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Simulating regimes of chemical disturbance and testing impacts in the ecosystem using a novel programmable dosing system

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4 Simulating regimes of chemical disturbance and 5 testing impacts in the ecosystem using a novel 6 programmable dosing-system

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24

25 *AUTHOR EMAIL ADDRESS:* m.browne@unsw.edu.au26 **Summary**

- 27 1. Pollution is a global issue at the frontier between ecology, environmental science,
28 management, engineering and policy. Legislation requires experiments to determine
29 how much contamination an ecosystem can absorb before there are structural or
30 functional changes. Yet, existing methods cannot realistically simulate regimes of
31 chemical disturbance and determine impacts to assemblages in ecosystems. This is
32 because they lack ecologically relevant species and biotic interactions, are logistically
33 difficult to set-up, and lack environmentally relevant regimes of chemical and abiotic
34 disturbance that organisms experience in polluted areas.
- 35 2. We solved these long-standing environmental, logistical and ecological problems by
36 developing a programmable dosing-system. This dosing-system simulates, *in situ*,
37 regimes of chemical disturbance to assemblages by manipulating the concentration,
38 duration, timing and frequency of pollutants to which they are exposed.
- 39 3. Experiments with priority pollutants (the metal copper and the biocide Chlorpyrifos)
40 and mussel assemblages revealed consistent plumes of contamination within mussel
41 beds. Mussels at the sources of experimental plumes of copper created by the dosing-
42 system accumulated 670% more copper in their tissues compared to mussels 0.5-50 m
43 away. In addition, when mussels were exposed to increasing concentrations of metal
44 there was a concomitant increase in the amount of metals in the tissues of mussels.
45 Combining the dosing-system with an established hierarchy of ecotoxicological assays
46 revealed mussel assemblages exposed to copper and/or Chlorpyrifos (had 40-70%

47 fewer worms, whilst biocides caused 81% fewer amphipods and mussels to filter 48%
48 less water. Combinations of copper and/or Chlorpyrifos had no effects on the
49 abundance of crabs, the respiratory functions of assemblages or the viability of
50 molluscan haemocytes.

51 4. As global contamination accelerates we discuss how this technological advance will
52 enable a diverse array of ecologists, managers and policy-makers to understand and
53 reduce pollution.

54

55 **Key-words:** *In situ*, levels of biological organization, cell, assemblage, filtration, pollution,
56 multiple stressors, environmentally relevant, ecologically relevant.

57

58 Introduction

59 Global infiltration of chemical contaminants causes global reductions in biodiversity (Johnston
60 & Roberts 2009) that jeopardize the useful functions and services that ecosystems provide.
61 Mitigating these problems is difficult because humans use >99 million different chemicals
62 (with 15,000 new chemicals added each day) and <0.3% of these chemicals have any form of
63 regulation (CAS 2015). Legislation (EC 2006; EPA 1980) requires experiments to determine
64 the quantity of a pollutant (known in the US as a “*Total Maximum Daily Allowance*” or
65 elsewhere as “*Environmental Quality Standard*”) an organism or its ecosystem can safely
66 absorb before there is a change in its structure and function. For 30 years, however,
67 experimental methods to determine these “critical loads” (Groffman *et al.* 2006) have been
68 criticised for (i) lacking ecologically relevant species and biotic interactions (Kimball & Levin
69 1985; Underwood 1995; Underwood & Peterson 1988), (ii) being logistically difficult to set-
70 up, and (iii) lacking environmentally relevant regimes of chemical, abiotic and biotic
71 disturbance that organisms experience in polluted areas. We explain these long-standing
72 global issues and provide a novel technological solution in the form of a programmable dosing-
73 system that simulates, *in situ*, regimes of chemical disturbance to assemblages.

74

75 Ecological problems

76 Laboratory experiments provide useful information about the uptake and the sensitivity of
77 organisms to contaminants; however, toxicological testing in laboratories is not the ecological
78 testing of toxicology (Underwood & Peterson 1988) that is required under commitments to
79 international legislation (EC 2006; EPA 1980) for three reasons. Firstly, experiments are
80 usually based on a single or few species that survive easily in the artificial conditions of the
81 laboratory and are thought to be sensitive to contaminants but are not always endemic in
82 polluted habitats. Secondly, experiments rarely study more than one species at a time and it is
83 well known that individual organisms often respond to contaminants in different ways to

84 populations and assemblages (Browne *et al.* 2015). Thirdly, important processes and
85 interactions that normally occur under natural conditions are lacking in the laboratory (e.g.
86 recruitment, competition, emigration; Underwood 1995; Underwood & Peterson 1998).
87

88 *Logistical problems*

89 It is impractical to do realistic experiments with assemblages in most laboratories so the
90 solution has been to use field experiments to determine whether effects actually arise over and
91 above the natural variation that is explicitly excluded in the laboratory (Kimball & Levin 1985).
92 In laboratory experiments (including micro/mesocosms) concentrations of contaminants are
93 manipulated and naturally varying confounding variables (e.g. temperature, salinity, light,
94 food) are kept artificially constant. In contrast, field experiments expose organisms to natural
95 variations in all environmental variables (e.g. tidal-cycle, spates, storms, droughts, predation,
96 competition) except the one of interest (e.g. dose of contaminant), which is manipulated *in situ*.
97 Where direct comparisons have been made, findings from field experiments have often been
98 inconsistent with those from laboratories (e.g. Thompson, Norton & Hawkins 1998). Closer
99 examination reveals that laboratory experiments can suffer from significant artefacts (e.g.
100 transferring organisms from habitats and caring for them in laboratories; Honkoop *et al.* 2003;
101 Dissanayake, Galloway & Jones 2008) that alter the performance and survival of organisms.
102 For instance, differences in their diet (Dissanayake *et al.*, 2009), architecture of aquaria (e.g.
103 glass vs plastic tanks; Teuten *et al.* 2007), water-quality (e.g. salinity, pathogens, gradients of
104 temperature) can affect whether or not a contaminant transfers into their tissues and impacts
105 the functional well being of an organism (Camus *et al.*, 2000; Fischer 1986; Parry & Pipe
106 2004; Roast *et al.*, 2001). To overcome some of these ecological, physiological and logistical
107 problems researchers do field experiments, here assemblages are exposed to doses of
108 contaminants *in situ* using a range of dosing-systems. These include fences that trap
109 contaminants (e.g. oil; McGuinness 1990), paints (e.g. anti-foulants; Johnston & Webb 2000)

110 and impregnated plaster-blocks that dissolve with the movement of water (e.g. Morrisey et al.
111 1995; Cartwright, Coleman & Browne 2006) or electronic pumps that deliver pulses of
112 contaminants (Roberts *et al.* 2008). These dosing-systems, however, also suffer from logistical
113 and financial problems. Fences cannot contain all contaminants and may alter biotic
114 interactions, the plaster itself can affect the physiology and behaviour of animals (Cartwright,
115 Coleman & Browne 2006) and existing pumps have to be operated manually. Where necessary
116 procedural controls are included, they can be costly and time-consuming. Furthermore,
117 running field experiments can be difficult due to tides and spates in aquatic habitats, poor
118 weather, a lack of electricity and running-water, and a lack of security can make the risk of
119 losing sophisticated and expensive equipment unacceptably large, especially in busy areas.
120 Therefore, there is a need for inexpensive, safe and autonomous technologies for manipulating
121 the dose of contaminants that can be deployed across aquatic and terrestrial habitats.

122

123 *Environmental problems*

124 Monitoring shows organisms in habitats experience complex regimes of chemical disturbance
125 with the concentrations, timing and frequency of toxicants to which these organisms are
126 exposed (e.g. transient contamination events through seasonal urban run-off; Maekpeace,
127 Smith & Stanley 1995; Church, Granato & Owens 1999). These regimes are expected to
128 become more complicated due to accelerated chemical production and climatic change
129 (Schiedek *et al.* 2007). This is problematic because variation (spatial and temporal) in
130 disturbances is known to cause variation in ecological impacts (Benedetti-Cecchi 2003) and
131 existing field and laboratory technologies are not able to expose organisms to environmentally
132 relevant regimes of chemical disturbance. Therefore, to fulfill legislative commitments (EC
133 2006; EPA 1980) to determine whether or not the ecosystem can safely absorb one or more
134 contaminants, scientists need technologies that allow them to expose assemblages to
135 environmentally relevant treatments based on observations (from monitoring) about the

136 concentrations, timing and the frequency at which assemblages in habitats are exposed to
137 chemical toxicants.

138

139 *The technological solution to understanding pollution*

140 To overcome the ecological, logistical and environmental problems with the existing dosing-
141 systems described above we developed a novel - programmable and automated – dosing-
142 system that allows ecologically and environmentally relevant testing of single and multiple
143 pollutants on natural systems by being able to manipulate: (i) the number of chemicals, (ii)
144 their concentration and the (iii) duration, (iv) frequency and (v) timing of influxes over
145 extended periods. As metals and biocides make up over 50 % of the priority pollutants we
146 explored their single and combined impacts on the structure and functions of mussel
147 assemblages. These assemblages are ideal because mussels are important ecosystem engineers
148 with clear links between their “health” and ecological impacts (Browne *et al.* 2015). For
149 instance, their shells create habitat and their feeding transfers sediments, algae, nutrients and
150 energy from the water-column into the mussel bed allowing them to support diverse
151 assemblages (e.g. Cole & MaQuadid 2010). Separate laboratory studies have shown that
152 exposure to pollutants can damage their cells (i.e. fragmented plasma membranes, smaller
153 lysosomes) and organs (i.e. gills, gut) and this has been linked to reduced feeding, respiration,
154 growth and survival, and less diverse assemblages (Browne *et al.* 2015). No field experiment
155 has, however, tested these predictions by manipulating the presence and absence of metals
156 and/or biocide in an ecologically and environmentally relevant manner.

157

158 **Materials and methods**

159 *Dosing-system and location of study*

160 The dosing-system was built by attaching up to ten 640 L tanks (IBC;
161 www.smithsofthedean.co.uk), 100 separate lengths of 100 m (10 km in total) polyethylene

162 tubing (diameters: 4 mm internally, 6 mm externally) coiled on plywood reels (200 mm
163 flange/barrel-diameters; www.northeastreel.com) for dispensing, digital-timers (TM-619,
164 Wenzou Changzin Electronics, www.aquavolt.com.au), batteries (Banner, AccuPro 12v-7A)
165 and charger as an electrical back-up in case the power-supply failed (Fig 1E-H). The dosing-
166 system discharges $4.88 \pm 0.5 \text{ L h}^{-1}$ to each plot and neither coiling the tubes ($F_{1,8}=1.22$, $P =$
167 0.30) nor using 3 different chokes (plastic collars made from irrigation micro-sprayers that
168 reduce the internal diameter of the tube; $F_{3,36}=0.86$, $P=0.47$) significantly reduced the rate of
169 discharge. At full capacity, the system can deliver controlled doses of contaminants to either
170 (i) 200 individual experimental plots, with 25 different treatment-combinations, or (ii) by
171 combining smaller plots, an area the size of 100 m^2 could be dosed, and (iii) temporally, the
172 duration of dosing could be manipulated to the nearest minute, hour, day, week, month or
173 indefinitely. This is because the timer(s) can be programmed to automatically start and stop at
174 the pre-programmed times to simulate transient and more prolonged discharges of
175 contaminants without the need for persons to be at experimental sites to operate it.
176 With permission from Malahide Marina and Fingal County Council we completed a series of
177 experiments to determine the capacity of the dosing-system to manipulate levels of
178 bioaccumulation and test the ecotoxicological impacts of pollutants (metals and/or biocides) on
179 multiple levels of biological organization (cell to assemblage). The system was deployed on
180 the floating pontoons on the Northern and Southern pontoons of Malahide Marina (Ireland;
181 Figure 1) as the marina has running-water, electricity, and a security system that prevents
182 vandalism and theft. Here mussel-assemblages living on the vertical-side of the floating
183 pontoons were used as part of our experiments.

184

185 *Experiment 1: size of plume*

186 A copper solution ($0.423 \text{ mg Cu L}^{-1}$ seawater, $\text{CuSO}_4 \cdot (\text{H}_2\text{O})_5$, CAS N^o 7758-99-8) was
187 continuously added to single patches (each $>1\text{m} \times 0.5\text{m}$) of mussels in two sites for 14 d

188 (northern and southern pontoons). Mussels at these sites live on the floating pontoons where
189 they form larger patches that cover the entire length of the floating pontoons. Copper was used
190 as a tracer in this study because it is recognized globally as a priority pollutant (EC 2006; EPA
191 1980), with inputs from run-off, discharges (e.g. sewage, storm-water), corrosion of
192 infrastructure and anti-fouling paint on boat-hulls (Makepeace, Smith & Stanley 1995).
193 Mussels (*Mytilus edulis*) growing on the floating pontoons of the marina allowed us to measure
194 bioaccumulation of pollutants discharged from the dosing-system. These organisms are
195 relatively resistant to many pollutants that accumulate in their tissues, thereby increasing
196 concentrations to levels more easily detected than those in the environment (Widdows &
197 Donkin 1992). This has allowed researchers to use the species complex as ‘sentinel’ organisms
198 to investigate global levels of contamination (Widdows & Donkin 1992). Mussels (after 14 d)
199 and water samples (100 mL; after 7 and 14 d) were collected at 0, 0.25, 0.5, 1, 2, 4, 6, 10, 50 m
200 in both directions along the pontoons from experimental point-sources (Figure 4). Mussels
201 were left in seawater (Instant Ocean; 18M Ω cm water with <0.015 ppm Cu) for 24 hr so that
202 they evacuated their guts and were then frozen in pre-cleaned 100 mL polyethylene bags.
203 Copper concentrations in tissues of mussel and seawater were determined using an Inductively
204 Coupled Mass Spectrometer (see chemical analysis).

205

206 *Experiment 2: manipulating concentrations of pollutant*

207 The dosing-system was deployed on the Northern Pontoon with three treatments (0, 50 and
208 1269 $\mu\text{g mg Cu L}^{-1}$ seawater) housed in separate tanks. The second treatment was
209 representative of concentrations from anti-fouling paints and the third treatment is the largest
210 mean concentration found in European stormwater (Makepeace, Smith & Stanley 1995).
211 Solutions from each tank were added to randomly chosen mussel patches (110 mm in
212 diameter) for 6 weeks and concentrations of copper quantified as previously mentioned. To

213 avoid cross-contamination, each replicate/treatment combination was >1.5 m apart, in this and
214 the remaining experiments.

215

216 *Experiment 3: impacts across levels of biological organization*

217 To determine whether the dosing-system can be combined with a suite of ecotoxicological
218 assays to determine changes in the structure and functions of assemblages we used our system
219 to discharge from five tanks (one tank with seawater; two tanks with 7.7 mg copper sulphate L⁻¹
220 seawater; two tanks with 250 mg chlorpyrifos L⁻¹ seawater) to five groups of experimental
221 mussel assemblages (control; +seawater; +metal; +biocide; +metal+biocide). One day later we
222 quantified changes in the structure (numbers of worms, amphipods and crabs) and functions
223 (viability of haemocytes from mussels, filtration and respiration) of mussel assemblages using
224 established techniques (see ecotoxicological assays). We chose one day for logistical reasons
225 and because previous ecotoxicological work that has used this duration (see Wu et al 2005;
226 Canty et al. 1997) to show metals, biocides and other pollutants causing these types of impacts
227 across levels of biological organization.

228

229

230 *Chemical analysis*

231 This was performed under ISO 9001:2008 protocols at Plymouth University. All glassware and
232 plastic containers were cleaned in a phosphate free degreasing soap-free detergent (10 %
233 solution of Decon 90), then rinsed in water (18 MW cm conductivity), followed by immersion
234 in a nitric acid (10 % Aristar grade by VWR) bath for 24 hr using 18M Ω cm water and rinsed
235 again in 18 MΩ cm water. All items were dried in a particle-free environment and stored in
236 sealed and cleaned polyethylene bags until required. To preserve dissolved Cu and to prevent
237 bacterial/microalgae growth in the samples of seawater, 100 μL of concentrated hydrochloric

238 acid (Sigma-Aldrich, AR grade) was added and samples were stored at <5 °C in the dark. The
239 frozen mussels were freeze-dried (Labconco, Freezone 6) in their storage containers for 3 d.
240 The length, width and height of the shell-valves were measured to the nearest mm using
241 callipers. Mussels were then shucked by removing the mussel from its shell and the mass of
242 the freeze-dried mussel measured (Scalehouse, ALD114CM) and recorded. Each freeze-dried
243 mussel was then transferred to a polyethylene container for digestion with concentrated nitric
244 acid (20-25 °C; 10 mL) (Sigma-Aldrich) for 4 d. The samples were then transferred to boiling
245 tubes for digestion at 100 °C for 48 hr (Skalar, Tecator 1016 Digester Heat-Block). Finally, the
246 acidic solutions containing the digested mussels were transferred to pre-cleaned polyethylene
247 screw-top 100 mL containers (Linnux, UK) and diluted to 10 mL with 18 MΩ cm water for
248 storage prior to analysis. The concentration of copper in the solutions containing the digested
249 mussel was determined using an Inductively Coupled Plasma Mass Spectrometer (X Series 2,
250 Thermo Fisher Scientific, Hemel Hempstead, U.K.). This instrument was operated in
251 ‘collision/reaction cell mode’, with 7 % H₂ in He as the collision/reaction gas, to negate the
252 effect of polyatomic interferences, e.g. ²³Na, ⁴⁰Ar and ²⁵Mg, ⁴⁰Ar on ⁶³Cu and ⁶⁵Cu respectively.
253 All mussel digests were diluted hundred-fold prior to analyses and In and Ir, added to give a
254 concentration of 10 mg per L in the diluted digests, were used as internal standards to account
255 for instrumental drift.

256

257 *Ecotoxicological assays*

258 The viability of haemocytes (cells within the haemolymph that function in the immune system
259 of invertebrates) from mussels was assessed as in Browne *et al.* (2008). This measured the
260 ability of their haemocytes to accumulate a red dye, with ‘healthier’ well-functioning cells
261 accumulating more dye than cells exposed to pollutants. Haemocytes were used because they
262 can be easily collected and play a major role removing harmful waste, supplying tissues with
263 nutrients and energy, healing wounds, degrading pathogens and adding minerals to their shell

264 Moreira et al. 2013). Previous research has shown that exposure to pollutants, however,
265 degrades these functions by damaging the plasma membrane and/or shrinking their lysosome.
266 Haemolymph (50 μ L) withdrawn from abductor muscles of three mussels was placed into
267 duplicate wells of a 96-well microtitre plates (pre-treated with Poly-Lysine). The plate was
268 then agitated (1400 rpm for 60 s) and then left for 50 min for the cells to adhere. Excess cells
269 were then discarded and the wells rinsed with phosphate buffer (pH 7.4). Neutral red dye
270 (0.4%) was then added and cells were incubated in the dark for 3 hr to prevent photolysis.
271 Wells were then washed with phosphate buffer again before a solution of 1% acetic acid/20%
272 ethanol was added to precipitate the dye. Absorbance was read at 550 nm using a
273 spectrophotometer, protein was quantified and results were presented as optical density per
274 gram protein.

275 Respiration transfers oxygen from seawater to the tissues of organisms and carbon dioxide in
276 the opposite direction. To measure this, mussel-patches were incubated *in situ* and in the dark
277 using an opaque chamber. Changes in dissolved oxygen ($\text{mg O}_2 \text{ L}^{-1}$) were measured using a
278 probe (HQ20 Hach Lange Ltd portable LDO™, Loveland, USA). To ensure that
279 measurements were taken at the correct time points, a linearity test was undertaken to test how
280 long it took the volume of water to be depleted of oxygen. Concentrations of oxygen in the
281 water were measured every two minutes over an hour to identify the period during which there
282 was a linear decrease of oxygen in the water. On the basis of this test, measurements of
283 oxygen were taken after 10 and 20 minutes. The initial ten minutes as defined by the linearity
284 test has been shown to also allow for acclimatisation of the assemblages and to ensure
285 photosynthesis had ceased after covering with an opaque chamber (Noël *et al.* 2010). The
286 chamber was also fitted with a pump to ensure water was circulated and that concentrations of
287 oxygen were homogenous and representative. Rates of oxygen uptake by the assemblages were
288 estimated using the equation; $\Delta [\text{O}_2] \text{ dark} / \Delta t \text{ dark}$, $\Delta [\text{O}_2] \text{ dark}$ is the difference in dissolved
289 oxygen concentration between measurements taken respectively at the beginning and end of

290 the dark period and Δt dark is the time difference between these measurements. Respiration
291 calculated for each individual plot every hour and expressed as $\text{mg.O}_2\text{.L}^{-1}\text{.hr}^{-1}$ (Noël *et al.*
292 2010).

293 We then tested the capacity of assemblages to clear particulate matter from seawater *in-situ*
294 using the same purpose-built chamber. 5 mL of a solution containing microalgae (*Isochrysis*
295 *galbana*) was injected into the chamber. The solution of microalgae was prepared such that it
296 gave a concentration of algal cells of 12 -15,000 per 0.5 mL in the chamber. To ensure that
297 microalgae remained suspended in solution, the chamber was fitted with a circulation pump.
298 After initial introduction of algal cells, 20 mL samples of seawater from within the chamber
299 were taken at three time intervals: 0 (T0), 15 (T1) and 30 (T2) min respectively. The numbers
300 of particles retained in samples were counted using flow-cytometry and clearance rates
301 calculated as the change in concentration per unit time using the following equation; clearance
302 rate = $V (\log_e C_1 - \log_e C_2)/t$, where V is the volume of water in the chamber and C1 and C2
303 are the algal concentration at the beginning and end of the time interval (t) (Noël *et al.* 2010).

304 To determine whether the structure of assemblages was affected by the experimental
305 treatments, the numbers of worms, amphipods and crabs found in each patch of mussel were
306 counted across the experimental treatments. Formal comparisons across the experimental
307 treatments were made using total numbers in these broad taxonomic groups and were not based
308 on the numbers of species present.

309

310 **Results**

311 The dosing-system consistently produced small concentrated plumes of contamination within
312 mussel beds on the outsides of both pontoons (Experiment 1) and the plume could not be
313 detected in water-samples from outside the mussel bed (Figure 2b; Table 2). Mussels at the
314 sources of experimental plumes created by the dosing-system accumulated 670% more copper
315 in their tissues compared to mussels 0.5-50 m away (Figure 2a; Table 1). This shows

316 experimental units in ecotoxicological experiments with this dosing-system need to be
317 separated by more than 0.5 m to avoid cross-contamination. There were no detectable
318 increases in the concentrations of copper in seawater outside the exposed mussel patches,
319 indicating that the copper was either taken in by the mussels, transported away from the
320 pontoon or precipitated out. In addition, when mussels were exposed to increasing
321 concentrations of copper, there was a concomitant increase in the amount of copper in the
322 tissues of mussels (Experiment 2; Figure 2c). Combining the dosing-system with an
323 established hierarchy of ecotoxicological assays (Experiment 3; Figure 3) revealed mussel
324 assemblages exposed to metals and/or biocides having 40-70% fewer worms ($F_{1,12} = 8.11$, $P <$
325 0.05 ; including the species *Pomatoceros lamarcki*, *Cirratulus cirratus*, *Nereis pelagica*,
326 *Neanthes irrorata*, *Capitella capitata* and *Manayukia aesturina*) and those exposed to biocides
327 having 81% fewer amphipods ($F_{1,12} = 6.51$, $P < 0.05$; including the species *Ampithoe*
328 *gammaroides*, *A. rubricata*, *Corophium volutator*, *Leucothoe spinicarpa*, *Gammarus chevreuxi*,
329 *G. zaddachi*, *G. duebeni*, *G. salinus*, *Hyale nilssoni*, *Melita palmata*, *M. obtusata* and
330 *Sunamphitoe pelagica*) with mussels that filtered 48% less water ($F_{1,12} = 8.11$, $P < 0.05$).
331 Combinations of pollutants had no effects on the abundance of crabs, the respiratory functions
332 of assemblages or the viability of molluscan haemocytes.

333

334 **Discussion**

335 The dosing-system allows the number of chemicals, their concentrations and the duration,
336 frequency, timing and spatial scale of influxes to be manipulated *in situ*. By combining the
337 dosing-system with a hierarchy of ecotoxicological assays we showed the practical value of our
338 system for generating environmentally and ecologically relevant data about the capacity of
339 pollutants to affect the structure and functions of assemblages. In our experiments the
340 respiratory functions of mussel-assemblages did not change despite reductions in the numbers
341 of organisms contributing to the respiration. Because there was no observed mortality in the

342 mussels across experimental treatments, it seems plausible that the capacity of mussel-
343 assemblages to respire (by consuming oxygen and releasing carbon dioxide) is probably
344 largely regulated by mussels, rather than the other organisms. Given that short-term exposure
345 to the pollutants elicited behavioural and ecological changes in the assemblages with no
346 reductions in respiration and the viability of molluscan haemocytes (an established biomarker
347 used in programmes of monitoring) work is clearly needed to test the capacity of sub-lethal
348 biomarkers of pollution to forecast these ecological impacts (Forbes, Palmqvist & Bach 2006;
349 Browne *et al.* 2015). Our dosing-system would allow scientists for the first time to
350 experimentally determine whether environmental assessments made at lower levels of
351 biological organisation indicate the progress of recovery processes at higher levels.
352 This is important because the use of biomarkers is largely based on single-compound
353 exposures carried out in the laboratory and compounds are assumed to have additive or
354 independent effects in habitats with mixtures of contaminants (Forbes, Palmqvist & Bach
355 2006). In our experiment, combinations of pollutants had no effects on the abundance of crabs,
356 the respiratory functions of assemblages or the viability of molluscan haemocytes. Exposure to
357 Chlorpyrifos, however, caused assemblages to filter 48% less water. A number of models
358 could explain this observation (e.g. mussels could be sensitive to Chlorpyrifos, in terms of
359 chemoreception or paralysis, so do not filter during its presence; whilst copper could affect the
360 bioavailability and toxicity of Chlorpyrifos to mussels) but they require further experiments
361 with detailed chemical analyses. Experiments are also needed to determine the ecological
362 and/or toxicological mechanisms causing the observed reductions in the numbers of worms and
363 amphipods in our experiment.

364 Pollution is often a transboundary problem and given our dosing-system's flexibility and
365 environmental realism (Table 3) it could be used across continents and habitats (i.e. freshwater,
366 marine and terrestrial habitats), as part of coordinated and distributed experiments (Fraser *et al.*
367 2012), to simulate and test the impacts of current (Church, Granato & Owens 1999) or

368 predicted regimes of chemical disturbance. Experiments in terrestrial habitats would be
369 straightforward providing there was sufficient access to electricity and water, and the spatial
370 extent of the plume in soil, air and organisms was quantified to ensure adequate spacing of
371 experimental plots. Our dosing-system therefore has practical applications and implications
372 for scientists, managers and policy-makers, in relation to international environmental trends,
373 risk assessment and the evaluation and design of environmental regulations to prevent
374 pollution. For instance, by identifying the location, nature and extent of ecotoxicological
375 impacts caused by particular pollutants at a global scale, it will reduce the considerable
376 uncertainty about the types and concentrations of pollutants that can be safely absorbed by
377 ecosystems. This will enable managers to prioritize resources for cleaning-up degraded
378 habitats and allow policy-makers to develop legislation that eliminates the production of
379 problematic chemicals in favour of safer alternatives (Rochman et al. 2013). Consequently,
380 our dosing-system will allow a broad-range of scientists (ecologists, industrial ecologists,
381 ecotoxicologists, conservation biologists, environmental scientists), managers and policy-
382 makers to fulfil legislative requirements to understand, forecast and reduce pollution in a
383 changing world.

384

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393

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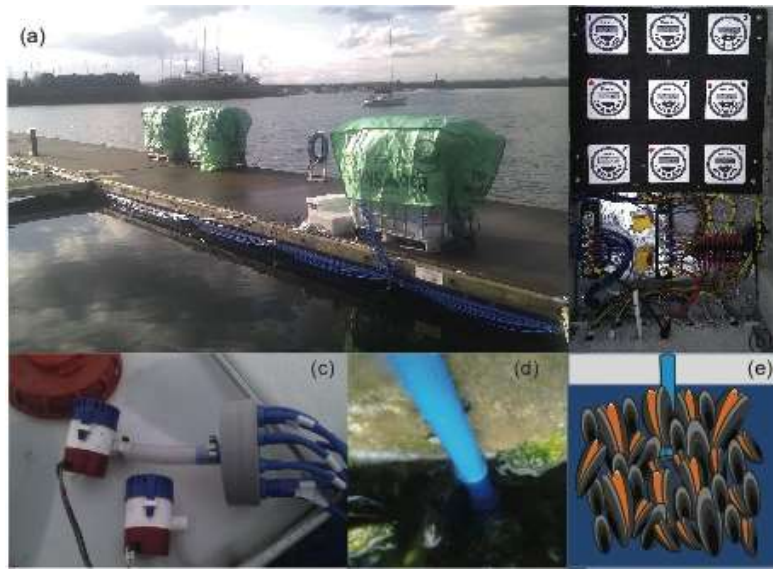
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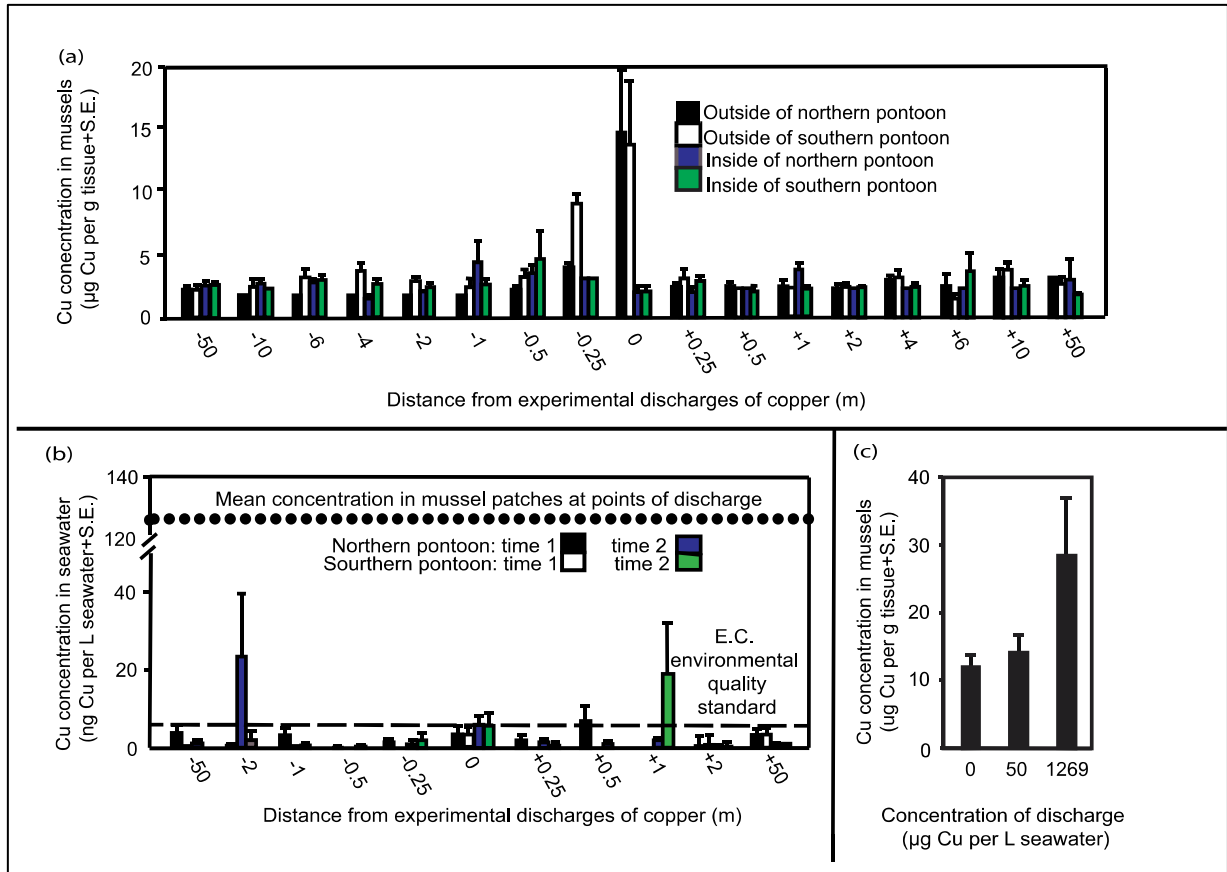
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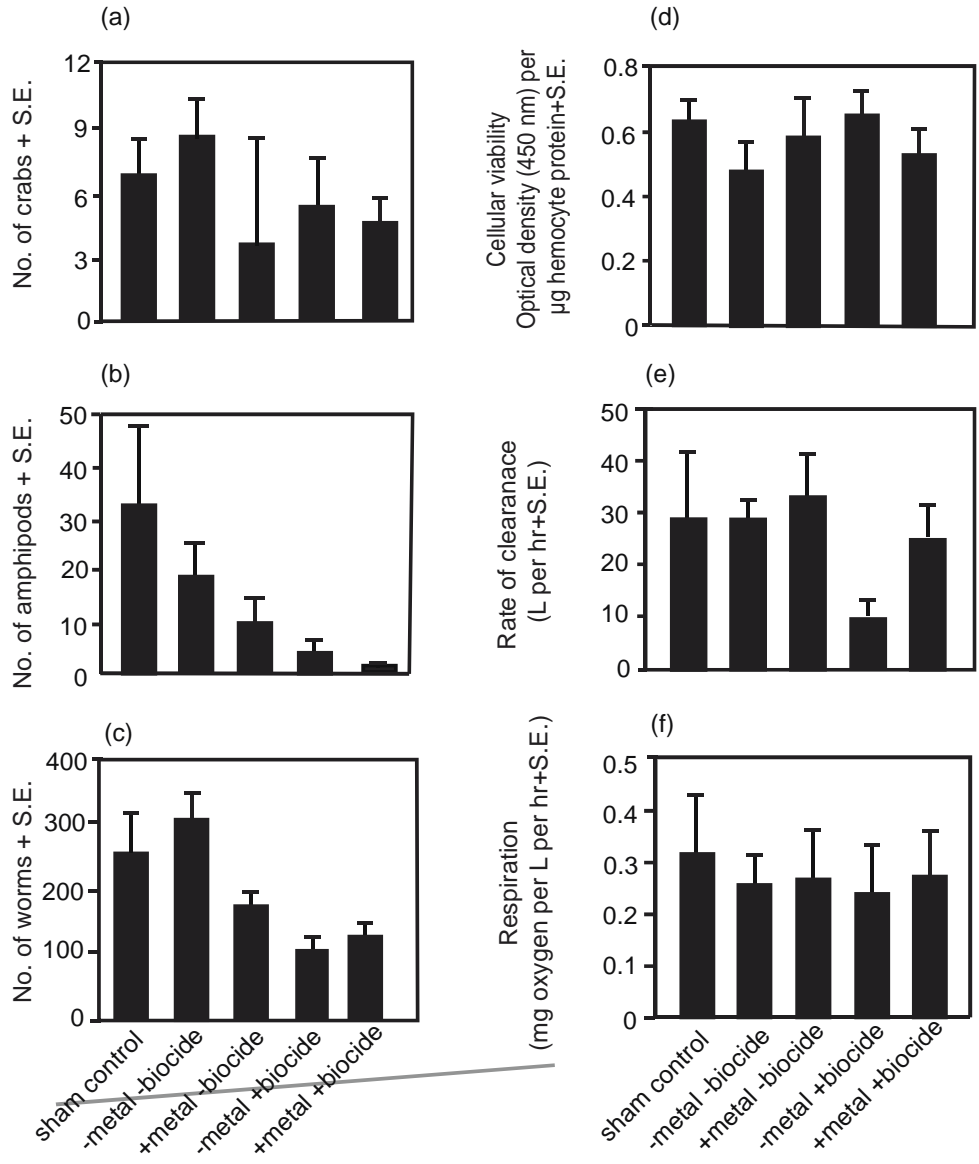
499 FIGURES



500
501 Figure 1. Dosing system (a) set-up at Malahide Marina (53N 27' 17.5; 6E 9' 12.46) with the
502 white box containing the programmable timers (b) each connected to bilge-pumps which
503 themselves are connected to the blue-tubing via the grey manifolds (c). Through this, mussel
504 assemblages on the vertical-side of the pontoon received controlled doses of pollutants (d, e).
505

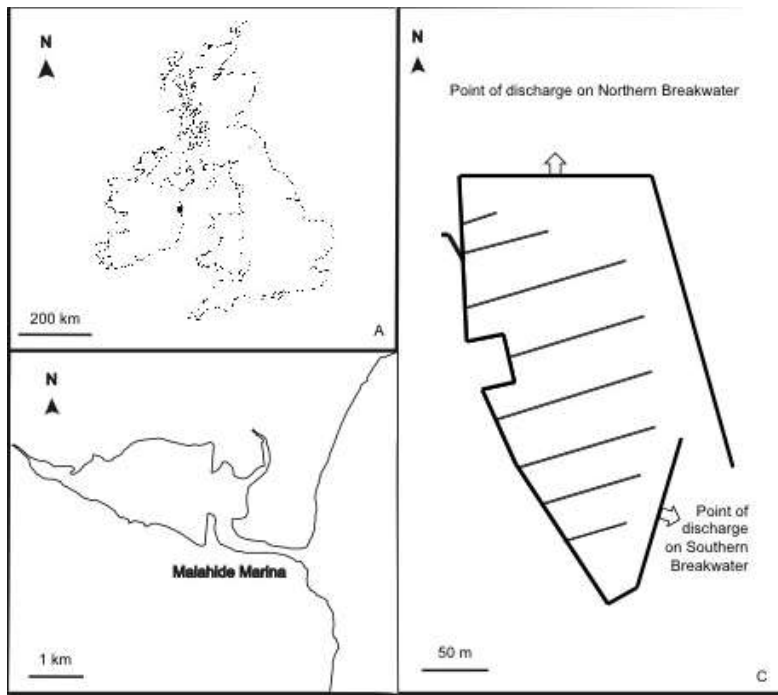


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 507
 508 Figure 2. Spatial distribution of copper in mussels (a) and seawater (b) at different distances
 509 from experimental discharges (Experiment 1). Not all water samples were analyzed because
 510 the plume could not be detected in water-samples outside of the mussel bed in either direction
 511 (Table 1). In contrast, mussels within the plume had over 670% more copper in their tissues
 512 compared to mussels 0.5-50 m away ($F_{16,136}=3.34$, $P<0.001$; Table 2). (c) Experimental
 513 discharges with increasing concentrations of copper caused increasing concentrations of copper
 514 in the tissues of mussels (Experiment 2).
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516
 517 Figure 3. Impacts of metal and/or biocide on the structure (numbers of crabs, amphipods,
 518 worms; a-c) and functions of mussel assemblages (viability of molluscan haemocytes, capacity
 519 of assemblage to filter the water and respire; d-f).

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523

524 Figure 4. Location of experiments within the British Isles and Malahide Marina.

525

526 **Tables**

527 Table 1. Spatial distribution of copper in the tissues of mussels collected at different distances
 528 from the 2 experimental discharges from the dosing-system. Analysis of variance with 3
 529 factors, including 'Pontoon' (Po) was fixed with 2 levels (North, South); 'Side' (Si) was fixed
 530 and orthogonal with 2 levels (Inner, Outer) and 'Distance' (Di) from experimental discharge
 531 was fixed and orthogonal with 17 levels (+50, +10, +6, +4, +2, +1, +0.5, +0.25, 0, -0.25, -0.5, -
 532 1, -2, -4, -6, -10, -50). There were 3 replicates of each combination. Statistical significance is
 533 denoted by $P < 0.01^{**}$ and 0.001^{***}

535 Source	df	MS	F
536 Potoon = Po	1	1.94	0.58
537 Side = Si	1	24.21	7.25**
538 Distance = Di	16	25.32	7.58***
539 Br x Si	1	5.10	1.53
540 Br x Di	16	3.12	0.93
541 Si x Di	16	28.78	8.62***
542 Br x Si x Di	16	1.84	0.55
543 Res	136	3.34	

544
 545 Cochran's test C = 0.33**
 546 SNK tests Outside of pontoon: 0 m > + 0.25 m > all other distances
 547 0 m: Outside of pontoon > Inside of pontoon
 548 + 0.25 m: Outside of pontoon > Inside of pontoon
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563 Table 2. Spatial distribution of copper in seawater outside of the mussel patches collected at
 564 different distances from the 2 experimental discharges from the dosing-system. Analysis of
 565 variance with 3 factors, including ‘Time’ (Ti) was random and orthogonal with 2 levels (9th
 566 and 14th September 2010), ‘Pontoon’ (Po) was fixed and orthogonal with 2 levels (North,
 567 South); ‘Distance’ (Di) from experimental discharge was fixed and orthogonal with 11 levels
 568 (+50, +10, +6, +4, +2, +1, +0.5, +0.25, 0, -0.25, -0.5, -1, -2, -4, -6, -10, -50). There were 5
 569 replicates of each treatment and statistical significance is denoted by $P < 0.001^{***}$.

570

571	Source	df	MS	F
572	Time = Ti	1	93.64	2.61
573	Pontoon = Po	1	60.07	5.69
574	Distance = Di	10	81.48	0.58
575	Ti x Po	1	10.55	0.29
576	Ti x Di	10	140.97	3.93***
577	Po x Di	10	97.02	0.94
578	<u>Ti x Si x Di</u>	<u>10</u>	<u>102.83</u>	<u>2.87***</u>
579	Res	176	35.88	

580

581 Cochran’s test C = 0.50**
 582 SNK tests Time 2 on Northern Breakwater: +2 m > all other distances
 583 Time 2 on Southern Breakwater: -1 m > all other distances

584

585

587 Table 3. The relative advantages of existing methods used in laboratory and field experiments. Shading refers to potential challenges

588 within set-ups that can cause persistent (grey shading), frequent (blue) or no problems (green).

589

Advantages of different types of experimental set-ups		Laboratory			Field				
		Static	Flow-through	Microcosm	Fences	Plaster-blocks	Paint	Watering cans, pumps	Our dosing-system
Ecotoxicological relevance (biological scales where impacts can be tested)	<i>Suborganismal</i>								
	<i>Organismal</i>								
	<i>Population</i>								
	<i>Assemblage</i>								
	<i>Ecosystem</i>								
Logistical and environmental advantages	<i>Includes natural environmental variation</i>								
	<i>Can test for stress(es) due to translocation of organisms</i>								
	<i>No water-quality problems</i>								
	<i>No husbandry required</i>								
Capacity to simulate and manipulate chemical regime	<i>Concentration</i>								
	<i>Number</i>								
	<i>Timing</i>								
	<i>Frequency</i>								
	<i>Automatic dosing</i>								

