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# **Accepted Manuscript**

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1	Neutral red retention time assay in determination of toxicity of nanoparticles
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20	Keywords: Mytilus, metal oxide, lysosome, membrane stability, neutral red, NRRT
21	
	1

#### Abstract 23

24	The neutral red retention time (NRRT) assay is useful for detecting decreased
25	lysosomal membrane stability in haemocytes sampled from bivalves, a phenomenon
26	often associated with exposure to environmental pollutants including nanomaterials.
27	Bivalves are popular sentinel species in ecotoxicology and use of NRRT in study of
28	species in the genus Mytilus is widespread in environmental monitoring. The NRRT
29	assay has been used as an in vivo test for toxicity of carbon nanoparticles (Moore MN,
30	Readman JAJ, Readman JW, Lowe DM, Frickers PE, Beesley A. 2009. Lysosomal
31	cytotoxicity of carbon nanoparticles in cells of the molluscan immune system: An in
32	vivo study. Nanotoxicology. 3 (1), 40-45). We here report application of this assay
33	adapted to a microtitre plate format to a panel of metal and metal oxide nanoparticles
34	(2ppm). This showed that copper, chromium and cobalt nanoparticles are toxic by this
35	criterion while gold and titanium nanoparticles are not. As the former three
36	nanoparticles are often reported to be cytotoxic while the latter two are thought to be
37	non-cytotoxic, these data support use of NRRT as a general in vitro assay in
38	nanotoxicology.
39	

#### 40 **1. Introduction**

41 The unusual properties of nanomaterials provide them with several possible routes to toxicity in biological systems. Their small size sometimes enables them to cross important 42 43 biobarriers e.g. skin, blood-brain, intestine, maternal-foetus (Tedesco and Sheehan, 2010; Elsaesser and Howard, 2012; Jiang et al., 2014). Their very large surface area to volume ratio 44 enables a greater proportion of atoms to be displayed on the particle surface compared to 45 corresponding macromaterials (Nel et al., 2009; Nel et al., 2013). Moreover, specific 46 functional groups on nanoparticle surfaces may facilitate biospecific interactions allowing a 47 range of possible biological effects (Hoet et al., 2004; Moore, 2006; Klaper et al., 2014). 48 Nanomaterials can also translocate within the human body into other systems such as 49 50 circulatory and lymphatic vessels (Gwinn and Vallyathan, 2006; Buzea et al., 2007; Elsaesser and Howard, 2012). Thus, nanoparticles have significant potential to cause adverse health 51 52 effects in humans and other organisms upon prolonged exposure.

53 Because of increasing commercial production and use of nanomaterials, issues of their accumulation and fate in the environment and their possible effects on 54 55 ecosystems arise (Moore, 2006; Tedesco and Sheehan, 2010; Ivask et al., 2014). The majority of human habitation worldwide is within 100km of coastlines and the aquatic 56 environment collects domestic, agricultural, shipping and industrial runoffs from 57 these coastal zones. This makes aquatic ecosystems particularly at risk to potential 58 toxicity of nanomaterials of anthropogenic origin. Invertebrates are key elements of 59 60 the aquatic food chain and mussels are amongst the most abundant of these (Baun et 3

al., 2008). As filter-feeders, mussels are exquisitely selective in the particle sizerange which they ingest (Defossez and Hawkins, 1997; Ward and Kach, 2009) and
can bioconcentrate metals and organic pollutants within their tissues. This has led to
their widespread study in ecotoxicology (Moore, 1985; Widdows and Donkin, 1992)
and filter-feeders have been suggested as especially attractive targets for probing the
environmental fate of nanomaterials (Moore, 2006; Ward and Kach, 2009; Canesi et
al., 2012).

Lysosomes are important subcellular organelles that contain many hydrolytic 68 enzymes, carry out protein degradation and detoxify some foreign compounds. At the 69 cellular level, lysosomal digestion pathways include phagocytosis, endocytosis and 70 autophagy. The lysosomal membrane protects the cytosol, and therefore the rest of the 71 cell, from leakage of degradative enzymes. However, malfunctioning of lysosomes 72 73 and their accumulation of toxic pollutants have been linked to lysosomal storage diseases and result in lysosomal injury and oxidative damage, in some cases leading 74 to cell death (Moore et al., 2007). The neutral red retention time (NRRT) assay takes 75 advantage of this phenomenon by measuring decreased time of retention of a dye, 76 neutral red (ACS no. 553-24-2), within phagocytic haemocytes of a range of aquatic 77 organisms including mussels, crustaceans and fish (Regoli, 1992; Tedesco et al, 2008; 78 79 Lowe et al 1995; Svendsen et al, 2004). In the popular sentinel species, *Mytilus edulis*, hemocytes are essential immune system components (Rickwood and Galloway, 80 81 2004). NRTT has been reported as a useful indicator of the organism's overall health

82 status because animals exposed to pollutants often have compromised lysosomal stability (Moore et al., 2009; Borenfreund and Puerner 1985; Piola et al., 2013). A 83 spectrophotometric version of the assay was developed by Babich and Borenfreund 84 (1990) and a microscopic slide observation method was developed by Moore et al., 85 (2009). This assay takes advantage of the tendency of haemocytes to take up 86 nanoparticles most probably by either phagocytosis or macro-endocytosis and 87 involves exposing haemocytes to nanoparticles on a microscope slide (Moore et al., 88 2009). In this short report, we have adapted this methodology to a microtitre plate 89 format enabling high-throughput screening of large numbers of replicates, doses and 90 91 nanoparticles simultaneously (Fig. 1). As proof of principle, we have assessed a panel of metal and metal oxide nanoparticles with this assay. 92

#### 94 **2. Materials and Methods**

#### 95 2.1. Mytilus edulis sampling

96	<i>M. edulis</i> individuals (4-6cm shell-length) were collected from an intertidal site in
97	Cork Harbour, Ireland (location: 51.49°N, 8 18°W; Lyons et al., 2003). All Animals
98	were acclimated in tanks for a week with a 12 h light/dark cycle at a temperature of
99	15°C and 34–36‰ salinity, fed and with regular changing of water.

- 100
- 101 *2.2.Nanoparticle suspension preparation*

Metal or metal oxide nanoparticles (copper oxide, titanium dioxide, gold, 102 chromium oxide and cobalt oxide) of nominal sizes <50nm were purchased from 103 Sigma-Aldrich (Dorset, UK). Nanopowders (10mg) were suspended in 10 ml of 20 104 mM citric acid adjusted to pH 7, and sonicated for 1h using a tip sonicator. A stepped 105 microtip was used and the total power transferred to the suspension was 2.4W 106 (determined by the calorimetric method). Ultrasound was applied as 15s pulses with 107 15s breaks between them (Taurozzi et al., 2010). The suspensions were left at  $60^{\circ}$ C 108 overnight and were then filtered using a 220nm pore size cellulose acetate filter 109 (Millipore, Watford UK). 110

Haemolymph samples were freshly extracted for NRRT assay as described by 113 Moore et al. (2009). In the present work, haemolymph from each of five animals was 114 115 extracted from adductor muscle using a 20 gauge hypodermic needle fitted on a 1 ml syringe containing 100ul tris buffered saline buffer, which was pooled to provide a 116 total volume of 2 ml haemolymph solution. Three biologically independent replicates 117 were used (i.e. haemolymph was taken from 3x5 individual animals). Samples were 118 constantly vortexed to resuspend the haemolymph and prevent aggregation. 119 Haemolymph was then evenly aliquoted (500 µL) followed by exposure to 120 nanoparticles at a final concentration of 2 ppm for 1 h at ambient temperature (20°C). 121 Tubes were gently shaken every 5 min to optimise exposure. The above procedure 122 was applied to a panel of metal or metal oxide nanoparticles and a control sample was 123 treated identically but without the presence of nanoparticle. 124

125

#### 126 2.4.Neutral red retention time (NRRT) assay

Following nanoparticle exposure, 100  $\mu$ l haemolymph from all six treatment groups was loaded into individual wells of a 96-well microtitre plate (Sarstedt, Wexford Ireland). This was performed with three independent biological replicates. Fifty  $\mu$ l stock neutral red dye solution (200  $\mu$ M) was then added. Four plates were used in parallel for time-points 15, 30, 60 and 90 min. All plates were placed in the dark allowing 15, 30, 60 or 90 min, respectively, for dye uptake. Dye and medium were quickly removed from the plates after incubation and washed with 150  $\mu$ L



#### 141 **3. Results and Discussion**

#### 142 *3.1. Neutral red retention time assay of metal oxide nanoparticles*

Haemolymph from *M. edulis* was exposed to a panel of metal or metal oxide 143 144 nanoparticles at a final concentration of 2ppm (Fig. 1). Lysosomal membrane stability was tested by measuring NRRT at four different time points; 15, 30, 60 and 90 min. 145 Results were analysed and statistically compared to the control group using a one-way 146 anova test with confidence limit of 95% (Figure 2). Lysosomal membrane stability 147 showed a significant decrease (p<0.05) upon exposure to copper, cobalt and 148 chromium nanoparticles at all time-points tested, indicating toxic effects on 149 lysosomes of these nanomaterials. However, no significant effects were observed on 150 151 exposure of titanium or gold nanoparticles, suggesting they are less toxic by the criterion of this in vitro assay. 152

153

#### 154 *3.2.Toxicity of metal or metal oxide nanoparticles*

The particles selected for this study have previously been reported to display a range of toxicity in biological systems. Titanium dioxide nanoparticles (which are widely used commercially as a component of sunscreens) are generally regarded as less toxic to aquatic species (Federici et al, 2007). However, it should be noted that, in mice, NO and tumour necrosis factor alpha production were elicited after exposure to titanium dioxide nanoparticles (<10nm). This finding suggested that both damage to

the cell structure and macrophage dysfunction may occur, leading to reduction in both 161 non-specific and specific immune responses in some individual animals (Liu et al 162 2010). Copper oxide and chromium oxide nanoparticles are notorious for their toxic 163 effects, and have been implicated in toxicity to non-target organisms (Ivask et al, 164 2014), reduction of immune status (Zha et al 2009), damage to animal tissues (Chen et 165 al, 2006; Griffitt et al, 2007), and induction of reactive oxygen species (Fahmy and 166 Cormier, 2009; Horie et al 2011). Cobalt oxide nanoparticles readily enter cultured 167 human cells where they are found to have a negative effect on cell viability (Papis et 168 al., 2009). They have been reported to induce primary DNA damage in a 169 concentration-dependent manner. Various redox enzyme activities were decreased 170 after treatment with cobalt nanoparticles, suggesting potential toxic risk and inhibition 171 172 of antioxidant capacity (Jiang et al, 2012).

173

#### 174 *3.3.Potential for high-throughput assay*

The assay format reported here includes minimisation of biological variation in haemocyte populations by pooling haemolymph across five individual animals. Moreover, three independent replicates gave essentially identical results and allowed reproducible discrimination across the nanoparticle panel studied. Use of 96-well microtitre plates makes possible high-throughput analysis of large numbers of samples, replicates and concentrations within the time-scale suggested by Moore et al.



186

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- 287

## 289 Figure legends

- 290 **Figure 1** Schematic overview of NRTT assay.
- 291 Figure 2 Neutral red retention time (NRRT) assay in response to a panel of
- 292 nanoparticles. Neutral red dye extracted from exposed haemocytes was measured
- spectrophotometrically at 570nm in a plate reader (\*p < 0.05 versus control values).





Figure 1

- Neutral red retention time assay used haemolymph of five pooled mussels.
- Assay was miniaturised for reading in a plate reader, facilitating many samples and replicates.
- Copper, chromium and cobalt nanoparticles were toxic while gold and titanium were not.