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1 The Regulation of Copper Stress Response Genes in
2 the Polychaete *Nereis diversicolor* during prolonged
3 Extreme Copper Contamination

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9 KEYWORDS: Polychaete, Copper, Transcriptomics, *Nereis diversicolor*, Pollution.

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16 **Abstract**

17 Polychaetes are frequented in toxicological studies, one reason being that some members
18 occupy shallow burrows in sediments and are maximally exposed to the contaminants that
19 accumulate within them. We have been studying one population of the polychaete *Nereis*
20 (*Hediste*) *diversicolor* exhibiting inheritable tolerance to extreme copper contamination in
21 estuarine sediment. Using transcriptome sequencing data we have identified a suite of genes with
22 putative roles in metal detoxification and tolerance, and measured their regulation. Copper
23 tolerant individuals display significantly different gene expression profiles compared to animals
24 from a nearby population living without remarkable copper levels. Gene transcripts encoding
25 principle copper homeostasis proteins including membrane copper ion transporters, copper ion
26 chaperones and putative metallothionein-like proteins were significantly more abundant in
27 tolerant animals occupying contaminated sediment. In contrast, those encoding antioxidants and
28 cellular repair pathways were unchanged. Non-tolerant animals living in contaminated sediment
29 showed no difference in copper homeostasis-related gene expression but did have significantly
30 elevated levels of mRNAs encoding Glutathione Peroxidase enzymes. This study represents the
31 first use of functional genomics to investigate the copper tolerance trait in this species and
32 provides insight into the mechanism used by these individuals to survive and flourish in
33 conditions which are lethal to their conspecifics.

34

35 **Introduction**

36 It is widely accepted that the release of anthropogenic waste into natural waters can,
37 through natural selection, effect fundamental changes on species whose habitat preferences leave
38 them vulnerable to exposure. Waste metals released from mining and quarrying frequently enter
39 aquatic systems through drainage and run-off, and are deposited into the sediment of rivers beds
40 and estuaries during transit to the sea ¹. Species that occupy these sediments are, therefore,
41 highly exposed and in many cases develop an evolutionary tolerance ². These ecological traits
42 offer a unique opportunity to study the methods used by organisms to adapt to extreme
43 conditions of metal excess when normal metal homeostasis is overwhelmed.

44 In some cases the magnitude of contamination events can instigate changes to a
45 population on an evolutionarily short time-scale (e.g. a few hundred years). In this study we refer
46 specifically to the infaunal Polychaete *Nereis (Hediste) diversicolor* (Harbour Ragworm). The
47 life history traits of this species may enhance its ability to quickly adapt to inhospitable
48 conditions, for example, its occupation of the intertidal zone make it naturally tolerant of
49 fluctuating environmental conditions, and its distribution within relatively isolated populations
50 with restricted gene flow may prolong the existence of specific adaptations even after the
51 selective forces have been removed ³.

52 One population, found within Restronguet Creek, a branch of the Fal Estuary, Cornwall, UK
53 occupies sediment that contains extraordinary concentrations of copper and to a lesser extent
54 arsenic, tin and zinc ⁴⁻⁶; a product of the local mining heritage dating back to the 1700's. Copper
55 (Cu), at 5073 parts per million (ppm) ⁴, is an order of magnitude more concentrated than in other
56 measured estuarine sediments in the south west of England and other contaminated areas world-
57 wide ^{7, 8}. Accordingly, the sediment is demonstrably toxic to macro-fauna ^{4, 9}, with the exception

58 that a population of *N. diversicolor* has flourished⁴. The individuals are tolerant to the acute
59 toxic effects of copper, zinc and cadmium^{4, 10-12}, but generally have a smaller size and a reduced
60 fecundity as a trade off towards the metabolic cost of tolerance¹³. The trait is not readily lost or
61 gained in the laboratory^{4, 12, 14} and is heritable in the absence of metal¹¹ indicating an underlying
62 genetic adaptation.

63 Hypotheses for the molecular basis of the Cu tolerance include (1) the recruitment and
64 enhancement of existing Cu handling pathways, leading to an increased rate of detoxification; (2)
65 the enhancement of cellular repair pathways to repair copper-induced cell damage; and (3)
66 physiological traits including an increase in mucus production by tolerant animals to reduce
67 exposure¹². It has been shown that the tolerant individuals take up ambient Cu^{15, 16} and produce
68 insoluble detoxificatory deposits within their tegument^{12, 17, 18}, suggesting enhanced
69 detoxification. Specimens collected from Restronguet Creek have up to 91% of the total body Cu
70 concentration present in the insoluble fraction, compared with up to 30% in organisms collected
71 from sites with unremarkable Cu levels¹⁹. Pook²⁰ found a significant increase in the reduced
72 glutathione (GSH) pool supporting a model in which copper is associated with GSH after
73 entering the cell and Cu-thiol complexes are then metabolised in lysosomes, leading to the
74 formation of the detoxificatory deposits. It is unclear whether this process might be sufficient to
75 prevent copper toxicity or if additional adaptation to copper-induced cell damage might also be
76 important; for example, the role of antioxidant systems and cellular repair pathways is unclear.
77 Furthermore, the mucus secreted by *N. diversicolor* only adsorbs small amounts of dissolved
78 metal ions and may not represent a major defence mechanism¹⁵.

79 Here we describe the first functional genomics-based study of the mechanism enabling *N.*
80 *diversicolor* to survive under perpetually Cu stressed conditions. Contextual data, including

81 metal concentration and copper tolerance, was collected by analysing samples from Restronguet
82 creek in tandem with those collected at nearby reference sites. We report the first transcriptome
83 sequence data for *N. diversicolor*, and the identification of a suite of gene transcripts orthologous
84 to those encoding Cu homeostasis and general stress response proteins including antioxidants
85 and DNA repair enzymes. Transcriptional regulation was measured using real-time PCR,
86 enabling for the first time the regulation of Cu homeostasis to be described in Restronguet Creek.
87

88 **Materials and Methods**

89 **Animal Collection and Husbandry**

90 *N. diversicolor* were collected by hand in April / May 2012 at 3 locations within the Fal
91 estuary system; Restronguet Creek (+50° 12' 32.97", -5° 5' 21.13"), Mylor Bridge, which lies
92 approximately 3.2 Km to the south (+50° 11' 1.05", -5° 4' 31.15") and Cowlands Creek, which
93 lies approximately 3.5 Km to the east (+50° 13' 39.53", -5° 2' 36.47"). Specimens with a wet
94 weight of 400-600 mg were used for experiments. Animals were either (1) snap-frozen in liquid
95 nitrogen on site or (2) maintained in glass aquaria containing clean sediment (Specialist
96 Aggregates; this material contained only trace amounts of metal) and artificial seawater (ASW;
97 Tropic Marin; 16 PSU) at 11 °C with a 12 hour light:dark photoperiod.

98 **Transmission Electron Microscopy**

99 The tegument was removed from anaesthetised animals (in 5 % ethanol / ASW) and fixed
100 in 3 % glutaraldehyde (in 0.2 M phosphate buffer, pH 7.2) for 2 hours; stained with 1 % osmium
101 tetroxide (aqueous) for 1 hour; dehydrated in alcohol; embedded in medium viscosity resin
102 (TAAB Laboratories Equipment Ltd); and thin sections (80 nm) were collected on an ultra-
103 microtome. Sections were pre-stained with uranyl acetate and Reynold's lead citrate (for
104 conventional viewing) or collected on gold grids (unstained) for energy dispersive X-ray
105 analysis. Stained sections were viewed with the JEOL 1400 TEM. Unstained sections for X-ray
106 analysis were viewed on a Philips/FEI CM12 TEM operating at 80kV fitted with an X-ray
107 microanalysis detector (EM-400 Detecting Unit, EDAX UK). Images were recorded using a SIS
108 MegaView III digital camera. Analysis of metal containing granular deposits within the tissue
109 was performed using ImageJ²¹, using the sizing and densitometry functions.

110

111 **Metal Measurements**

112 Sediment cores were collected at each study site (2-5 cm depth), dehydrated in a vacuum
113 oven and sieved with a 500 µm mesh. Then, 0.5 g was acid-digested in near-boiling Aqua Regia
114 for 1 hour. A certified reference material (LGC 6137 Estuarine Sediment) was processed in
115 parallel.

116 *N. diversicolor* (frozen) were ground into a fine powder and 0.5 g was acidified with 10
117 % nitric acid and microwave digested at 200 °C for 10 minutes (ETHOS EZ, Milestone). A
118 certified reference material (NRC TORT-2 Lobster hepatopancreas) was processed in parallel.

119 The concentration of As, Cu, Sn and Zn in the digests was measured using a Varian 725
120 ES ICP-OES (Agilent). A five point calibration curve was constructed using an appropriate
121 series of multi-element calibration standards and quality assurance was provided for by the
122 analysis of a second multi-element standard prepared from a different stock solution and by the
123 analysis of the certified reference materials.

124 **Copper Toxicity Assay**

125 Animals were placed individually into acid washed glass troughs (15 cm x 5 cm x 5 cm)
126 containing 200 mL of ASW and a 10 cm length of 6 mm internal diameter polyvinylchloride
127 tubing (refuge); 11°C; and 12 hour photo-period. After 24 hours the water was replaced with a
128 solution of CuCl₂, renewed every 24 hours. Animals were considered dead if their anterior
129 segments didn't respond to a mechanical stimulus.

130 **RNA Extraction**

131 Animals were rinsed in 0.2 µm filtered ASW, snap-frozen and ground into a fine powder.
132 Total RNA was extracted using Trizol (Ambion) and the RNeasy Mini Kit (Qiagen). Quality
133 assurance was provided for by resolving the RNA on a non-denaturing agarose gel to visualise a

134 sharp ribosomal RNA band, and by measuring the absorbance of the sample using a Nanodrop
135 spectrophotometer (Thermo Scientific). Residual genomic DNA was digested with RNase free
136 DNase (RQ1, Promega) prior to clean-up using the RNeasy Mini Kit. Complete DNA digestion
137 was confirmed by a null result from a Taq polymerase-based PCR amplification reaction using
138 the digest sample as a template.

139 **454 Sequencing**

140 RNA was collected from 12 animals; 6 from Restronguet Creek (copper tolerant) and 6
141 from Cowlands Creek (copper sensitive). The samples were combined in equal proportions and a
142 cDNA library was prepared from 2 µg using the MINT First Strand cDNA Synthesis Kit with
143 oligodT primers, followed by cDNA amplification and Duplex Specific Nuclease-based library
144 normalisation using the TRIMMER and EncycloPCR kits (Evrogen). Normalised cDNA was
145 fragmented by nebulisation followed by adapter sequence ligation and emulsion PCR, and the
146 cDNA library was sequenced on full GS FLX and GS FLX+ Titanium series sequencing plates
147 according to the standard protocol of Roche (Eurofins Genetic Services Ltd).

148 The raw 454 data reads were assembled *de novo*. Adapter sequence trimming, quality
149 filtering and the initial assembly of sequence contigs were carried out using GS De Novo
150 Assembler (version 2.3, Roche). Additional assemblies were generated using Mira²² and Cap3
151²³. For each unique contig open reading frames were filtered using DeconSeq²⁴ and annotated
152 using the blastx programme conducted against the non-redundant protein sequence database
153 (NCBI) with an E-value cut-off of 10⁻⁵.

154 **Gene Expression Analysis**

155 Gene expression analysis was performed using real-time PCR (see below). For the *in situ*
156 analysis we collected at least 20 undamaged individuals from each site and immediately snap-

157 froze them in liquid nitrogen to stabilize the mRNA pool. For each site the frozen specimens
158 were combined and homogenized for RNA isolation. This was repeated on 3 different days,
159 sampling at low tide. For analysis in the absence of metal we placed individuals from each site
160 into clean sediment (as described above) and removed at least 20 individuals every 7 days.
161 Replicate measurements were taken from 3 different holding tanks.

162 **Real Time PCR**

163 Complementary DNA was prepared by oligodT-primed reverse transcription using the
164 Thermoscript RT-PCR System (Invitrogen). The relative abundance of gene transcripts between
165 different individuals was determined by real-time PCR using the CFX96 Real-time PCR
166 Detection System (Biorad) and SYBR green DNA detection chemistry. Transcript sequence-
167 specific primers (designed using Primer 3²⁵) amplify 150-250 bp within the 5' region of the
168 target sequence; see supporting information. All RT-PCR reactions were performed in triplicate,
169 each reaction containing 12.5 μ L of IQ SYBR supermix (Biorad), 7.5 μ L of nuclease free water,
170 2 μ L of forward and reverse primers (10 pmol/ μ L) and 1 μ L of template cDNA; the total reaction
171 volume was 25 μ L.

172 The thermal cycling program was 95°C for 5 minutes followed by 40 cycles of 95°C for
173 15 seconds, 55-60°C for 30 seconds and 72°C for 1 minute. At the end of the programme product
174 specificity was analysed from a dissociation curve (55°C to 95°C at 1°C increments for 5 seconds
175 each). Threshold cycle (C_t) values were determined within the exponential portion of the reaction
176 curves. Relative transcript abundance was calculated by normalising the C_t values for the target
177 to the C_t values for a panel of endogenous reference transcript sequences. A panel of 7
178 endogenous reference sequences were tested, and those that gave consistent results were selected
179 for the analysis of the real-time PCR data; the data presented represents the mean value obtained

180 using each valid reference. The relative transcript abundance was calculated from E^n , where n
181 was the difference between the C_t values obtained and E is the efficiency of the primer set used
182 for the amplification reaction. Primer efficiency was derived as $E = 10^{\frac{-1}{-m}}$ where m is the slope
183 of a linear fit for C_t values obtained from a 10-fold dilution series of cDNA template in Real
184 Time PCR reactions.

185

186 **Results and Discussion**

187 **Heavy metal contamination in the Fal Estuary System and Copper Tolerance in *Nereis***
188 ***diversicolor***

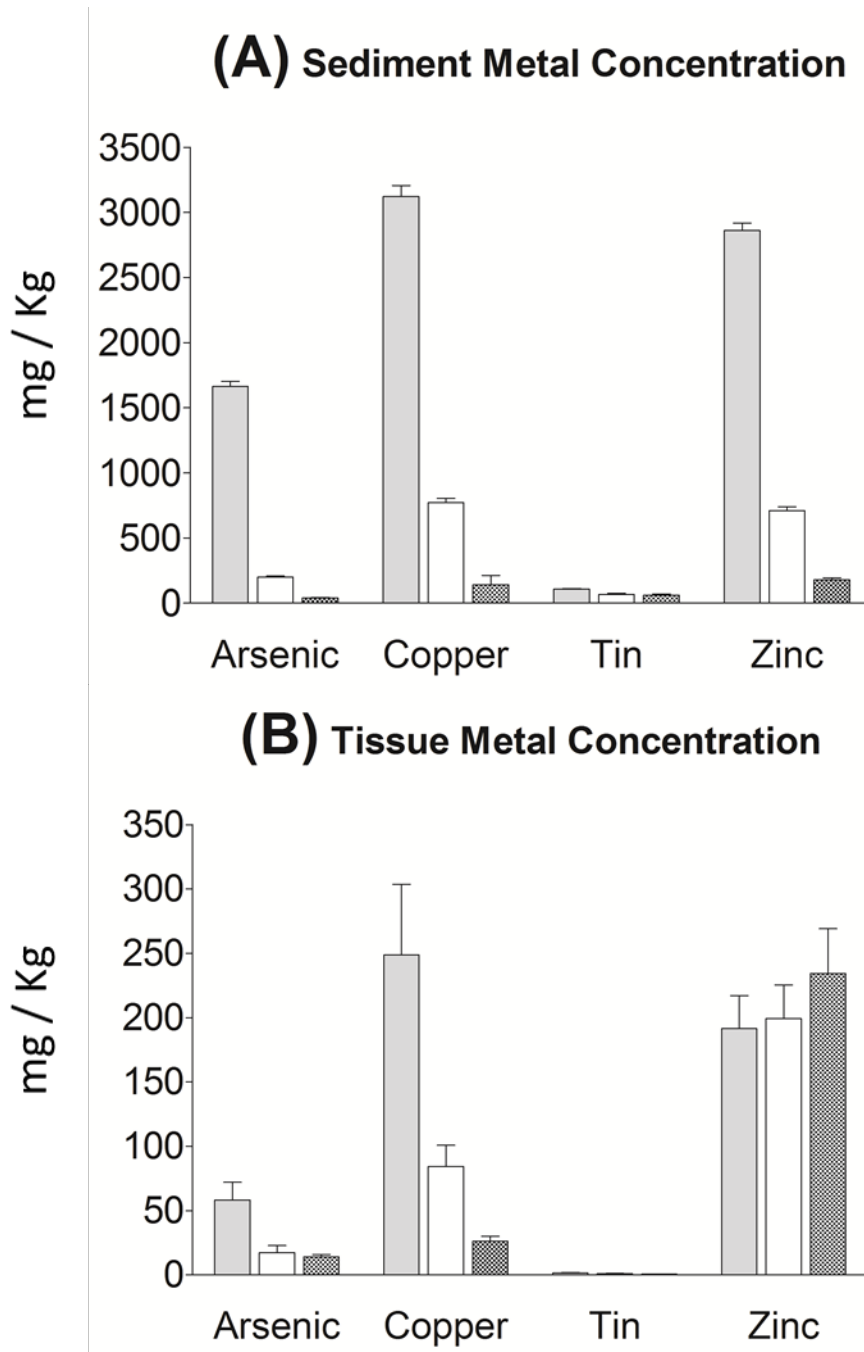
189 We sampled 3 estuarine locations in the south west of England, which contained a rich
190 population of *Nereis diversicolor* (more than 1,000 burrows / m²). These were Restronguet Creek
191 (RC), where metal contamination is famously high ⁴, Mylor Bridge (MB) and Cowlands Creek
192 (CC). The concentration of Arsenic (As), Copper (Cu), Tin (Sn) and Zinc (Zn) in sediment
193 samples was determined by ICP-OES to be in the order Cu>Zn>As>Sn and RC>MB>CC, as
194 shown Figure 1A. At the time of sampling RC sediment contained more Cu than in any
195 previously measured estuarine location within south west England ⁶, the Rio Tinto in Peru ⁷ and
196 the Aznalcollar Tailings in the USA ⁸. We use the Kelly Indices (former GLC guidelines) for
197 contaminated soil (IRCL 59/83) to describe the extent of Cu contamination at each site; RC
198 being unusually heavily contaminated; MB being heavily contaminated; and CC being
199 uncontaminated and therefore used as a reference site for the gene expression experiments. To
200 ascertain the effect of the metal pollution on the ecological diversity of the sediment the benthic
201 macro-fauna was measured according to the method of Simpson ²⁶. Biodiversity at RC was
202 significantly reduced in line with previous determinations ⁴, specifically in the upper reaches
203 where we only found *N. diversicolor* (data not shown).

204 In *N. diversicolor* tissue As, Cu and Sn were accumulated to the same order as the
205 ambient concentration, however tissue Zn concentration showed no significant difference
206 between the populations (t-test, p = > 0.05) despite clear differences in the sediment Zn
207 concentration at each site, Figure 1B. Historical comparison with measurements made by Bryan
208 and Gibbs in 1983 ⁴ (for sediment) and Berthet *et al* in 2003 ¹² (for tissue), wherein the same

209 sampling locations were accessed, indicate that despite a modest increase in the sediment Cu
210 concentration the tissue Cu concentration has decreased by more than an order of magnitude.
211 Recent work by Rainbow *et al*^{15, 16} to quantify the kinetics of Cu uptake by *N. diversicolor*
212 indicates that Cu accumulation rate is proportional to the biologically available copper in their
213 immediate environment, and therefore we conclude that the bioavailable portion of the Cu in RC
214 may have decreased dramatically in the last few decades. Historical data for As, Sn and Zn was
215 unavailable for comparison. Possible mechanisms for the decline in sediment Cu include tidal
216 “flushing” together with reduced inputs after cessation of local mining operations, or changes to
217 the sediment chemistry (e.g. the concentration of dissolved organic matter).

218

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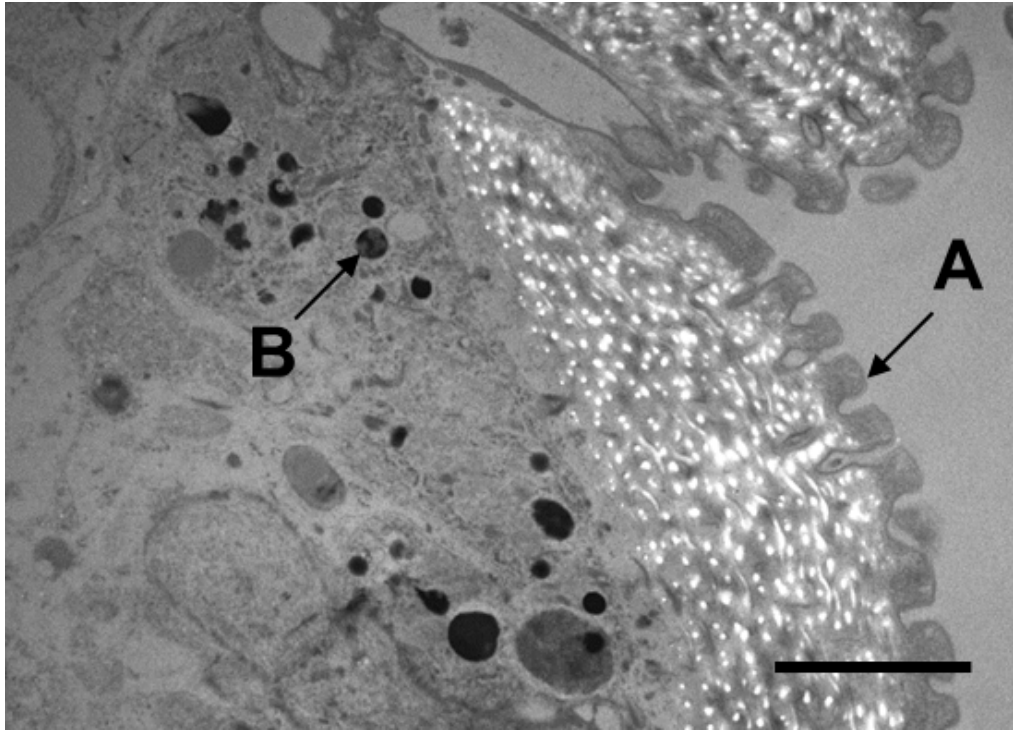


220

221 **Figure 1.** Results of the ICP-OES determination of metal concentration in (A) the sediment and
222 (B) the tissue of *N. diversicolor* in Restronguet Creek (grey bars), Mylor Bridge (white bars) and
223 Cowlands Creek (shaded bars). The error bars represent the standard error on the mean from \geq
224 25 sediment samples or \geq 10 individuals.

225 The fate of the bio-accumulated copper can be investigated using transmission electron
226 microscopy and energy dispersive X-ray spectroscopy, which shows that copper co-localises
227 with sulphur in granular deposits close to the tegument epicuticle ^{12, 18}, as shown in Figure 2. In
228 similar work carried out in 2003 Mourneyrac *et al* ¹² found that Cu tolerant animals from RC
229 presented with an increase in the number and density of the Cu-containing granules indicating a
230 role as detoxificatory Cu stores and a potential role in Cu tolerance. In this study, however,
231 image analysis showed no difference in the number of granules, their size or their density
232 between the populations tested. This corresponds to our comparison of the tissue concentration
233 measured in this study compared to another study carried out using the same sampling location in
234 2003 ¹⁹.

235



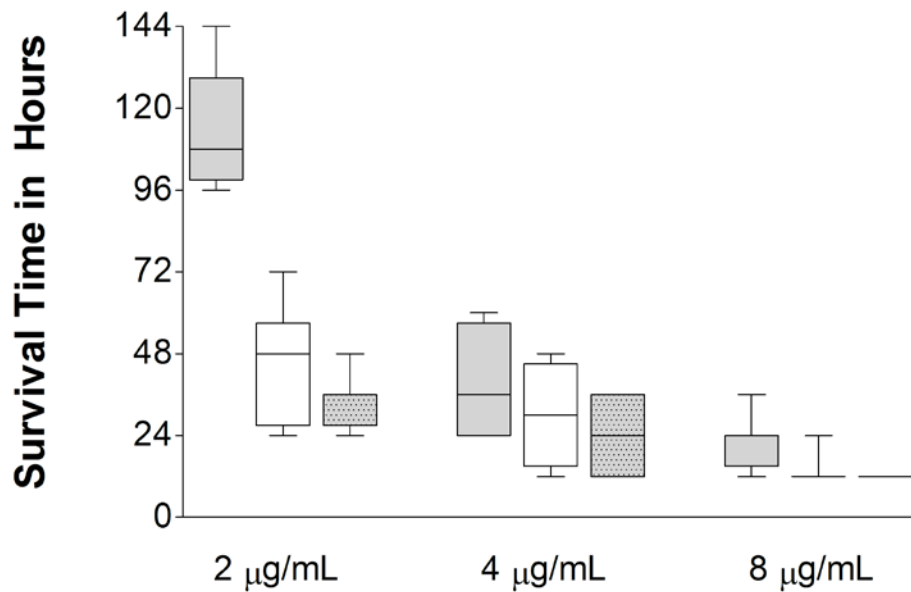
237

238 **Figure 2.** A Transmission Electron Micrograph of a thin sectioned *N. diversicolor* Tegment,
 239 excised from a specimen collected at Restronguet Creek, showing the Epicuticle Layer (A) and
 240 the underlying tissue with Cu / S containing granules (B). Sections were analysed unstained by
 241 energy dispersive X-ray spectroscopy to measure the elemental composition of the granular
 242 structures, which were predominantly copper and sulphur (data provided in the supplementary
 243 information). A typical image is shown. Bar = 2 μ m.

244

245 The Cu contamination in RC has driven the selection of a Cu tolerance phenotype that
 246 was first reported over 30 years ago ⁴. To confirm that this adaptation was still present we
 247 compared the Cu tolerance of animals from the differentially contaminated sediments by placing
 248 individuals into a solution of CuCl_2 and measuring their survival over several days. At a Cu

249 concentration of 2 $\mu\text{g/mL}$ the individuals found at RC were significantly (two-way ANOVA $P \leq$
250 0.05) more tolerant to the CuCl_2 than those found at either MB or CC, as shown in Figure 3,
251 despite the decrease in the tissue burden of Cu indicating that the selective pressure may have
252 diminished. Crucially, the RC model for the study of rapid adaptation and tolerance was still
253 available for gene expression studies.



254
255 **Figure 3.** *N. diversicolor* from the different sites ($n \geq 12$) were placed individually into solutions
256 of CuCl_2 . The survival of individuals from Restronguet Creek (unusually heavy Cu
257 contamination; grey bars), Mylor Bridge (heavy Cu contamination; white bars) and Cowlands
258 Creek (uncontaminated; shaded bars) was measured every 12 hours. The data is presented as a
259 Box-Whiskers plot.

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262

263 **Sequencing and Transcriptome Assembly**

264 To enable a functional genomics-based investigation of the mechanisms underlying the
265 Cu tolerance trait we generated transcriptome sequence datasets using the 454 pyrosequencing
266 technology. A normalised cDNA library was prepared from a pool of the Cu tolerant and Cu
267 sensitive *N. diversicolor* found in RC and CC respectively. The library was sequenced using two
268 sequencing plates (GS FLX Titanium Series, Roche), generating 2,151,516 sequence reads with
269 an average read length of 440 bp and 307 bp of which 1,923,224 (89.4 %) were used for *de novo*
270 assembly of the *N. diversicolor* transcriptome. Read assembly was carried out using multiple
271 assembly algorithms (Newbler, Mira and Cap3; ^{22, 23, 27}) to maximise the potential coverage ²⁸.
272 All sequence data underwent post-assembly filtering using DeconSeq ²⁴ and unique transcript
273 sequences of greater than 300 bp in length were queried against the non-redundant protein
274 database using blastx with an E-value cut-off of 10^{-5} . Manual annotation, where necessary, was
275 carried out using local tblastn searches of the transcriptome assembly data with query sequences
276 corresponding to target genes or proteins. Transcriptome coverage was assessed using Transrate
277 ²⁹ against a set of “eukaryotic conserved” proteins (CEGMA ³⁰). The assembly statistics, blast
278 results for each assembly and transcriptome coverage are summarised in Table 1 (additional
279 information is available in the supplementary material, Table S2). The annotated sequence
280 assemblies were used to identify orthologues to copper homeostasis systems and associated
281 detoxificatory and repair pathways including glutathione metabolism, antioxidants and DNA
282 repair proteins.

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Table 2. Transcriptome Assembly and Annotation Statistics.

Assembly Statistics			
Assembler	Mira	Newbler	Cap3
Unique Contigs (< 300 bp)	60,335	31,770	59,769
Largest (bp)	6801	6996	12,565
Number > 1 Kbp	11,873	14,720	18,343
Mean Contig Length	751	1,065	887
N₅₀	837	1,195	1,018
Number with ORF	16,151	15,442	20,690
Annotation Results			
Blast Database	uniprot_sprot	uniprot_sprot	uniprot_sprot
Blast Program	BlaxtX	BlaxtX	BlaxtX
E<10⁻⁵	23964	15318	25426
E<10⁻¹⁰	20496	13555	21954
E<10⁻³⁰	10339	8301	11897
Transcriptome Coverage			
RBH* (as % of Reference)	0.21 %	0.33 %	0.47 %
Reference Coverage	0.26 %	0.34 %	0.39 %
Number of Chimeras	0	0	0

*Reciprocal Blast Hits

286

287

288 Copper Genes

289 We identified transcript sequences with orthology to copper transporters CTR1 (solute
290 carrier family 31 protein) and ATP7A (P-Type ATPase family protein), which function to move
291 copper across the plasma membrane, into or out of the cell respectively³¹. The ATP7A protein
292 has an additional role in directing intracellular copper into the trans-Golgi network for

293 incorporation into newly synthesised cuproproteins and may relocate to eliminate copper from
294 the cell under conditions of copper excess³². The ATP7 protein in humans has 2 isoforms,
295 ATP7A and ATP7B, but is represented by a single homolog in lower organisms including insects
296³³; and in our polychaete sequences we found evidence for a single ATP7 gene, which had more
297 similarity to the ATP7A isoform. Any free Cu⁺ in the cell is highly toxic, so Cu trafficking is
298 mediated by specialised chaperone proteins³⁴. We identified Nereid orthologues to the protein
299 CCS (Copper Chaperone to Superoxide dismutase³⁵), which delivers Cu⁺ cofactor ions to SOD1
300 (Superoxide dismutase), and ATOX1 (Antioxidant protein 1), which interacts with ATP7³⁶ to
301 support copper export or compartmentalisation.

302 Additional systems augment copper homeostasis. Cellular thiols including those
303 presented on glutathione and metallothionein / metallothionein-like proteins (MTLPs) sequester
304 excess Cu⁺ away from the metabolism and contribute to the formation of detoxificatory stores³⁷,
305³⁸. In addition to the identification of the principle components of glutathione metabolism we
306 found evidence for 2 MTLP genes, encoding a typical (< 200 amino acids) and atypically large
307 predicted protein. To our knowledge MTLPs in this species have, until this data, been
308 unconfirmed. Interestingly, the discovery of 2 putative MTLP genes encoding different sized
309 proteins sheds light on the work by Poirier *et al* who found a bi-modal distribution of Cu within
310 heat stable, size fractionated *N. diversicolor* cytosol¹⁸. Additional analysis of the large MTLP
311 sequence reveals a 404 amino acid protein (confirmed proteomically; data not shown) containing
312 5 repeats each comprising 10 cysteine residues with the consensus GCGC-X₅-G-XX-CC-X-G-
313 XX-C-X₁₁-G-XX-G-XX-C-XX-CX₇CX₅CX₇CX₅GX₄K; the total cysteine composition was 13
314 %. We also looked for the major antioxidant proteins, which neutralise the effect of Cu⁺-
315 catalysed formation of reactive oxygen species. In addition to Catalase, *N. diversicolor* has at

316 least 2 isoforms of superoxide dismutase which are identical, or nearly identical to those found in
317 other polychaete species including *Perinereis nuntia*. Finally, we looked for DNA repair
318 pathway proteins for which we found a complement of candidate sequences. The sequences
319 described in this study are summarised in Table 2.

320

321

322

Table 2. Transcript sequences used in this study.

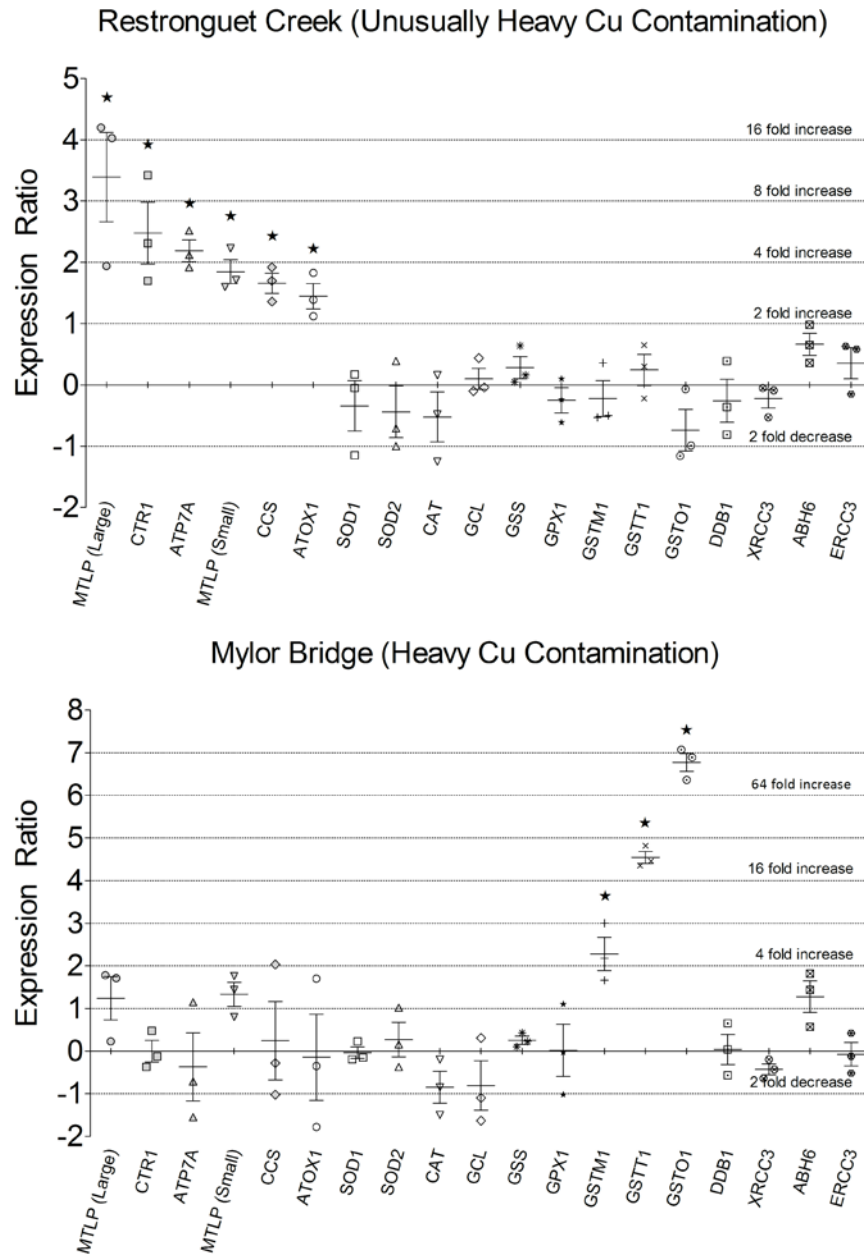
Name ¹	Protein Description	%ID ²	Organism ³	Genbank	E ⁴
Copper Transporters					
ATP7A	Wilson's disease protein; ATPase family	68	<i>Daphnia pulex</i>	EFX90405.1	10 ⁻³⁹
CTR1	High affinity copper uptake transporter; Solute carrier family	36	<i>Harpegnathos saltator</i>	EFN85594.1	3x10 ⁻²⁴
Copper Chaperones					
ATOX1	Copper transport protein	51	<i>Crassostrea ariakensis</i>	AEJ08756.1	
CCS	Copper chaperone for superoxide dismutase	51	<i>Salmo salar</i>	NP_001133786.1	
Metallothionein-like proteins					
-	Cd/Se Metallothionein	55	<i>Schistosoma mansoni</i>	XP_002575981.1	10 ⁻¹³
-	Atypical Metallothionein-like protein	26	<i>Trichomonas vaginalis</i>	XP_001321197.1	9x10 ⁻⁰⁹
Antioxidants					
SOD1	Cu/Zn Superoxide Dismutase	100	<i>Perinereis nuntia</i>	ADM64420.1	3x10 ⁻¹⁰
SOD2	Mn Superoxide Dismutase	80	<i>Perinereis nuntia</i>	ADM64421.1	10 ⁻⁹⁹
CAT	Catalase	78	<i>S. kowalevskii</i>	XP_002738841.1	10 ⁻⁹⁹
Glutathione metabolism					
GCL	Glutamate cysteine ligase	92	<i>Laeonereis acuta</i>	AAV48595.2	10 ⁻¹²²
GSS	Glutathione synthetase	40	<i>Tribolium castaneum</i>	XP_968070.1	2x10 ⁻⁷²
GPX1	Glutathione peroxidase	68	<i>Hyriopsis cumingii</i>	ACY72387.1	4x10 ⁻⁴¹
GSTM1	Glutathione S-transferase (Mu)	56	<i>Reishia clavigera</i>	ACD13785.1	3x10 ⁻³⁵
GSTT1	Glutathione S-transferase (Theta)	84	<i>Neanthes succinea</i>	ABQ82132.1	10 ⁻¹¹¹
GSTO1	Glutathione S-transferase (Omega)	46	<i>Neanthes succinea</i>	ABR24228.1	2x10 ⁻⁴⁸
DNA Repair					
DDB1	Damage DNA binding protein 1	78	<i>Aedes aegypti</i>	XP_001655231.1	5x10 ⁻⁴⁶
XRCC3	DNA excision repair protein	39	<i>Bos taurus</i>	NP_001071585.1	10 ⁻¹¹
ABH6	DNA alkylation repair protein	58	<i>Danio rerio</i>	NP_001005390.1	10 ⁻⁶⁷
ERCC3	DNA repair helicase	74	<i>Ixodes scapularis</i>	XP_002399857.1	7x10 ⁻⁹⁹

323 ¹Name in Homo sapiens; ²% Identity with top blast result; ³Top blast result; ⁴E-value.

324 **Expression *in situ***

325 The expression of these principle copper homeostasis genes was measured *in situ* (snap-
326 freezing the animals at site to stabilise the mRNA pool), by comparing the abundance of mRNA
327 transcripts using real-time PCR. We compared representatives ($n \geq 20$) of the 2 Cu-exposed
328 populations, at RC and MB, with the unexposed reference population at CC on 3 different days
329 during 2011. The results are compiled into a single figure for clarity, as shown in Figure 4.
330 Unsurprisingly, *N. diversicolor* found in RC had significantly increased (one way ANOVA, $p <$
331 0.05) levels of gene transcripts corresponding to the copper transporters ATP7 and CTR1 and
332 copper chaperones ATOX1 and CCS. These genes have an established role in protecting the cell
333 from excess copper and their up-regulation in RC probably reflects the elevated sediment copper
334 as shown in Figure 1A. Up-regulation of CTR1, which encodes a copper uptake transporter, is
335 consistent with previous data indicating that the animals living in RC have increased Cu uptake
336 rates despite the Cu-contamination^{15, 16}. Our current theory is that the enhanced Cu uptake can
337 protect the organism from Cu damage by increasing the rate of elimination of Cu in their
338 immediate surroundings as it moves through detoxificatory pathways including the formation of
339 the Cu / S containing granules^{12, 18}. This process makes *N. diversicolor* a candidate for the
340 bioremediation of contaminated sediments, as noted by others³⁹.

341 Putative MTLP transcripts were also significantly more abundant in RC; specifically the
342 atypically large MTLP sequence showed up-regulation by up to 16-fold compared to individuals
343 from the reference site. Conversely, we found no significant difference between RC and the
344 reference site in the amount of mRNA transcripts corresponding to the antioxidant proteins and
345 the DNA repair enzymes.



346

347

348 **Figure 4.** The *in situ* abundance of transcripts in representatives of the Cu contaminated
 349 populations in comparison to the uncontaminated reference population. The analysis was
 350 performed on a pool of at least 20 individuals on 3 different days. The horizontal line shows the
 351 mean expression ratio versus the reference, the error bars are the SEM from 3 replicate
 352 measurements (days) and the symbols show the values obtained for each replicate. Statistically
 353 significant differences (one way ANOVA, $p < 0.05$) are marked with a star.

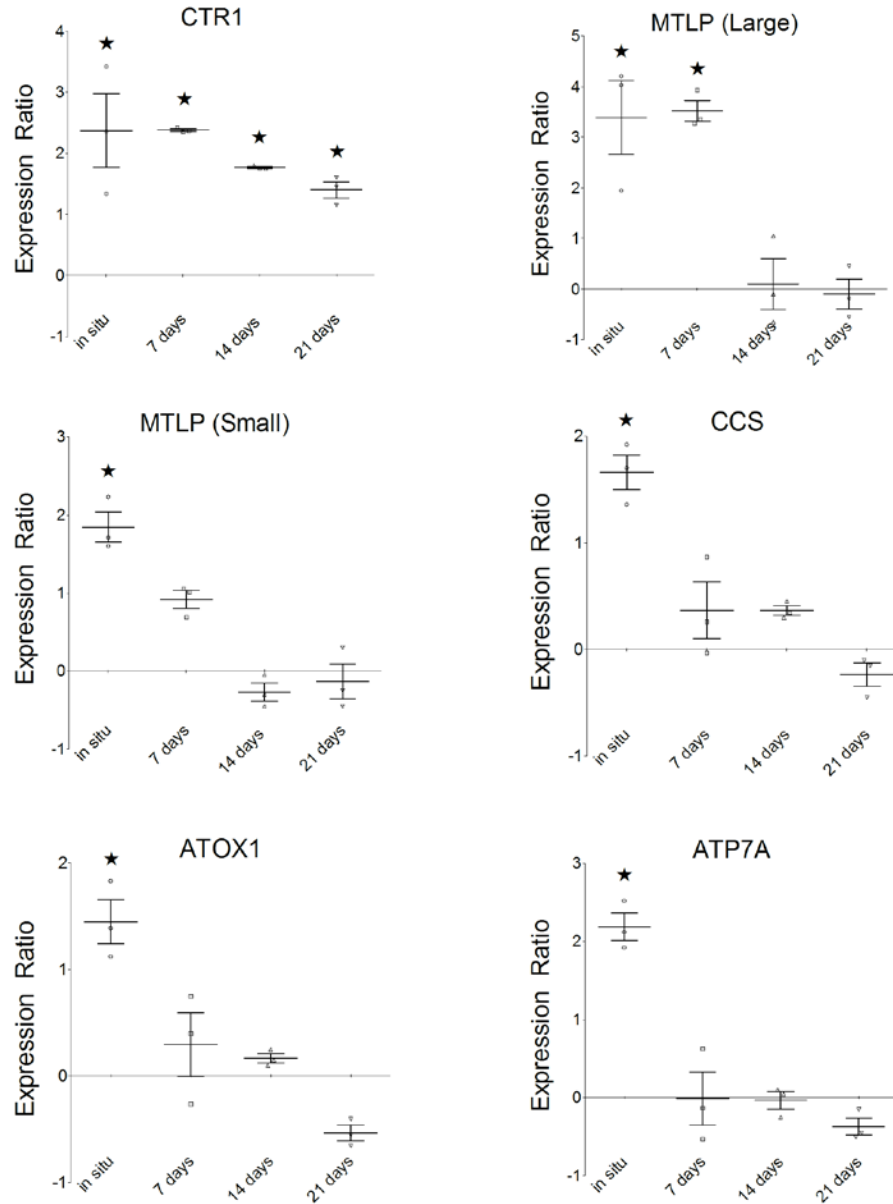
354 It is curious that the gene expression data gave no evidence for the activation of these
355 systems which have an established and significant role in protecting the cell from the effects of
356 heavy metals ⁴⁰. Our previous biochemical assay data ²⁰, comparing the activity of Superoxide
357 dismutase and Catalase between animals in RC and the reference population, agreed with our
358 genetic data that the exposed / tolerant individuals do not experience additional oxidative stress.
359 In contrast, exposure of *Laeonereis acuta* (Nereididae), to Cu led to increased activities of
360 Superoxide dismutase, Catalase, and Glutathione S-transferase ⁴¹, and enhanced antioxidant
361 functions have been associated with copper exposure in a number of other annelid species (for
362 example ⁴²). In this instance it is possible that the hyper activation of the Cu detoxificatory
363 pathways, conferring the tolerance trait, is sufficient to mitigate any Cu-catalysed radical
364 formation. It then follows that DNA repair pathways are similarly unaffected by the high
365 ambient Cu levels.

366 The population in MB had also been exposed to a high concentration of Cu in the
367 sediment but this was far lower than the extreme concentration measured in RC, and these
368 individuals did not display a significant increase in their ability to tolerate Cu indicating that the
369 selective pressure posed by this contamination is lower (compared to RC). We found no
370 significant differences in the expression of the aforementioned copper homeostasis genes
371 between individuals collected from MB and the reference site. It is, however, noted that the
372 abundance of some gene transcripts was highly variable on the different days with, for example,
373 the putative ATOX1 gene transcript fluctuating between approximately 4-fold up- and down-
374 regulated compared to the reference population. In contrast to our data for RC, we found that the
375 animals collected from MB displayed significant up-regulation of Glutathione S-transferase
376 (GST) enzymes, by up to approximately 128-fold, however other antioxidants including

377 Superoxide dismutase and Catalase were unchanged compared to the reference. The GST
378 enzymes have important roles in cellular detoxification and anti-lipid peroxidation processes and
379 are frequented as biomarkers of environmental pollution. However, the response of GST
380 expression to metals remains unclear; some studies reporting that they are induced by metal-
381 contamination ⁴³ and others that they are repressed ⁴⁴.

382 **Genetic Response to Cu depletion**

383 We measured the stability over time of the observed differences in transcriptional
384 regulation between the tolerant (RC) and non-tolerant (CC / reference) animals when in the
385 absence of Cu. To achieve this, a second series of real-time PCR experiments were carried out
386 wherein animals from each population were placed into tanks containing clean sediment and
387 ASW for up to 3 weeks prior to RNA extraction. The relative abundance of mRNA transcripts
388 matching the copper homeostasis genes (i.e. those showing significant regulation between the
389 populations *in situ*) was measured every 7 days by sacrificing a random sample of the individuals
390 within each tank. The observed *in situ* differences in transcript abundance were reduced after 7
391 days in the clean sediment. There was no significant difference in the abundance of the
392 transcripts after 14 days, with the exception that the transcript encoding CTR1 remained
393 significantly more abundant in the Cu-tolerant animals from RC over the duration of the
394 experiment. The CTR1 gene encodes a Cu uptake membrane transporter and may supply Cu to
395 sustain normal cellular processes in addition to detoxificatory pathways.



396

397 **Figure 4.** A comparison of the abundance of select transcripts in representatives of the Cu
 398 contaminated population from RC in comparison to the uncontaminated reference population.
 399 The analysis was performed on a pool of at least 20 individuals from 3 different holding tanks.
 400 The horizontal line shows the mean expression ratio versus the reference, the error bars are the
 401 SEM from 3 replicate measurements (tanks) and the symbols show the values obtained for each
 402 replicate. Statistically significant differences (one way ANOVA, $p < 0.05$) are marked with a
 403 star.

404 Interestingly, the copper toxicity assay, as described in Figure 3, was repeated using
405 animals subjected to the 21 day experiment (see supplementary information Figure S2) wherein
406 the animals from RC remained significantly more tolerant to CuCl₂. This supports the hypothesis
407 that the mechanism for Cu tolerance in the RC population can persist in the absence of Cu, and is
408 encoded by an underlying genetic adaptation.

409 The use of next generation sequencing in environmental science has expanded rapidly,
410 facilitating functional genomics-based investigations using non-model (i.e. non-sequenced)
411 organisms. This study was intended as a primer to facilitate molecular understanding of the
412 processes that enable a species to rapidly adapt to pressure from anthropogenic contaminants
413 such as waste metals. In summary, we compared 3 populations of *N. diversicolor*, either living
414 with or without prolonged Cu stress, and either with or without ecological tolerance. The results
415 indicate that whilst the tolerant individuals display up-regulated expression of Cu homeostasis
416 genes, they do not make a measurable response to the product of Cu toxicity; namely oxygen
417 radicals and cellular (DNA) damage. This supports the hypothesis that the Cu tolerance trait is
418 facilitated by the activity of a detoxificatory pathway as opposed to some enhanced cellular
419 repair. There is mounting evidence for the role of the detoxificatory granules in metal tolerance
420 in invertebrates, and further work should be carried out to identify the genes and proteins that
421 function in their formation. The non-tolerant but heavily exposed individuals in MB did not
422 display a measurable change in the expression of Cu homeostasis genes compared to individuals
423 from the reference population, but did produce significantly greater quantities of gene transcripts
424 corresponding to antioxidants; specifically Glutathione S-transferase enzymes. Whilst this
425 response cannot be attributed to the metal contamination alone, the absence of the response to

426 Cu, as observed in the tolerant population, is good evidence that the Cu homeostasis systems in
427 this species are central to the molecular basis for the tolerance.

428

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432

433 **Supporting Information:** The supporting information submitted with this manuscript provides
434 details of the primer sequences used for real-time PCR experiments including those specific for
435 internal references (Table S1), transcriptome assembly and validation statistics (Table S2),
436 Energy Dispersive X-ray Spectroscopy analysis data for the identification of Cu- and S-
437 containing granules in *N. diversicolor* thin sections (Figure S1), and the results of a Cu toxicity
438 assay for animals that had been placed in clean sediment for 21 days (Figure S2). This material is
439 available free of charge via the Internet at <http://pubs.acs.org>.

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