and REo/RC, representing the energy flux from Q_A^- into the electron transport chain and reduction of PSI terminal acceptors on the electron acceptor side, this adds to the concept of electron transfer being the possible cause of reduced performance. The reduction in quantum yields and ratios $\Psi(Eo)$, $\varphi(Eo)$, $\delta(Ro)$ and $\varphi(Ro)$ also suggest inhibition of electron movement between Q_A and the acceptor side of PSI.

20N has been seen to act as a uncoupler of oxidative phosphorylation in mitochondria of pea plants (Macri et al., 1996). Uncoupling can also occur in chloroplasts, the oxygen evolving complex (OEC) can be uncoupled and lead to inhibition of the re-oxidation of Q_{A}^{-} (He et al., 2018). This would incur the electron transport inhibition effects seen and the decrease in reduction of PSI electron acceptors. However, if uncoupling of the OEC was occurring the Fv/Fo value is sensitive to this and no significant difference for Fv/Fo was detected in our study.

406 Another possibility for MoA is based upon are similarity of our results to those seen in pea 407 leaves treated with (3-(3',4'-dichlorophenyl)-1,1-dimethylurea (DCMU) (Farias et al., 2016), and reflect their finding of performance indices being a more sensitive than both quantum 408 409 yield of PSII φ (Po) and Fv/Fm which were unaffected. The reduction in movement of 410 electrons into the electron transport chain can cause the over excitation of PSII as seen with photosynthetic herbicides including DCMU (Giardi and Pace, 2005). By binding to Q_B; the 411 412 plastoquinone domain, in the D1 protein of chloroplasts, photosynthesis is inhibited with 413 more energy being absorbed than can be transported into the electron transport chain (Gatidou et al., 2015). 414

A potential issue for plants when too much light energy is being absorbed is oxidative stress.
If ZON was acting upon the Q_B region of the D1 protein in the chloroplasts, this region is

417 involved in controlling the electron transport chain and thus limiting the normal production of singlet oxygen. In the presence of ZON the protein quenching of singlet oxygen would be 418 inhibited and could lead to oxidative stress (Kreiger-Liszkay, 2005). In this study there was no 419 effect on catalase activity and TBARS content decreased in ZON exposures, being significant 420 421 in the highest test concentration. This was probably due to the reduced growth of plant tissue, 422 supporting the conclusion of the absence of overt oxidative stress in Lemna spp. under these 423 experimental conditions. However, excess energy can be transferred to non-photosynthetic 424 pathways as a protective mechanism against reactive oxygen species formation. The DIo/RC flux increased in the highest ZON treatment indicating light energy dissipating in the form of 425 heat. These preliminary data show a strong basis to work from with ZON effect electron 426 427 transport, further measures such as light adapted state chlorophyll fluorescence including 428 non-photochemical quenching (NPQ) are a key area to consider to demonstrate this further 429 and show excess energy is being diverted away from the electron transport chain to prevent oxidative stress during the ZON exposure. Furthermore, additional endpoints should consider 430 the point at which electron transport is inhibited, whether as we have suggested it is around 431 432 or after Q_A or whether something is occurring prior to this in the PSII reaction centre at P680 433 or pheophytin.

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435 5. Conclusions and regulatory context

This laboratory study finds ZON to be less toxic to *Lemna* sp. than other mycotoxins reported in literature. With no previous freshwater mycotoxin studies including algae as a test organism, the higher sensitivity of *P. subcapitata* as compared with macrophytes observed in this study demonstrates the importance of using a multi-species approach in ecotoxicology

and when defining environmental safety levels. Suitable conditions for fungal growth on crops, of increased precipitation, suggest surface waters are a vulnerable ecosystem to mycotoxin contamination via run off from fields. Observed phytotoxicity values for freshwater algae and macrophytes generated here show no immediate risk, with the acute NOEC for microalgae 1000 times higher than the maximum concentration reported to date in environmental samples.

446

447 Regarding extrapolation of mycotoxin aquatic phytotoxicity data to other groups of organisms (eg cyanobacteria or seaweeds), the Adverse Outcome Pathway (AOP) approach is a valuable 448 framework (Ankley et al., 2010; Burden et al., 2015). Currently, AOP information for 449 450 mycotoxin-induced phytotoxicity is lacking, with our results showing some indications of phytotoxicity associated with perturbed chlorophyll fluorescence parameters. Mechanistic 451 452 toxicity data are important in understanding the impacts of mycotoxins on aquatic organisms 453 given their widespread occurrence (Gromadzka et al., 2009; Kolpin et al., 2014). The current preliminary data for macrophytes needs further study to understand the mechanism of ZON 454 455 induced phytotoxicity and cytotoxicity since they were not consistent with regard to the concept of photo oxidative stress being due to ZON-induced electron transport inhibition. 456

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475 **References**

- Abbas, H.K., Johnson, B.B., Shier, W.T., Tak, H., Jarvis, B.B. and Boyette, C.D. 2002.
 Phytotoxicity and mammalian cytotoxicity of macrocyclic trichothecene mycotoxins
 from *Myrothecium verrucaria*. *Phytochemistry* 59, 309-313
- Abbas, H. K., Yoshizawa, T. and Shier, W.T. 2013. Cytotoxicity and phytotoxicity of
 trichothecene mycotoxins produced by *Fusarium* spp. *Toxicon*. 74, 68-75
- 481 Aebi, H., 1984. Catalase in vitro. Methods in Enzymology 105, 121-126
- 482 Appenroth, K.J., Stockel, J., Srivastava, A. and Strasser, R.J. 2001. Multiple effects of chromate
- 483 on the photosynthetic apparatus of *Spirodela polyrhiza* as probed by OJIP chlorophyll a
 484 fluorescence measurements. *Environ. Pollut.* 115, 49-64
- 485 Ankley, G.T., Bennett, R.S., Erickson, R.J., Hoff, D.J., Hornung, M.W., Johnson, R.D., Mount,
- 486 D.R., Nichols, J.W., Russom, C.L., Schmieder, P.K., Serrrano, J.A., Tietge, J.E., and
- 487 Villeneuve, D.L., 2010. Adverse outcome pathways: a conceptual framework to support
- 488 ecotoxicology research and risk assessment. *Environ. Toxicol. Chem.* 29, 730-741
- 489 Bakos, K., Kovacs, R., Staszny, A., Sipos, D. K., Urbanyi, B., Muller, F., Csenki, Z. and Kovacs, B.

2013. Developmental toxicity and estrogenic potency or zearalenone in zebrafish (*Danio rerio*). *Aquat. Toxicol.* 136-137, 13-21

- 492 Bartelt-Hunt, S.L., Snow, D.D., Kranz, W.L., Mader, T.L., Shapiro, C.A., van Donk, S.S., Shelton,
- 493 D.P., Tarkalson, D.D. and Zhang, T.C. 2012. Effect of growth promotants on the
- 494 occurrence of endogenous and synthetic steroid hormones on feedlot soils and in runoff
- 495 from beef cattle feeding operations. *Environ. Sci. Technol.* 46, 1352–1360

- Beers, R.F. and Sizer, I.W., 1952. A spectrophotometric method for measuring the breakdown
 of hydrogen peroxide by catalase. *Journal of Biological Chemistry* 195, 133-140
- 498 Burden, N., Sewell, F., Andersen, M.E., Boobis, A., Chipman, J.K., Cronin, M.T., Hutchinson,
- T.H., Kimber, I., and Whelan, M., 2015. Adverse outcome pathways can drive non-animal
 approaches for safety assessment. *J. Appl. Toxicol.* 35, 971-975
- Canao-Sancho, G., Marin, S. Ramos, A.J. and Sanchis, V. 2012. Occurrence of zearalenone, an
 oestrogenic mycotoxin, in Catalonia (Spain) and exposure assessment. *Food Chem. Toxicol.* 50, 835-839
- Choi, C.J., Berges, J.A., and Young, E.B. 2012. Rapid effects of diverse toxic water pollutants on
 chlorophyll a fluorescence: variable responses among freshwater microalgae. Water
 Res. 46, 2615-2626.
- EC, 2006. Regulation setting maximum levels for certain contaminants in foodstuffs. European
 Commission regulation number 1881/2006 on 19th December 2006
- 509 Environment Agency, 2007. The direct toxicity assessment of aqueous environmental samples
- using the juvenile *Daphnia magna* immobilisation test. *Methods for the Examination of*Waters and Associated Materials.
- 512 Esterbauer, H., and Cheeseman, K.H., 1990. Determination of aldehydic lipid peroxidation
- products: malonaldehyde and 4-hydroxynoneal. *Methods in Enzymology* 186, 407–421
- 514 Farias, M.E., Martinazzo, E.G. and Bacarin, M.A. 2016. Chlorophyll fluorescence in the 515 evaluation of photosynthetic electron transport chain inhibitors in the pea. *Rev. Ciênc.*
- 516 *Agron.* 47, 178-186

- 517 Gatidou, G., Stasinakis, S. and Iatrou, I. 2015. Assessing single and joint toxicity of three 518 phenylurea herbicides using *Lemna minor* and *Vibrio fischeri* bioassays. *Chemosphere* 519 119, 69-74
- Giardi, M.T. & Pace, E. 2005. Photosynthetic proteins for technological applications. *Trends Biotechnol.* 23, 257-263
- Gromadzka, K., Waśkiewicz, A., Goliński, P. and Świetlik, J. 2009. Occurrence of estrogenic
 mycotoxin Zearalenone in aqueous environmental samples with various NOM content.
 Water Res. 43, 1051-1059
- He, L., Yu, L., Li, B., Du, N. and Guo, S. 2018. The effect of exogenous calcium on cucumber
 fruit quality, photosynthesis, chlorophyll fluorescence, and fast chlorophyll fluorescence
 during the fruiting period under hypoxic stress. BMC Plant Biol. 18
- Jarošová, B., Javůrek, J., Adamovský, O. & Hilscherová, K. 2015. Phytoestrogens and mycoestrogens in surface waters - their sources, occurrence and potential contribution
- to estrogenic activity. *Environ. Int.* 81, 26-44
- 531 Kolpin, D.W., Schenzel, J., Meyer, M.T., Phillips, P.J., Hubbard, L.E., Scott, T.M., and Bucheli,
- 532 T.D. 2014. Mycotoxins: Diffuse and point source contributions of natural contaminants 533 of emerging concern to streams. *Sci. Total Environ.* 470 & 471, 669-676
- 534 Krieger-Liszkay, A. 2005. Singlet oxygen production in photosynthesis. J. Exp. Bo. 56, 337-346
- Lahive, E., Halloran, J.O., and Jansen, M.A.K. 2011. Differential sensitivity of four *Lemnaceae*species to zinc sulphate. *Environ. Exper. Bot.* 71, 25-33
- 537 Le Guevel, R. and Pakdel, F. 2001. Assessment of oestrogenic potency of chemicals used as
- growth promoter by *in vitro* methods. *Hum. Reprod.* 16, 1030-1036

539	Macri, F., Vianello, A., Braidot, E., Petrussa, E. and Mokhova, E.N. 1996. Zearalenone – induced
540	uncoupling in plant mitochondria is sensitive to 6-ketocholestanol. IUBMB 39, 1001-
541	1006

- 542 Marnett, L.J., 1999. Lipid peroxidation–DNA damage by malondialdehyde. *Mutation Research*543 424, 83–95
- Mirkovic, T., Ostroumov, E.E., Anna, J.M., Van Grondelle, R., Govindjee, G.D. and Scholes, G.D.
 2017. Light Absorption and Energy Transfer in the Antenna Complexes of Photosynthetic
- 546 Organisms. *Chemical reviews*. 117, 249-293
- 547 Misra, A.N., Srivastava, A. and Strasser, R.J. 2001. Utilization of fast chlorophylla fluorescence
 548 technique in assessing the salt/ion sensitivity of mung bean and Brassica seedlings. *J.*549 *Plant Pysiol.* 158, 1173-1181
- OECD, 2006. Test Guideline 221: Lemna sp. Growth Inhibition Test. Organisation for Economic
 Cooperation and Development Guideline for Testing of Chemicals.
- OECD, 2011. Test Guideline 201: Freshwater Alga and Cyanobacteria, Growth Inhibition Test.
 Organisation for Economic Cooperation and Development Guideline for Testing of
 Chemicals.
- Paixão, S.M., Silva, L. Fernandes, A., O'Rourke, K. Mendonça, E. and Picado, A. 2008.
 Performance of a miniaturized algal bioassay in phytotoxicity screening. *Ecotoxicology*17, 165-171
- Rashedi, M., Sohrabi, H.R., Ashjaazdeh M.A. and Rahimi, E. 2012. Zearalenone contamination
 in barley, corn, silage and wheat bran. *Toxicol. Ind. Health* 28, 779-782

- 560 Schwartz, P., Thorpe, K.L., Bucheli, T.D., Wettstein, F.E. and Burkhardt-Holm, P. 2010. Short-561 term exposure to the environmentally relevant estrogenic mycotoxin zearalenone 562 impairs reproduction in fish. *Sci. Total Environ*. 409, 326–333
- 563 Schwartz, P., Bucheli, T.D., Wettstein, F.E. and Burkhardt-Holm, P. 2013. Life-cycle exposure
- to the estrogenic mycotoxin zearalenone affects zebrafish (*Danio rerio*) development and reproduction. *Environ. Toxicol.* 5, 276-289
- Singh, S., Singh, S., Ramachandran, V. and Eapen, S. 2010. Copper tolerance and response of
 antioxidative enzymes in axenically grown *Brassica juncea* (L.) plants. *Ecotoxicol. Environ. Saf.* 73, 1975-1981
- Smutná, M., Priebojová, J., Večerková, J. & Hilscherová, K. 2017. Retinoid-like compounds
 produced by phytoplankton affect embryonic development of *Xenopus laevis*. *Ecotoxicol. Environ. Saf.* 138, 32-38
- 572 Soto, P., Gaete, H. and Hidalgo, M.E. 2011. Assessment of catalase activity, lipid peroxidation,
- 573 chlorophyll-a, and growth rate in the freshwater green algae *Pseudokirchneriella* 574 *subcapitata* exposed to copper and zinc. *Lat. Am. J. Res.* 39, 280-285
- 575 Strasser RJ, Srivastava A, Tsimilli-Michael M (2000) The fluorescence transient as a tool to
- 576 characterize and screen photosynthetic samples. In: Yunus M, Pathre U, Mohanty P (eds)
- 577 Probing photosynthesis: Mechanism, regulation and adaptation. Taylor and Francis,
 578 New York London, pp 445–483
- Vannini, C., Domingo, G., Marsoni, M., Fumagalli, A., Terzaghi, R., Labra, M., De Mattia, F.,
 Onelli, E. and Bracale, M.. 2011. Physiological and molecular effects associated with
 palladium treatment in *Pseudokirchneriella subcapitata*. Aquat. Toxicol 102, 104-113.

582	Vanhoutte, I., Mets, L.D., Bouve, M.D., Uka, V., Mavungu, J.D.D., Saeger, S.D., Gelder, L.D. and
583	Audenaert, K. 2017. Microbial detoxification of deoxynivalenol (DON), assessed via
584	Lemna minor L. bioassay, through biotransformation to 3-epi-DON and 3-epi-DOM-1.
585	<i>Toxins</i> . 9, 63

- Vesonder, R.F., Labeda, D. and Peterson, R.E. 1992. Phytotoxic activity of selected water soluble metabolites of *Fusarium* spp. against *Lemna minor* L. (Duckweed).
 Mycopathologia 118, 185-189
- 589 Wu, X., Jiang, J. & Hu, J. 2012. Determination and occurrence of retinoids in a eutrophic lake
- 590 (Taihu Lake, China): cyanobacteria blooms produce teratogenic retinal. *Environ. Sci.*591 *Technol.* 47, 807-814
- 592 Yusuf, M.A., Kumar, D., Rajwanshi, R., Strasser, R.J., Tsimilli-Michael, M., Govindjee, Sarin, N.B.

593 2010. Overexpression of γ-tocopherol methyl transferase gene in transgenic *Brassica*

juncea plants alleviates abiotic stress: Physiological and chlorophyll a fluorescence measurements. *Biochim Biophys Acta Bioenerg*. 1797, 1428-1438

- Zain, M.E., 2011. Impact of mycotoxins on humans and animals. *Journal of Saudi Chemical Society* 15, 129–144
- Zhang, L., Su, F., Zhang, C., Gong, F. and Liu, J. 2016. Changes of photosynthetic behaviours
 and photoprotection during cell transformation and astaxanthin accumulation in *Haematococcus pluvialis* grown outdoors in tubular photobioreactors. *Int. J. Mol. Sci.* 18,
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- Figure 1. Growth curves of *P. subcapitata* exposed to zearalenone (CAS number 17924-92-4)
- in a 72 h static study at $23.8 \pm 1^{\circ}$ C with 0.2 mg Zinc/L as a positive control.
- Figure 2. Chlorophyll fluorescence parameters measured in *L. minor* after 7 d exposure to
- zearalenone (CAS number 17924-92-4) in a static study at 24 ± 1°C (concentration measured
- using UV-Vis spectrometry with an LOD of 0.18 mg ZON/L). Values are normalised to the
- 611 control group 0 (< 0.18 mg ZON/L).
- Figure 3. Catalase and TBARS content (± SD) measured in *L. minor* after 7 d exposure to
- 513 zearalenone (CAS number 17924-92-4) in a static study at $24 \pm 1^{\circ}$ C.
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- 617 LIST OF TABLES
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- 4) in a 72 h static study at $23.8 \pm 1^{\circ}$ C with 0.2 mg Zinc/L as a positive control.
- Table 2. Growth responses of *L. minor* exposed to zearalenone (CAS number 17924-92-4) in a
- 622 7 d static study at $24 \pm 1^{\circ}$ C.

Table 1. Growth responses of *P. subcapitata* exposed to zearalenone (CAS number 17924-92-

623	Table 3. Maximal quantum efficiency of Photosystem (Fv/Fm) measured in <i>P. subcapitata</i>
624	(mean ± SD) after 72 h exposure to zearalenone (CAS number 17924-92-4) in a static study at
625	24 \pm 1°C (concentration measured using UV-Vis spectrometry with an LOD of 0.18 mg ZON/L).
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Figure 1. Growth curves of *P. subcapitata* exposed to zearalenone (CAS number 17924-92-4)

in a 72 h static study at $23.8 \pm 1^{\circ}$ C with 0.2 mg Zinc/L as a positive control.



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Figure 2. Chlorophyll fluorescence parameters measured in *L. minor* after 7 d exposure to zearalenone (CAS number 17924-92-4) in a static study at $24 \pm 1^{\circ}$ C (concentration measured using UV-Vis spectrometry with an LOD of 0.18 mg ZON/L). Values are normalised to the control group 0 (< 0.18 mg ZON/L).

Footnote - Fv/Fm = maximal quantum efficiency of Photosystem II; TFm = time to reach 639 maximum chlorophyll fluorescence; Area = proportional to the pool size of the electron 640 acceptors Qa on the reducing side of Photosystem II; Fv/F₀ = quantum yield of the 641 642 photochemical and non-photochemical processes; ABS/RC = absorption of light energy per 643 reaction centre; DIo/RC = energy dissipation per reaction centre), TRo/RC (the energy trapping rate per reaction centre; TRo/RC = energy trapping rate per reaction centre; ETo/RC = 644 photosynthetic electron transport rate per reaction centre; REo/RC = reduction of acceptors 645 in PSI per reaction centre; $\varphi(Po)$ = maximum quantum yield of primary photochemistry; $\Psi(Eo)$ 646 = probability of a trapped exciton moving an electron past $Q_{A^{-}}$ to the electron transport chain; 647

- 648 $\varphi(Eo) = quantum yield of electron transport from Q_A⁻; <math>\delta(Ro) = probability an electron from the$
- 649 intersystem reduces PSI terminal electron acceptors and $\varphi(Ro)$ = quantum yield of reduction
- of PSI terminal electron acceptors; PI_{ABS} = performance index of photosynthetic efficiency and
- 651 PI_{Total} = energy conservation for reduction of PSI terminal acceptors respectively.



Figure 3. Catalase and TBARS content (\pm SD) measured in *L. minor* after 7 d exposure to zearalenone (CAS number 17924-92-4) in a static study at 24 \pm 1°C.

Footnote – significant difference only seen between the TBARS content in the dilution water
control and 14.4 mg ZON/L.

- 660 **Table 1.** Growth responses of *P. subcapitata* exposed to zearalenone (CAS number 17924-
- 661 92-4) in a 72 h static study at 23.8 ± 1°C (concentration measured using UV-Vis spectrometry
- with an LOD of 0.18 mg ZON/L) with 0.2 mg Zinc/L as a positive control.
- 663

Nominal	Mean endpoint	Mean inhibition of	algal growth (%)
concentration,	at 72h		664
measured in brackets	Cell density	Average Specific	Yield 665
(mg ZON/L)	(cells/ml x 10^5)	Growth Rate (ASGR)	005
0 (< 0.18)	3.1 ± 0.14	-	_ 666
0.032 (< 0.18)	2.7 ± 0.12	4	¹⁵ 667
0.1 (< 0.18)	2.8 ± 0.14	3	11
0.32 (0.23)	2.6 ± 0.45	5	17ª ⁶⁶⁸
1.0 (0.83)	1.6 ± 0.14	16ª	48ª 669
3.2 (3.12)	1.1 ± 0.08	24ª	64ª
0.2 (Zn)	2.3 ± 0.24	7ª	26°
EC ₂₀ (± 95 % CI)	-	1.72 (1.25 - 2.4)	0.19 (0.08 - 0.363)1
EC ₅₀ (± 95 % CI)	-	> 3.2	0.92 (0.74 - 1. <mark>8</mark>)/2
NOEC	-	0.23	0.1
LOEC	-	0.83	0.23

674 Footnote - ^a Significantly different (P < 0.05) from control treatment

675 Summary effect values calculated with measured values where possible.

677 **Table 2.** Growth responses (± SD) of *L. minor* exposed to zearalenone (CAS number 17924-92-

4) in a 7 d static study at $24 \pm 1^{\circ}$ C (concentration measured using UV-Vis spectrometry with

Nominal	Mean measu	red endpoints	Calculated inhibition of growth (%)			
concentration, measured in brackets			Average Specific Growth Rate		Yield	
(mg ZON/L)	Frond number	Frond area	Frond number	Frond area	Frond number	Frond area
		(mm ²)		(mm ²)		(mm ²)
0 (< 0.18)	23 (± 2.6)	115.3 (± 11.6)	-	-	-	-
0.1 (< 0.18)	23 (± 1.6)	116.8 (± 14.1)	1	-1	0	-2
0. 32 (0.36)	21 (± 3.2)	100.7 (± 23.6)	5	7	10	14
1.0 (1.1)	22 (± 2.6)	98.4 (± 28.7)	3	10	5	17
3.2 (3.4)	23 (± 3.4)	122.6 (± 16.8)	-1	-4	0	-8
10 (11.4)	11 (± 1.8)	45.1 (± 6.3)	38ª	38ª	60ª	67ª
Positive control	10 (± 1.3)	33.4 (± 5.0)	39ª	53ª	65°	79ª
2 (1.4) mg Zn/L						
NOEC	-	-	3.4	3.4	3.4	3.4
LOEC	-	-	11.4	11.4	11.4	11.4
EC ₂₀ (± 95 % CI)			6.5 (3.5 - 11.3)	6.0 (3.5 - 11.3)	4.3 (3.5 - 11.3)	3.0 (3.5 - 11.3)
EC ₅₀ (± 95 % CI)	-	-	>11.4	>11.4	10.3 (3.5 - 11.3)	8.8 (3.5 - 11.3)

679 an LOD of 0.18 mg ZON/L).

680

681 Footnote - ^a Significantly different (P < 0.05) from control treatment

682 Summary effect values calculated with measured values where possible.

- **Table 3.** Maximal quantum efficiency of Photosystem (Fv/Fm) measured in *P. subcapitata*
- (mean ± SD) after 72 h exposure to zearalenone (CAS number 17924-92-4) in a static study at
- $24 \pm 1^{\circ}$ C (concentration measured using UV-Vis spectrometry with an LOD of 0.18 mg ZON/L).

Nominal	Maximal
concentration,	quantum
measured in	efficiency
brackets	(Fv/Fm)
(mg ZON/L)	
0 (< 0.18)	0.49 (± 0.002)
0.032 (< 0.18)	0.49 (± 0.001)
0.1 (< 0.18)	0.49 (± 0.003)
0.32 (0.23)	0.49 (± 0.004)
1.0 (0.83)	0.50 (± 0.005)
3.2 (3.12)	0.50 (± 0.011)
0.2 (Zn)	0.48 (± 0.009)