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Oxidative responsiveness to multiple stressors in the key Antarctic species, *Adamussium colbecki*:  
Interactions between temperature, acidification and cadmium exposure

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*Original*

Oxidative responsiveness to multiple stressors in the key Antarctic species, *Adamussium colbecki*:  
Interactions between temperature, acidification and cadmium exposure / Benedetti, Maura; Lanzoni,  
Ilaria; Nardi, Alessandro; D'Errico, Giuseppe; DI CARLO, Marta; Fattorini, Daniele; Nigro, Marco; Regoli,  
Francesco. - In: MARINE ENVIRONMENTAL RESEARCH. - ISSN 0141-1136. - STAMPA. - 121:(2016), pp. 20-  
30. [10.1016/j.marenvres.2016.03.011]

*Availability:*

This version is available at: 11566/239512 since: 2022-06-01T13:02:34Z

*Publisher:*

*Published*

DOI:10.1016/j.marenvres.2016.03.011

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note finali coverage

(Article begins on next page)

**Oxidative responsiveness to multiple stressors in the key Antarctic species, *Adamussium colbecki*:  
interactions between temperature, acidification and cadmium exposure**

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31 **Abstract:**

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High-latitude marine ecosystems are ranked to be among the most sensitive regions to climate change since highly stenothermal and specially adapted organisms might be seriously affected by global warming and ocean acidification. The present investigation was aimed to provide new insights on the sensitivity to such environmental stressors in the key Antarctic species, *Adamussium colbecki*, focussing also on their synergistic effects with cadmium exposure, naturally abundant in this area for upwelling phenomena. Scallops were exposed for 2 weeks to various combinations of Cd (0 and 40 µgL<sup>-1</sup>), pH (8.05 and 7.60) and temperature (-1 and +1°C). Beside Cd bioaccumulation, a wide panel of early warning biomarkers were analysed in digestive glands and gills including levels of metallothioneins, individual antioxidants and total oxyradical scavenging capacity, onset of oxidative cell damage like lipid peroxidation, lysosomal stability, DNA integrity and peroxisomal proliferation. Results indicated reciprocal interactions between multiple stressors and their elaboration by a quantitative hazard model based on the relevance and magnitude of effects, highlighted a different sensitivity of analysed tissues. Due to cellular adaptations to high basal Cd content, digestive gland appeared more tolerant toward other prooxidant stressors, but sensitive to variations of the metal. On the other hand, gills were more affected by various combinations of stressors occurring at higher temperature.

**Keywords:** Antarctic scallop, Multiple stressors, Climate Change, Cadmium, Oxidative stress, Antioxidant defences

50 **Introduction:**

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The concentration of CO<sub>2</sub> in the atmosphere has continuously increased from 278 ppm in pre-industrial time up to 400 ppm nowadays, representing one of the most important drivers of global climate change (IPCC, 2013). Approximately 30% of total CO<sub>2</sub> emissions has been absorbed by seawater, causing the well known process of ocean acidification, OA (Raven et al., 2005). Progressive changes in carbonate chemistry determined a decrease of 0.1 pH units compared to the pre-industrial levels, with a further expected reduction of 0.3 to 0.5 units by the end of the 21st century (Raven et al., 2005).

The Southern Ocean accounts for about 4% of the global uptake of CO<sub>2</sub> by the world oceans due to the high solubility of CO<sub>2</sub> at low temperature and mixing patterns from upwellings and deep water formation (Fabry et al., 2009). Similar characteristics lead to an increased rate of acidification and a more rapid shoaling of the saturation horizons (Fabry et al., 2009). In this respect, polar organisms, which have evolved in environmentally stable conditions, might be more vulnerable to climate change, in particular regarding calcification processes and variations of fundamental pathways like energy metabolism, growth, reproduction, larval development and oxidative stress. Compared to temperate models, however, only a few studies have investigated the possible effects of climate change on Antarctic marine species (Cubillos et al., 2007; McClintock et al., 2009; Moy et al., 2009; Seibel et al., 2012; Walker et al., 2013; Constable et al., 2014; Collard et al., 2015; Flynn et al., 2015).

In addition, while future scenarios of temperature and ocean acidification can be simulated from CO<sub>2</sub> emission models (IPCC, 2013), at this moment it is virtually impossible to predict the biological impact and synergistic effects of multiple stressors, which can be indirectly modulated by variations of environmental factors due to climate change. In this respect, it has been widely postulated that increased temperature and ocean acidification could influence chemical speciation and bioavailability of environmental pollutants, bioaccumulation processes, responsiveness of detoxification mechanisms, sensitivity and consequences on organisms health condition, but clear evidences of similar modulations are still lacking.

In the Ross Sea, local upwelling phenomena are responsible for a natural enrichment and elevated bioavailability of cadmium (Cd) which is accumulated in tissues of Antarctic invertebrates and fish at values typically 10-50 folds higher than those of similar temperate species (Nigro et al., 1997). Although this element does not apparently cause adverse effects on the organisms, in Antarctic fish it was shown to influence the responsiveness of fundamental metabolic pathways to other stressors, i.e. the cytochrome P450 biotransformation mechanisms, male vitellogenin expression and the antioxidant network (Regoli et al., 2005; Canapa et al., 2007; Benedetti et al., 2007; 2009). In the scallop *Adamussium colbecki* (Smith 1902), another key sentinel species of the Antarctic environment, the enhanced accumulation of Cd during algal bloom is paralleled by a general increase of antioxidant defences (Regoli et al., 2000; 2002). Beside the role in counteracting a natural increase of prooxidant pressure during phytoplankton blooms, oxyradical

85 metabolism and antioxidant defences have a fundamental role for polar organisms in adaptation  
86 mechanisms to high solubility of oxygen in cold seawater, elevated content of oxidizable poly-unsaturated  
87 fatty acids (PUFAs) in membranes, high cellular mitochondrial densities, and the need of long-term  
88 protection of proteins and RNAs due to their low turnover rate (Abele and Puntarulo, 2004).

89 Based on previous issues, the main aim of this study was to investigate whether variations of  
90 temperature and pH may singularly or synergistically affect the sensitivity of *A. colbecki* to Cd, highlighting a  
91 potentially reciprocal modulation of key cellular responses by multiple stressors. Scallops were exposed to  
92 various combinations of treatments including two different levels of temperature, pH and Cd  
93 concentrations, opportunely chosen as reflecting environmentally realistic or future scenarios for Antarctic  
94 marine environment.

95 Analyses of Cd bioaccumulation were integrated with a wide panel of early warning biomarkers and  
96 results were elaborated within a quantitative model (SediquaSoft) which, based on biological relevance  
97 and magnitude of observed variations, summarize a hazard index for biomarkers results (Piva et al., 2011;  
98 Benedetti et al., 2012).

99 Overall, this study was expected to provide new insights on mechanisms underlying the  
100 responsiveness of a model Antarctic species to variations of temperature and acidification, interactions  
101 occurring between multiple stressors, and potential consequences of climate change in areas characterized  
102 by elevated environmental pollution or geochemical anomalies.

## 103 **Materials and methods:**

### 104 *Experimental design*

105 Scallops, *A. colbeckii*, were sampled during the XXIX Italian Antarctic Expedition (2013-2014) from  
106 Terra Nova Bay (Ross Sea) and acclimatized to laboratory conditions for 10 days with running, unfiltered  
107 seawater at the controlled temperature of -1°C and pH 8.05. A total of 240 organisms were randomly  
108 distributed in eight tanks (150 L each) and exposed to one of the following experimental conditions: 1)  
109 control (CTRL), at environmental temperature (-1°C) and environmental pH (8.05); 2) Cd exposure (Cd), at  
110 40 µg/L of Cd, -1°C, pH 8.05; 3) acidified water condition (Ac), at pH 7.60 and -1°C; 4) warm exposure  
111 (Warm), at +1°C and pH 8.05; 5) acidified and Cd exposure (Ac + Cd), at pH 7.60, 40 µg/L of Cd and -1°C; 6)  
112 warm and Cd exposure (W + Cd), at +1°C, 40 µg/L of Cd and pH 8.05; 7) warm and acidified condition (W +  
113 Ac), at +1°C and pH 7.6; 8) warm, acidified and Cd exposure (W + Ac + Cd), at +1°C, pH 7.6 and 40 µg/L of  
114 Cd. After 14 days, organisms were sacrificed, haemolymph, gills and digestive glands were rapidly  
115 dissected, frozen in liquid nitrogen and stored at -80°C until analyses. A portion of haemolymph and gills  
116 was maintained in Carnoy's solution (3:1 methanol:acetic acid) for micronuclei frequency analyses. No  
117 mortality was observed during the experiments. For both chemical and biochemical analyses, 5 pools, each  
118 constituted by tissue of 6 specimens, were prepared for digestive glands, gills and haemolymph.

120 *Chemical analyses*

121 Cd concentration in scallops tissues was analysed according to previously described methods  
122 (Regoli et al., 2005). For every treatment, digestive glands and gills were dried at 70°C until constant weight  
123 and digested under pressure with nitric acid in microwave digester systems (CEM, Mars Systems). Quality  
124 assurance and quality control was assessed by processing blank samples and reference standard material  
125 (Mussel Tissue Standard Reference Material SRM 2977, National Institute of Standards and Technology). Cd  
126 was analysed by atomic absorption spectrophotometry with electrothermal atomization. The  
127 concentrations obtained for the standard reference material were always within the 95% confidence  
128 interval of certified values. Data are expressed as  $\mu\text{g/g}$  dry weight (mean values  $\pm$  standard deviations,  $n=5$ ).

129 *Biomarker analyses*

130 Sample preparation and analytical protocols have been fully detailed elsewhere (Regoli et al.,  
131 2000). Metallothioneins were analyzed in digestive glands and gills homogenized in 20 mM Tris-HCl buffer  
132 (pH 8.6), 0.5 M sucrose, 0.006 mM phenylmethylsulfonyl fluoride (PMSF), and 0.01%  $\beta$ -mercaptoethanol  
133 and centrifuged at 30,000g for 45 min. After acidic ethanol/chloroform fractionation of tissue supernatants,  
134 metallothioneins were quantified by a spectrophotometric assay using reduced glutathione (GSH) as  
135 standard.

136 For measurement of enzymatic antioxidants, tissues (digestive gland and gills) were homogenized  
137 (1:5 and 1:3 w:v ratio respectively) in 100 mM K-phosphate buffer (pH 7.5), 0.1 mM phenylmethylsulphonyl  
138 fluoride (PMSF), 0.1 mg mL<sup>-1</sup> bacitracin, 0.008 TIU mL<sup>-1</sup> aprotinin, 1 mg mL<sup>-1</sup> leupeptin, 0.5 mg/mL  
139 pepstatin, NaCl 2.5%, and centrifuged at 110,000g for 1 h at 4 °C. Measurements were made with a Varian  
140 (model Cary 3) spectrophotometer at a constant temperature of 18 °C. Catalase (CAT) was measured by the  
141 decrease in absorbance at 240 nm (extinction coefficient,  $\epsilon = 0.04 \text{ mM}^{-1} \text{ cm}^{-1}$ ) due to the consumption of  
142 hydrogen peroxide, H<sub>2</sub>O<sub>2</sub> (12 mM H<sub>2</sub>O<sub>2</sub> in 100 mM K-phosphate buffer pH 7.0). Glutathione reductase (GR)  
143 was determined from NADPH oxidation during the reduction of oxidized glutathione, GSSG ( $\lambda = 340 \text{ nm}$ ,  $\epsilon =$   
144  $6.22 \text{ mM}^{-1} \text{ cm}^{-1}$ ). The final assay condition were 100 mM K-phosphate buffer pH 7.0, 1 mM GSSG, and 60  
145 mM NADPH. Glutathione peroxidases (GPx) activities were assayed in a coupled enzyme system where  
146 NADPH is consumed by glutathione reductase to convert the formed GSSG to its reduced form (GSH). The  
147 decrease of absorbance was monitored at 340 nm ( $\epsilon = 6.22 \text{ mM}^{-1} \text{ cm}^{-1}$ ) in 100 mM K-phosphate buffer pH  
148 7.5, 1 mM EDTA, 1 mM dithiothreitol, 1 mM sodium azide (NaN<sub>3</sub>) (for hydrogen peroxide assay), 2 mM GSH,  
149 1 unit glutathione reductase, 0.24 mM NADPH, and 0.5 mM hydrogen peroxide or 0.8 mM cumene  
150 hydroperoxide as substrates, respectively, for the selenium-dependent and for the sum of Se-dependent  
151 and Se-independent forms. The rate of the blank reaction was subtracted from the total rate. Glutathione  
152 S-transferases (GST) were determined at 340 nm using 1-chloro-2,4-dinitrobenzene as substrate (CDNB).  
153 The assay was carried out in 100 mM K-phosphate buffer pH 6.5, 1.5 mM CDNB, 1 mM GSH ( $\epsilon = 9.6 \text{ mM}^{-1}$   
154  $\text{cm}^{-1}$ ).

155 Total glutathione was analyzed in samples of digestive gland and gill homogenized (1:5 and 1:3 w:v ratio  
156 respectively) in 5% sulfosalicylic acid with 4 mM EDTA, maintained for 45 min on ice and centrifuged at  
157 37,000g for 15 min. The resulting supernatants were enzymatically assayed (Benedetti et al., 2007).  
158 The total oxyradical scavenging capacity (TOSC) assay measures the overall capability of cellular  
159 antioxidants to neutralize different forms of artificially generated oxyradicals, thus inhibiting the oxidation  
160 of 0.2 mM  $\alpha$ -keto- $\gamma$ -methiolbutyric acid (KMBA) to ethylene gas (Regoli et al., 2000). Peroxyl radicals (ROO $\cdot$ )  
161 were generated by the thermal homolysis of 20 mM 2–2'-azo-bis-(2-methylpropionamide)-  
162 dihydrochloride (ABAP) in 100 mM K-phosphate buffer, pH 7.4. Hydroxyl radicals ( $\cdot$ OH) were produced by  
163 the Fenton reaction of iron-EDTA (1.8  $\mu$ M Fe<sup>3+</sup>, 3.6  $\mu$ M EDTA) plus ascorbate (180  $\mu$ M) in 100 mM K-  
164 phosphate buffer. Ethylene formation in control and sample reactions was analyzed at 10–12 min time  
165 intervals by gas-chromatographic analyses and the TOSC values quantified from the equation: TOSC=100-  
166 (JSA/JCA $\times$ 100), where JSA and JCA are the integrated areas calculated under the kinetic curves for samples  
167 (SA) and control (CA) reactions. For all the samples, a specific TOSC (normalized to content of protein) was  
168 calculated by dividing the experimental TOSC values by the relative protein concentration contained in the  
169 assay.

170 The content of malondialdehyde (MDA) was measured in homogenates of scallops digestive glands  
171 and gills derivatized with 1-metyl-2-phenylindole and spectrophotometrically determined after calibration  
172 against a malondialdehyde standard curve.

173 Acetylcholinesterase activity (AChE) was analyzed in hemolymph and gills: hemolymph was  
174 centrifuged at 3,000g for 5 min, while gills were homogenized in 100 mM Tris-HCl buffer (pH 7.2), 0.55 M  
175 sucrose and centrifuged at 10,000g for 10 min. Obtained supernatants were spectrophotometrically  
176 assayed by the Ellman's reaction at  $18 \pm 1$  °C,  $\lambda = 412$  nm,  $\epsilon = 13.6$  mM<sup>-1</sup> cm<sup>-1</sup>.

177 For the activity of Acyl CoA oxidase (AOX), samples were homogenized in 1 mM sodium bicarbonate  
178 buffer (pH 7.6) containing 1 mM EDTA, 0.1% ethanol, 0.01% Triton X-100 and centrifuged at 500g for 15  
179 min at 4 °C. The H<sub>2</sub>O<sub>2</sub> production was measured in a coupled assay by following the oxidation of  
180 dichlorofluorescein-diacetate (DCF-DA) catalyzed by an exogenous horseradish peroxidase (HRP). The  
181 reaction medium was 0.5 M potassium phosphate buffer (pH 7.4), 2.2 mM DCF-DA, 40  $\mu$ M sodium azide,  
182 0.01% Triton X-100, 1.2 U mL<sup>-1</sup> HRP in a final volume of 1 mL. After a preincubation at 25 °C for 5 min in  
183 the dark with an appropriate volume of sample, reactions were started adding the substrates palmitoyl-CoA  
184 at final concentrations of 30  $\mu$ M and 100  $\mu$ M for Acyl-CoA oxidase (AOX) and readings were carried out  
185 against a blank without the substrates at 502 nm.

186 Protein concentrations were measured according to Lowry method (1951), using bovine serum albumin  
187 (BSA) as standard. All biochemical biomarkers are expressed as mean values  $\pm$  standard deviations (n=5).

188 Lipofuscin content of tertiary lysosomes was determined on duplicate cryostat sections (8  $\mu$ m thick)  
189 of digestive gland. Slides were fixed in Beker's fixative (+2.5% NaCl) and stained by Schmrol reaction before

190 mounting in glycerol gelatine. Five measurements were made on digestive tubules of each section (two  
191 sections for mussel, 10 scallops for each experimental condition). Quantification of staining intensity was  
192 performed with Image-Pro® Plus 6.2 Analysis Software and then normalized to the area of digestive  
193 tubules.

194 The DNA integrity was evaluated at chromosomal level by measuring the micronucleus (MN)  
195 frequency in scallops haemocytes rapidly fixed in Carnoy's solution (3:1 methanol: acetic acid), dispersed  
196 on glass slides and stained with the fluorescent dye 4',6-diamidino- 2-phenylindole at 100 ng ml<sup>-1</sup>. For each  
197 experimental condition, 10 scallops were observed and for each specimen 2000 cells with preserved  
198 cytoplasm were scored to assess the presence of micronuclei, defined as round structures, smaller than 1/3  
199 of the main nucleus diameter, on the same optical plan and clearly (Benedetti et al., 2014).

200

## 201 Statistical analyses

202 Analysis of variance (1-way ANOVA) was applied to test significance of effects on biochemical  
203 parameters caused by temperature, acidification, Cd exposure and their reciprocal interactions. Level of  
204 significance was set at  $p < 0.05$ , homogeneity of variance was checked by Cochran C and mathematical  
205 transformation applied if necessary; post-hoc comparison (Newman-Keuls) was used to discriminate  
206 between means of values. Multivariate statistical analysis (non-metric multidimensional scaling analysis)  
207 was applied to biomarkers data to discriminate various experimental treatments.

208 Results on biological parameters in scallops were further elaborated within a previously developed  
209 quantitative and software-assisted model (SediquaSoft, Piva et al., 2011). Whole calculations, detailed  
210 flow-charts, rationale for weights and thresholds have been fully given elsewhere and successfully validated  
211 in field conditions, during the characterization and classification of risk from industrial and harbour  
212 sediments, natural hydrocarbon seepage in coastal areas, the recent Costa Concordia wreck at Giglio Island  
213 and the ecotoxicological effects of microplastics (Piva et al., 2011, Benedetti et al., 2012; 2014; Regoli et al.,  
214 2014; Avio et al., 2015; Bebianno et al., 2015). Briefly, depending on species and tissue, the model assigns  
215 to each biomarker a "weight" based on the relevance of biological endpoint, and a "threshold" for changes  
216 of biological significance which consider both inductions and/or inhibitions of various responses. For every  
217 analysed biomarker, the measured variation is compared to the threshold, then corrected for the weight of  
218 the response and the statistical significance of the difference compared to controls. Depending on the  
219 magnitude of the calculated effect, each biomarker response is assigned by the model to 1 of 5 classes of  
220 effect (from Absent to Severe); the calculation of the Hazard Quotient for biomarkers ( $HQ_{BM}$ ) does not  
221 consider the contribution of responses with an effect lower or equal to threshold (Absent or Slight),  
222 calculates the average for those with an effect up to two-fold compared to the threshold (Moderate), and  
223 adds the summation ( $\Sigma$ ) for the responses more than 2 fold greater than the respective threshold, i.e. Major  
224 or Severe (Piva et al., 2011):



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$$HQ_{BM} = \left( \frac{\sum_{j=1}^N Effect_W(j)_{1 < Effect(j) \leq 2}}{num\ biomark_{1 < Effect(j) \leq 2}} + \sum_{k=1}^M Effect_W(k)_{Effect(j) > 2} \right)$$

According to variations measured for various biomarkers, the model summarizes the level of cumulative HQ<sub>BM</sub> in one of five classes of hazard for biomarkers, from Absent to Severe (Piva et al., 2011).

**Results:**

Cd concentrations significantly increased in both digestive glands and in gills of Cd exposed scallops with values 3 and 5 folds higher than those measured in respective control groups (Fig. 1A, 1B). Co-occurring variations of other factors had negligible effects on Cd accumulation with slightly lower values of this element measured only in digestive glands of organisms co-exposed in acidic conditions (Ac+Cd): higher temperature did not cause additional effects on the uptake of this element in both the tissues (Fig. 1A, 1B).

Levels of metallothioneins did not vary in digestive gland (Fig. 1C), while a significant increase was observed in gills after exposure to Cd, without any difference when the element was dosed alone or in combination with higher temperature and/or lower pH (Fig. 1D).

The antioxidant status was assessed by integrating analyses of individual antioxidants with the total capability to neutralize different forms of oxyradicals. In digestive gland, catalase showed a trend toward lowered enzymatic activities with significant effects after co-exposures between Cd and higher temperature, with or without acidified conditions (Fig. 2A). Increased temperature affected also the responses of GST, while no variations were observed for GR and GSH (Fig. 2B-D). Glutathione peroxidases, particularly Se-dependent forms, were slightly inhibited by co-exposures to higher temperature with Cd and/or acidification (Fig. 2E-F). Variations of individual antioxidants were not paralleled by any significant change in the TOSC values toward either ROO· or ·OH (Fig. 2G-H).

Antioxidant responses in gills exhibited lower values of catalase after exposure to Cd with increased temperature and acidification, while various combinations between Cd, acidification and/or temperature were often more effective than individual stressors in affecting responses of GST, GSH, Se-dependent GPx (Fig. 3A-F). Antioxidant variations in gills were reflected by a slight increase of TOSC toward ROO· and HO· in scallops exposed to higher temperature with or without Cd (Fig. 3G-H).

Among cellular biomarkers, malondialdehyde showed a general increase in digestive gland after treatment to multiple stressors, but no variations in gills (Fig. 4A-B). On the other hand, lipofuscin was increased by Cd, low pH and their combination, while it decreased after exposure to higher temperature

257 with or without the other stressors (Fig. 4C). Frequency of micronuclei significantly increased in scallops  
258 exposed to all treatments with both individual and multiple stressors (Fig. 4D). Fluctuating changes of AOX  
259 were observed in digestive gland and gills with a few effects observed after exposure to treatments  
260 involving higher temperature and other stressors (Fig. 4E-F). Acetylcholinesterase was increased in  
261 digestive gland by Cd with or without acidification, while similar effects were induced in gills by higher  
262 temperature with or without Cd and acidification (Fig. 4G-H); the combination of the 3 stressors tended to  
263 reduce this enzymatic activity in both tissues.

264 The nMDS ordination of biomarker responses in a multidimensional space ( $d = 2$ ) analysis is given in  
265 Fig. 5. In digestive tissues results indicated a separation between treatments involving co-exposures at  
266 higher temperature and those with Cd and/or acidification at  $-1^{\circ}\text{C}$  (environmental temperature) (Fig. 5A); in  
267 gills, a more evident difference was observed between organisms exposed to individual stressors compared  
268 to their multiple combinations (Fig. 5B). The elaboration of the overall biomarkers results through weighted  
269 criteria summarized in a single hazard index the biological significance of cellular responses observed in  
270 tissues of *A. colbeckii* exposed to different experimental treatments (Fig. 6). In digestive gland, the  
271 elaborated Hazard Quotient was Moderate for scallops exposed to Cd alone and in combination with low  
272 pH and higher temperature, Slight for all the other treatments. On the other hand, gills appeared more  
273 sensitive to co-exposures between higher temperature and the other stressors (HQ Moderate), compared  
274 to individual stressors or their combination at  $-1^{\circ}\text{C}$  (HQ Slight).

## 275 Discussion:

276 High-latitude marine ecosystems are ranked to be among the most sensitive regions to climate  
277 change since highly stenothermal and specially adapted organisms might be seriously affected by global  
278 warming and ocean acidification. The present investigation was aimed to provide new insights on the  
279 sensitivity to such environmental stressors in the key Antarctic species, *A. colbeckii*, focussing also on their  
280 synergistic effects with Cd exposure. Cd accumulation is of special interest for marine organisms in the area  
281 of Terra Nova Bay due to a naturally elevated bioavailability of this metal enriched in water column by up-  
282 welling currents: during phytoplanktonic bloom, when the algae represent an important trophic source, Cd  
283 is transferred to both benthic species and pelagic food webs. As a consequence of this local peculiarity,  
284 tissue concentrations of Cd in organisms from Terra Nova Bay are up to 20 fold higher than those normally  
285 measured in similar temperate species (Nigro et al., 1997). The elevated Cd content in tissues, despite not  
286 directly toxic, was shown to influence metabolism of organic xenobiotics (Regoli et al., 2005; Benedetti et  
287 al., 2007), and probably interfere with endocrine receptor (ER) and expression of vitellogenin (VTG) in  
288 males of fish *T. bernacchii* (Canapa et al., 2007). In this respect, it was of interest to explore whether a  
289 similar peculiarity of Antarctic organisms might have reciprocal interactions with their sensitivity to the  
290 effects of climate change.

292 This study confirmed elevated basal concentrations of Cd in tissues of *A. colbecki*, but also the  
293 capability of this scallop to further accumulate the metal in response to enhanced bioavailability.  
294 Surprisingly, the Cd accumulation in both digestive gland and in gills was not affected by either  
295 temperature increase or acidification which were expected to influence metabolism and chemical  
296 availability respectively. In this respect, previous studies revealed that temperature increased metal uptake  
297 in some temperate and Arctic organisms (Baines et al., 2006; Sokolova and Lannig, 2008), while other  
298 investigations on temperate mussels (*Mytilus galloprovincialis*) did not support a similar modulation  
299 (Izagirre et al., 2014). Regarding acidification,  $PCO_2$  was shown to enhance Cd and Cu accumulation in  
300 oysters and clams, but the rather subtle and tissue-specific variations suggested that  $CO_2$ -dependent  
301 effects on the metal uptake of marine bivalves are complex and not predictable from the chemical models  
302 of the metal speciation in seawater (Rodriguez-Romero et al., 2014; Ivanina et al., 2015).

303 The increase of Cd concentration in tissues of *A. colbecki* was not reflected by variations of  
304 metallothioneins in digestive gland of exposed scallops, while levels of these proteins increased in gills. The  
305 lack of response in digestive tissues can be related to the elevated basal levels of metallothioneins,  
306 reflecting an adaptation mechanism to high natural concentrations of Cd in these Antarctic organisms.  
307 Despite we did not analyse subcellular distribution of Cd in exposed scallops, previous studies and  
308 metallothioneins characteristics suggest this element as mostly present in a soluble form (Viarengo et al.,  
309 1997; Regoli et al., 1997, 2002): this hypothesis is also supported by the typical amino acid composition of  
310 these proteins in *A. colbecki* that confer a preferential binding capacity for Cd (Ponzano et al., 2001). High  
311 levels of metallothioneins have also been shown to provide protection against oxidative stress through  
312 reaction of sulfhydryl groups with reactive oxygen species, thus further representing an important defence  
313 in polar marine species toward the high environmental prooxidant pressure.

314 Oxyl radical metabolism is of great importance for Antarctic organisms and in our study, a wide  
315 panel of antioxidant defences was integrated with measurement of Total Oxyl Radical Scavenging Capacity  
316 (TOSC) towards hydroxyl and peroxy radicals. Despite the limited effects on Cd bioaccumulation, exposure  
317 to multiple stressors appeared to modulate oxidative responsiveness to this element. In digestive gland, co-  
318 exposures to Cd and higher temperature caused a significant decrease of catalase and glutathione  
319 peroxidases, confirming the sensitivity of these enzymes in revealing a prooxidant pressure, and the  
320 importance of  $H_2O_2$  metabolism as a possible driver (Regoli and Giuliani, 2014). Similar effects, not  
321 compensated by a varied capability to neutralize both  $ROO\cdot$  and  $HO\cdot$ , resulted in a slight increase of  
322 peroxidative processes and consequent enhancement of malondialdehyde content: in this respect, the  
323 lowered values of lipofuscin after treatments involving higher temperature might be the consequence of  
324 enhanced excretion processes of oxidized membranes through tertiary lysosomes. Acidification caused

325 generally slighter oxidative effects in digestive gland, and only when acting as co-stressor with higher  
326 temperature and/or Cd exposure.

327 Sensitivity to oxidative stress was highlighted also in gills where Se-dependent glutathione  
328 peroxidases confirmed decreased values in response to multiple combinations of Cd, temperature and  
329 acidification. Glutathione S-transferases and levels of GSH were upregulated in these tissues, modulating  
330 the responsiveness to Cd in combination with temperature and/or acidification. GSH can act as a direct  
331 scavenger toward H<sub>2</sub>O<sub>2</sub> and, as cofactor of GST, contributes to removal of peroxidative products on  
332 damaged membranes: in this respect, no malondialdehyde accumulation was observed in gills which also  
333 exhibited a certain enhancement of the overall capability to neutralize both peroxy and hydroxyl radicals,  
334 particularly in response to higher temperature and Cd. The different responsiveness of digestive gland and  
335 gills to investigated multiple stressors confirm that variations of antioxidants to environmental stressors  
336 can not be generalized, responding to specific signals, interactions, and pathways that differ in various  
337 tissues or exposure conditions (Regoli et al., 2011). Also for antioxidants, the more limited fluctuations  
338 observed in digestive gland compared to gills are probably related to the higher basal levels of such  
339 defences. The elevated protection in digestive tissues was shown as an important strategy of this scallop to  
340 cope with fluctuations of prooxidant pressure in the Antarctic marine environment characterized by low  
341 temperature and high levels of dissolved oxygen, marked seasonality in photochemical activation of  
342 dissolved organic matter and food availability (Viarengo et al., 1995; Regoli et al., 1997; 2000; 2002).

343 Overall, our results confirm previous evidences that elevated temperature and acidification may  
344 represent prooxidant stressors, as a common consequence of the metabolic and acid–base disturbance in  
345 animals (Matoo et al., 2013). Elevated temperature was shown to cause oxidative stress in marine molluscs  
346 also through a mismatch between generation and detoxification of reactive oxygen species (ROS) ( Abele et  
347 al., 2001). On the other side, acidification was responsible for significant changes in gene expression and  
348 activity of antioxidant defences in the Arctic spider crab *Hyas araneus* and in the Eastern oyster,  
349 *Crassostrea virginica* (Tomanek et al., 2011; Harms et al., 2014). Different mechanisms have been  
350 suggested for prooxidant effects of acidification, including the reaction of CO<sub>2</sub> with peroxynitrite and  
351 formation of reactive carbonate and nitrogen species with elevated oxidizing potential (Dean, 2010).  
352 Elevated CO<sub>2</sub> and/or low environmental pH can also have indirect effects since molluscs have a limited  
353 capability for pH regulation, and sea water acidification would thus determine intracellular acidosis  
354 (Tomanek et al., 2011). Such intracellular conditions may negatively affect the efficiency of mitochondria by  
355 increasing the electron slip in ROS-generating mitochondrial complex I and II, and/or by partially inhibiting  
356 the flow through the downstream electron transport chain complex: in either case, these disturbances of  
357 the electron transport chain would result in elevated rates of ROS generation. Further, intracellular acidosis  
358 can cause the release of chelated transition metals such as Fe<sup>2+</sup> from intracellular store, thus favouring the

359 onset of oxidative stress through Fenton reactions and generation of hydroxyl radicals (Tomanek et al.,  
360 2011).

361 The responsiveness of *A. colbecki* to combinations of various stressors was confirmed by the  
362 enhanced levels of micronuclei observed after all treatments to individual and multiple stressors:  
363 considering the effects observed on antioxidants, the higher frequency of MN might confirm a certain  
364 unbalance of oxyradical metabolism but also suggest a modulation of cellular turnover: in this respect,  
365 beside an oxidative damage on DNA, micronuclei formation would be, at least partially, favoured by a  
366 higher mitotic rate. More limited variations occurred for Acyl CoA oxidase, indicating that higher  
367 temperature, in combination with Cd and/or acidification, may represent the primary stressor affecting  $\beta$ -  
368 oxidation of fatty acids and lipid metabolism. Finally, acetylcholinesterase was particularly modulated by Cd  
369 exposures with or without other stressors, confirming the neurotoxic potential of this element (Del Pino et  
370 al., 2014). Reported effects of Cd include block of cholinergic transmission by decrease of acetylcholine  
371 synthesis and release, inhibition of AChE activity and of postsynaptic transmission, blockage of cholinergic  
372 receptors (Del Pino et al., 2014).

373 When all the complex variations on biomarkers effects were elaborated through multivariate  
374 analysis or the quantitative hazard model (SediquaSoft) interesting trends could be summarized. The  
375 multivariate scaling analysis (nMDS) revealed that temperature influence the pattern of biological  
376 responses in digestive gland where the analysis tended to separate cold from warm exposed organisms. On  
377 the other hand, considering both the biological relevance and the magnitude of observed responses, the  
378 overall quantitative effects were always higher for organisms exposed to Cd, alone and in combination with  
379 other factors: after these treatments, the elaborated HQ was assigned as Moderate, compared to Slight in  
380 organisms exposed to higher temperature, acidification or their combination. This result would further  
381 confirm that the adaptation to the high basal content of Cd, which involves specific cellular strategies like  
382 elevated antioxidant defences, makes these tissues more tolerant toward other potentially prooxidant  
383 stressors like temperature or acidification: at the same time, however, such delicate homeostatic  
384 equilibrium, appears overwhelmed when cellular concentrations of the metal increase, not being further  
385 exacerbated by contemporary variations of temperature and pH. A different trend of responsiveness was  
386 observed in gills which, according to both nMDS and hazard model, generally appeared more sensitive to  
387 various combinations of stressors occurring at higher temperature.

388 In conclusion, this study provided new insights on the reciprocal, synergistic and modulatory effects  
389 of ocean warming, acidification and metals bioaccumulation in a key Antarctic invertebrate. Despite the  
390 elevated basal levels of antioxidants as adaptive mechanism to the high environmental prooxidant  
391 pressure, *A. colbecki* was sensitive to interactions between climate change and Cd bioavailability, with  
392 different sensitivity among analysed tissues toward different factors. Further studies are needed to better

393 understand long term effects, as the onset of earlier biological responses at transcriptional levels or the link  
394 with physiological status of the organisms, as well as the different sensitivity of polar organisms in  
395 comparison with similar temperate species.

396 **Acknowledgments:**

397 This study was financially supported by the Italian National Program on Antarctic Research (PNRA, project  
398 2013/AZ1.14). We thank to the logistic/technical staff and scuba divers of Mario Zucchelli Station of the  
399 XXIX Italian Antarctic Expedition.

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Figure 1. Cadmium concentrations (A-B) and levels of metallothioneins (C-D) in digestive glands and gills of *A. colbecki* exposed to different experimental conditions. Letters indicate significant differences between groups of means. Data are given as mean values  $\pm$  standard deviations,  $n = 5$ .

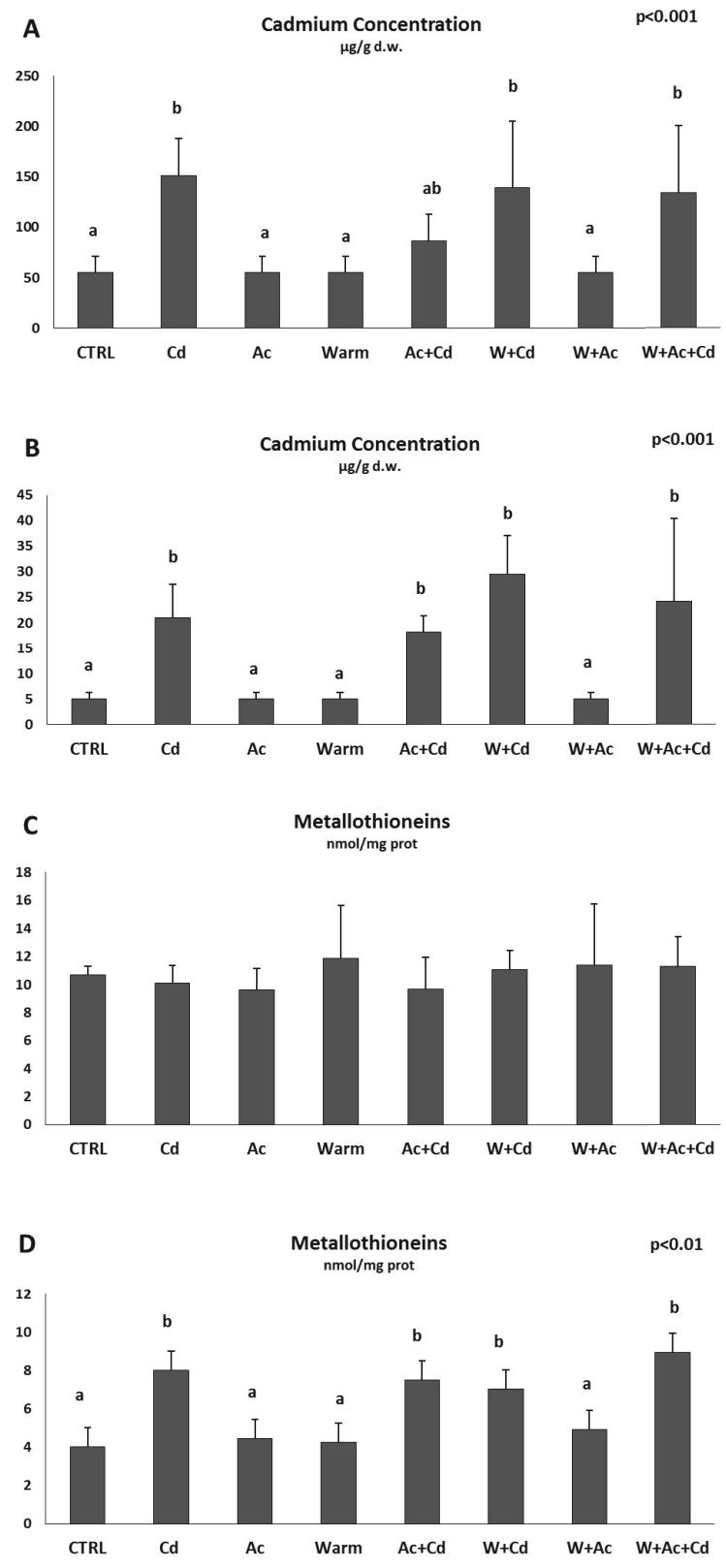
Figure 2. Activities of individual antioxidant enzymes and Total Oxyradical Scavenging Capacity toward peroxy and hydroxyl radicals in digestive gland of *A. colbecki* exposed to different experimental conditions. Letters indicate significant differences between groups of means. Data are given as mean values  $\pm$  standard deviations,  $n = 5$ .

Figure 3. Activities of individual antioxidant enzymes and Total Oxyradical Scavenging Capacity toward peroxy and hydroxyl radicals in gills of *A. colbecki* exposed to different experimental conditions. Letters indicate significant differences between groups of means. Data are given as mean values  $\pm$  standard deviations,  $n = 5$ .

Figure 4. Levels of malondialdehyde, lipofuscin, micronuclei, acyl CoA oxidase and acetylcholinesterase in tissues of *A. colbecki* exposed to different experimental conditions. Panels A, C, E: digestive gland; B, F: gills; D,G: haemolymph. Letters indicate significant differences between groups of means. Data are given as mean values  $\pm$  standard deviations,  $n = 5$ .

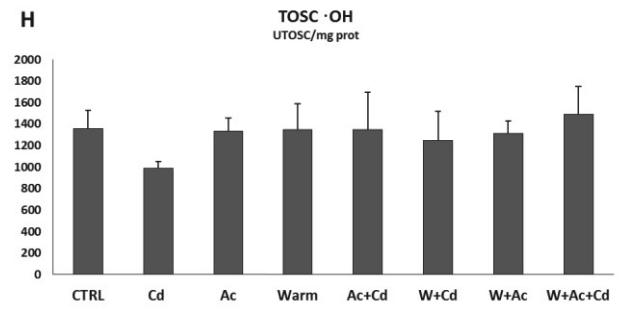
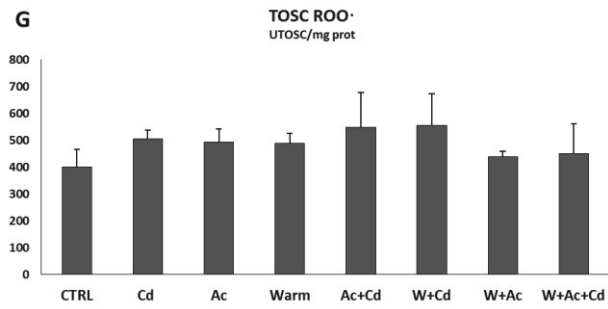
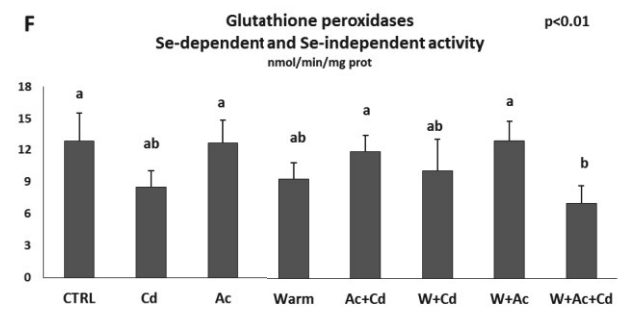
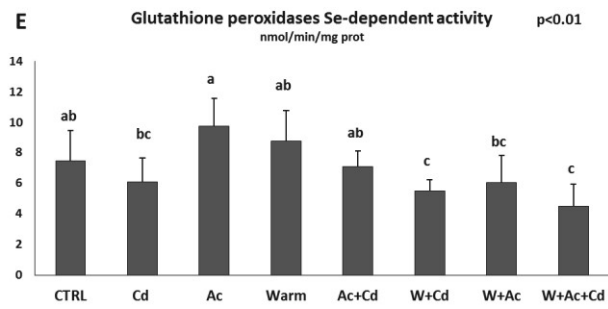
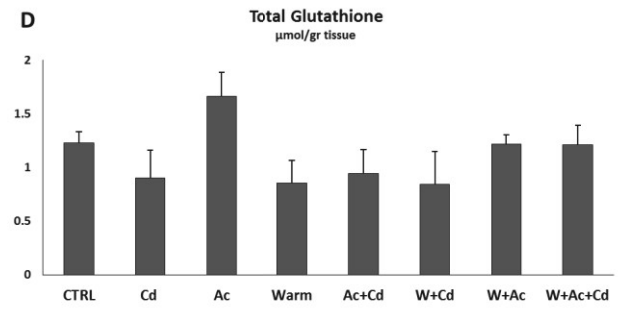
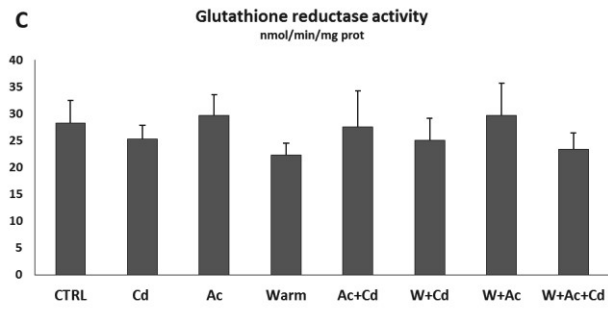
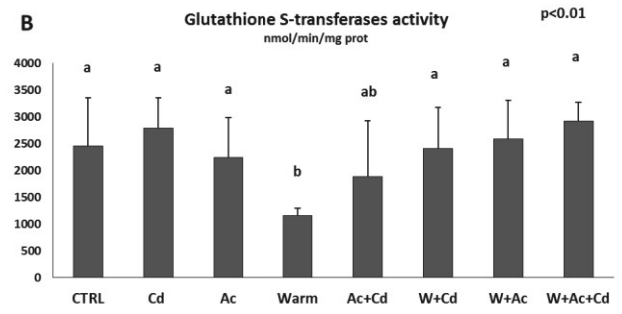
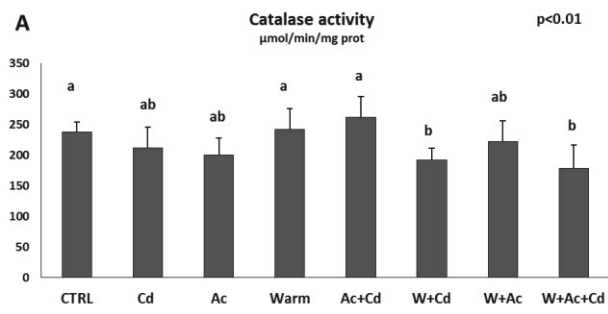
Figure 5. Non-metric multidimensional scaling analysis (MDS) ordination plots of biological responses in digestive glands (A) and in gills (B) of *A. colbecki* exposed to different experimental conditions. White points indicate samples exposed to environmental temperature (-1°C) and dark points indicate samples exposed to warm conditions (+1°C).

Figure 6. Weight of Evidence (WOE) classification of biomarkers data, in scallops exposed to different laboratory conditions. The quantitative Hazard Quotients (HQ) and the assigned classes of hazard are given.

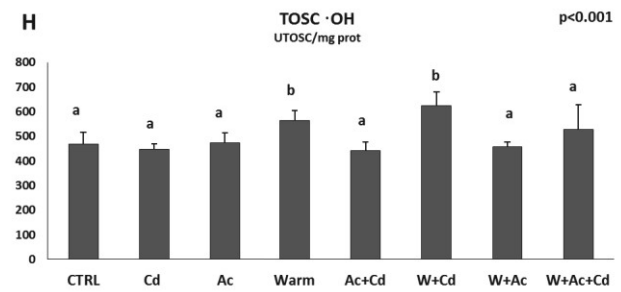
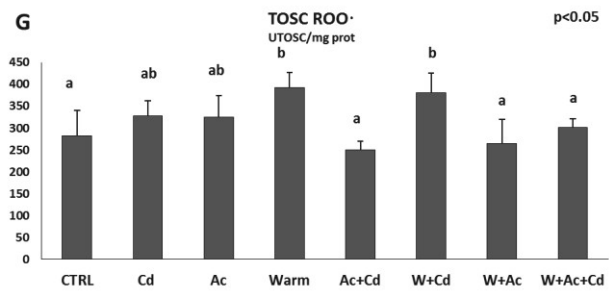
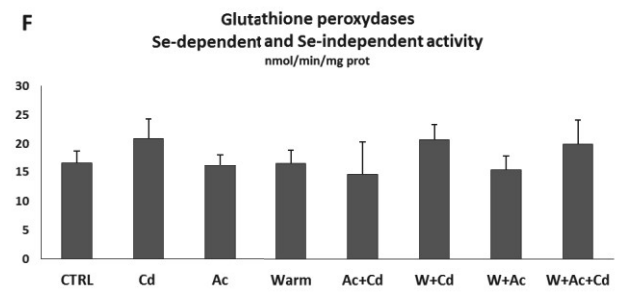
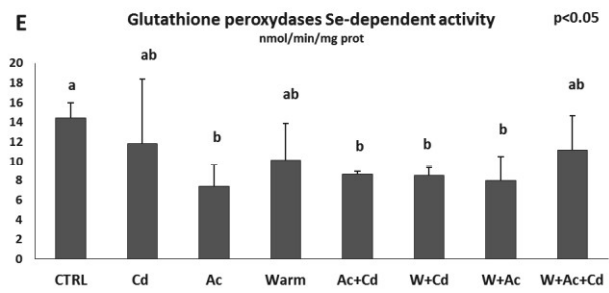
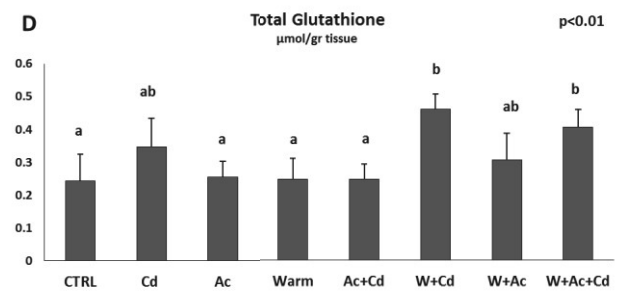
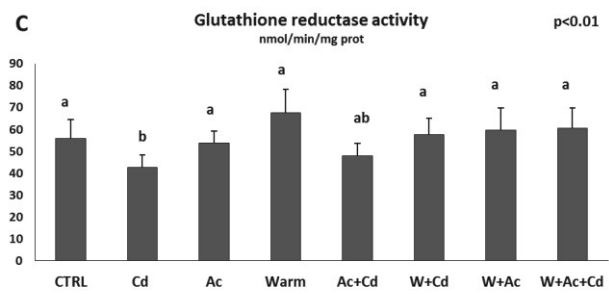
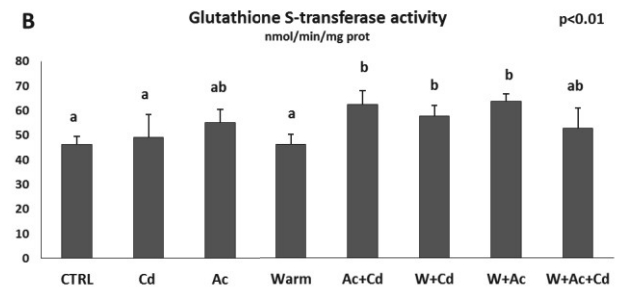
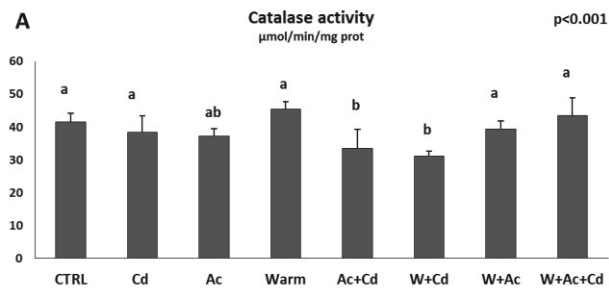


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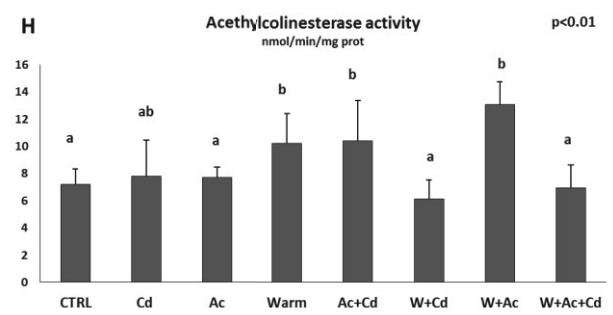
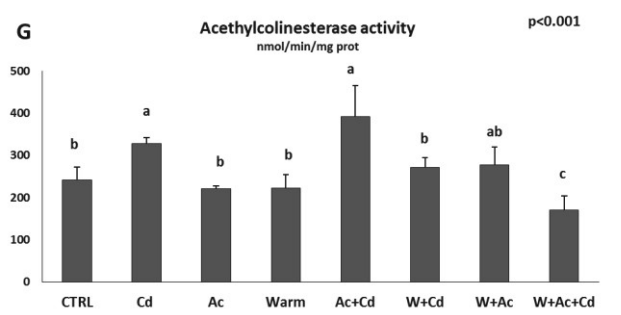
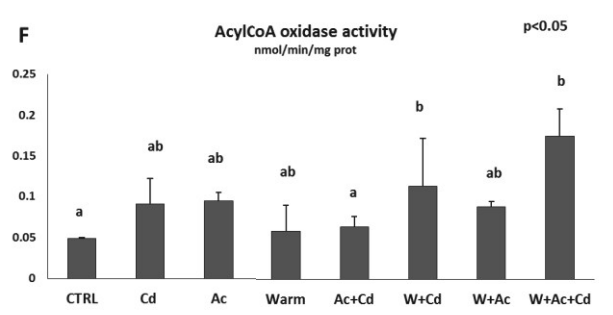
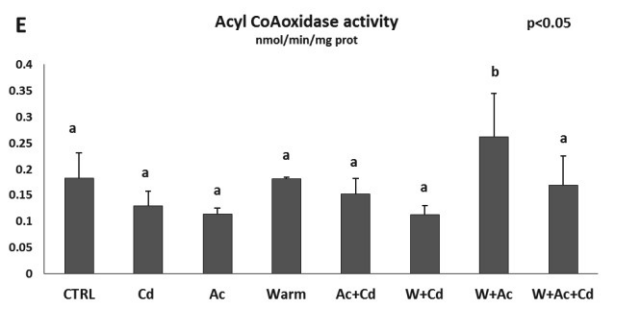
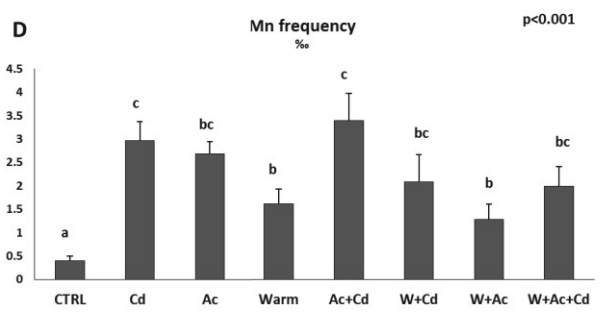
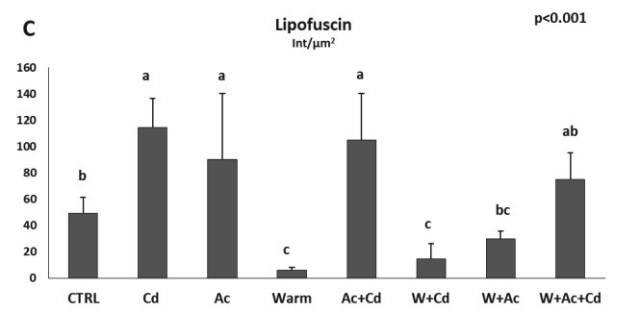
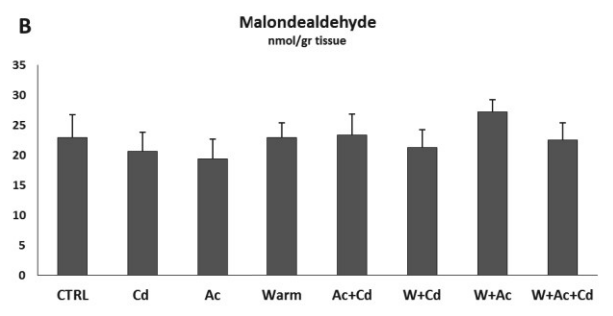
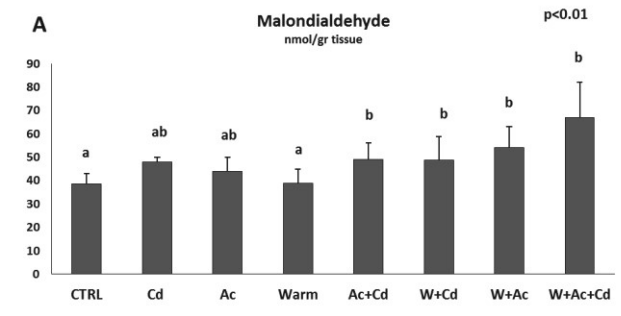


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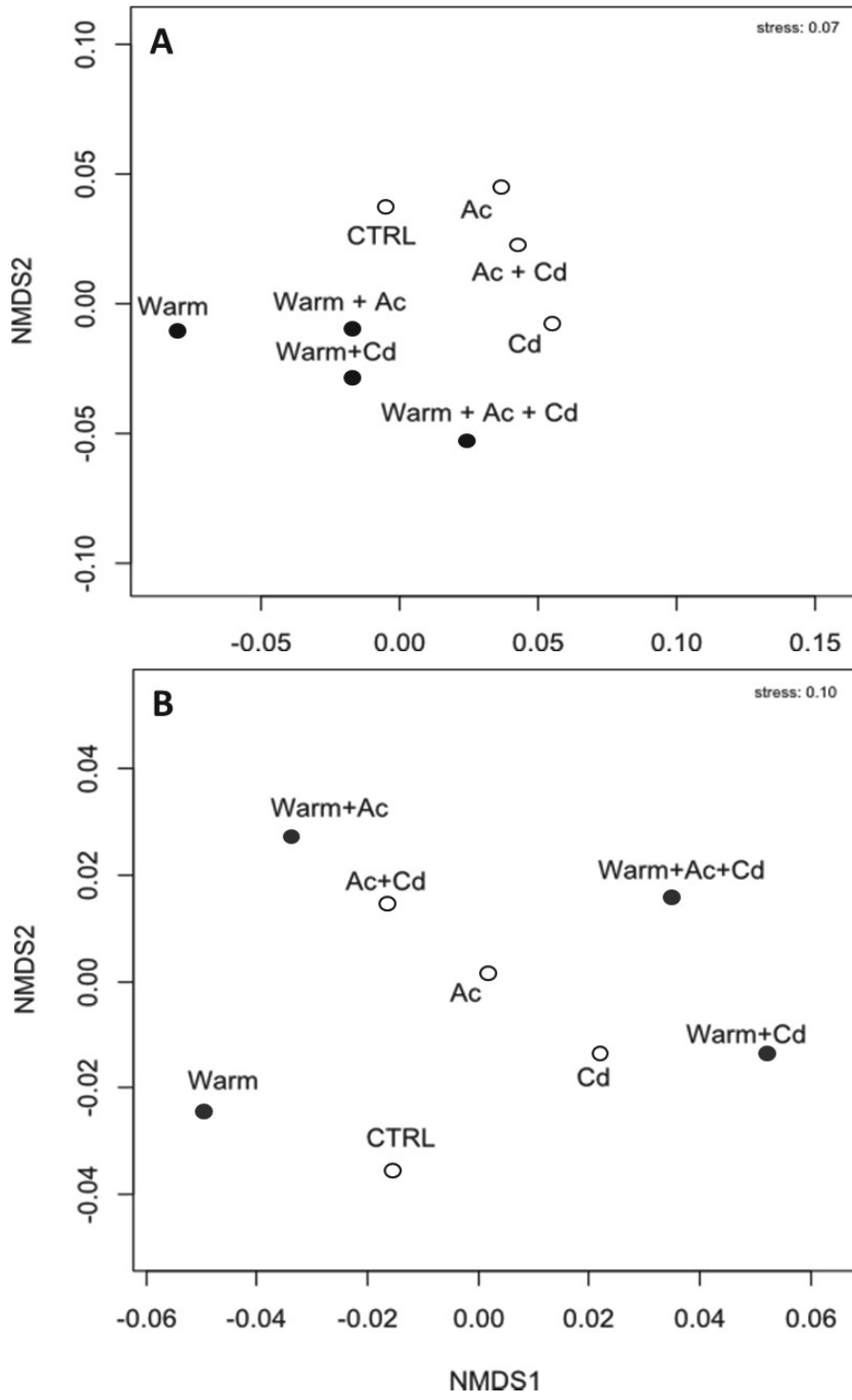
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Figure 3.



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















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22 Figure 5.

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Tissues	Experimental groups	Hazard Quotients (HQ)	Class of Hazard	Level
Digestive glands	Cd	8.9	Moderate	
Digestive glands	Ac	7.99	Slight	
Digestive glands	W	5.92	Slight	
Digestive glands	Ac+Cd	8.62	Moderate	
Digestive glands	W+Cd	6.13	Moderate	
Digestive glands	W+Ac	1.9	Slight	
Digestive glands	W+Ac+Cd	5.5	Moderate	
Gills	Cd	3.53	Slight	
Gills	Ac	5.75	Slight	
Gills	w	2.35	Slight	
Gills	Ac+Cd	1.61	Slight	
Gills	W+Cd	10.58	Moderate	
Gills	W+Ac	9.03	Moderate	
Gills	W+Ac+Cd	13.15	Moderate	

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25 Figure 6.

**Highlights:**

- Effects of multiple stressors were analysed in the Antarctic scallop *A. colbecki*
- Synergistic effects occurred between temperature, acidification and cadmium
- Oxidative responsiveness was evident in digestive gland and gills
- Cadmium was the primary stressor for digestive gland
- Gills were more sensitive to treatments at higher temperature