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Effects of ocean warming and acidification on accumulation and cellular responsiveness to cadmium in mussels Mytilus galloprovincialis: Importance of the seasonal status

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1	Effects of ocean warming and acidification on accumulation and cellular responsiveness to
2	cadmium in mussels Mytilus galloprovincialis: importance of the seasonal status.
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6	Alessandro Nardi ^a , Maura Benedetti ^{a,b} , Giuseppe d'Errico ^a , Daniele Fattorini ^a , and Francesco
7	Regoli ^{a,b*}
8	
9	
10	^a Dipartimento di Scienze della Vita e dell'Ambiente (DiSVA), Università Politecnica delle Marche,
11	Ancona, Italy
12	^b CoNISMa, Consorzio Interuniversitario per le Scienze del Mare, Roma, Italy
13	
14	
15	
16	
17	*Corresponding author: Prof. Francesco Regoli
18	Dipartimento di Scienze della Vita e dell'Ambiente (DiSVA),
19	Università Politecnica delle Marche,
20	Via Brecce Bianche 60131, Ancona, Italy
21	e-mail: f.regoli@univpm.it
22	
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24	bioaccumulation; cellular biomarkers
25	

Abstract

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Ocean warming and acidification could represent additional threat to marine organisms already coping with other anthropogenic impacts, such as chemical contamination in coastal areas. In this study, interactions between such multiple stressors and their synergistic effects in terms of accumulation, detoxification and biological effects of metals were investigated in the Mediterranean mussel Mytilus galloprovincialis. Organisms sampled during the winter period were exposed for 28 days to different combinations of two temperatures (10°C and 15°C), two pH/pCO₂ (8.20/~400µatm and 7.4/~3000µatm) and two doses of cadmium (0 and 20 µg/L). Cadmium concentrations increased in digestive glands and gills of metal-exposed mussels, and were further enhanced by co-exposure at higher temperature. Interactive effects of temperature and/or pH were observed on Cd-mediated metallothioneins induction, responsiveness of antioxidant system and onset of oxidative damages to lipids, with tissue-specific effects. Immunological effects showed a generalized sensitivity of lysosomal membrane stability toward the investigated stressors with major effects in co-exposed organisms. Cadmium and temperature affected phagocytosis efficiency and haemocytes population composition probably influencing the micronuclei frequency through varied mitotic rate. Several differences were highlighted between these results and those previously obtained from mussels exposed in summer, supporting the importance of season when addressing the tolerance of temperate organisms to variations of environmental factors. The elaboration of the whole biomarkers results through weighted criteria allowed to summarize specific hazard indices, highlighting tissue-specific sensitivity toward multiple stressors and the need of improving the knowledge on multiple stressors interaction.

1.Introduction

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Ocean warming and acidification are posing serious threats to marine ecosystems. Since the preindustrial values, ocean pH has dropped by 0.1 units (Gattuso and Lavigne, 2009) and global mean temperature has risen by almost 1°C (Hansen et al., 2016). According to projections of future scenarios, pH will further decrease down to 7.8 and temperature will rise of 2°C by 2100 (IPCC, 2013). These changes can induce direct effects on health status and physiological performance of marine organisms, but synergistic effects between multiple stressors may also occur with less documented consequences (Crain et al., 2008; Burton and Johnston, 2010; Kroeker et al., 2013). Metal contamination is a typical anthropogenic footprint in coastal areas (Doney, 2010; Bijima et al., 2013; Gilaranz et al., 2016) and both ocean warming and acidification, through different mechanisms, can individually influence distribution and fate of trace elements in sediments and seawater, as well as their bioaccumulation in marine organisms. Temperature may increase bioaccumulation of metals enhancing their solubility (Sokolova and Lannig, 2008), and the ventilation and feeding activity of marine organisms due to higher energy demand (Viarengo et al., 1988; Nichols and Playle, 2004; Baines et al., 2006; Cherkasov et al., 2007; Mubiana and Blust, 2007; Guinot et al., 2012; Negri et al., 2013; Coppola et al., 2018). On the other side, also CO₂-enrichment in seawater can alter the speciation and solubility of metals forming strong complexes with carbonate ions (Millero et al., 2009; Hoffmann et al., 2012; Stockdale et al., 2016) or with organic compounds (Gledhill et al., 2015). Increased fluxes of metals from sediments to seawater have been shown to occur at reduced pH values or simulating CO₂ leakages from sub-seabed (Riba et al. 2003; Ardelan et al., 2009; de Orte et al., 2014a, 2014b; Basallote et al., 2015), and a higher uptake due to acidification has been described in several marine invertebrates including bivalves (López et al., 2010; Ivanina et al., 2014; Götze et al., 2014; Rodríguez-Romero et al., 2014a; Shi et al., 2016; Velez et al., 2016), polychaetes

(Rodríguez-Romero et al., 2014b), and cephalopods (Lacoue-Labarthe et al., 2009, 2011).

Various cellular mechanisms have been suggested to influence the vulnerability of marine organisms to metals toxicity when also exposed to variations of temperature and pH/pCO₂. The interaction between temperature and metals has been demonstrated to cause impairment of mitochondrial function, pro-oxidative mechanisms, accumulation of lipid peroxidation products and damages to lysosomes and DNA (Sokolova, 2004; Kefaloyianni et al., 2005; Cherkasov et al., 2007; Mubiana and Blust, 2007; Sokolova and Lannig, 2008; Guinot et al., 2012; Negri et al., 2013; Attig et al., 2014; Banni et al., 2014; Gomiero and Viarengo, 2014; Izagirre et al., 2014; Múgica et al., 2015). Similarly, interactive effects between metals and acidification are addressed in impaired larval development, onset of DNA damage, pro-oxidative mechanisms and lower antioxidant efficiency, alterations of the immune function (Lewis et al., 2013, 2016; Roberts et al., 2013; Campbell et al., 2014; Ivanina et al., 2015, 2016; Siddiqui and Bielmyer-Fraser, 2015). Recently, cadmium (Cd) accumulation was shown to be unaffected by variations of temperature and pH in the digestive gland and gills of the Mediterranean mussel M. galloprovincialis and of the Antarctic scallop A. colbecki. At the same time, however, the effects of Cd on metal-binding proteins, immune system, antioxidant responses and oxidative stress biomarkers were strongly modulated by these co-factors, with different effects and magnitude depending on species and tissue (Benedetti et al., 2016; Nardi et al., 2017). Beside species- and tissue-mediated specificity, seasonality of environmental factors (notably temperature) and biological processes (i.e. reproductive cycle) could be responsible for the different capability of organisms to tolerate variations of temperature and pH related to climate change and their interactions with pollutants. In this respect, several cellular responses typically involved in defence mechanisms and resistance to stressors exhibit marked seasonal fluctuations in marine invertebrates and the ecophysiological performance of organisms is also influenced by their seasonal metabolic status and energy available for processes like reproduction and growth (Ringwood et al., 2002; Bocchetti and Regoli, 2006; Farcy et al., 2007; Bocchetti et al.,

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2008; Pereira et al., 2012).

Considering the complexity of interactions between environmental and biological factors, this study aimed to elucidate whether seasonality can influence the responsiveness of M. galloprovincialis to various combinations of temperature, pH and Cd: experiments were performed in winter and results compared with those obtained in a similar study carried out in summer season (Nardi et al., 2017). Beside Cd accumulation, a wide battery of biomarkers was chosen to reflect the network of cellular mechanisms underlying metal detoxification, oxyradical metabolism and onset of different forms of cellular toxicity (Regoli and Giuliani, 2014): these biomarkers included induction of metallothioneins, variations of single antioxidant defenses, total oxyradical scavenging capacity, accumulation of lipid peroxidation products in digestive gland and gills, while lysosomal membrane stability, phagocytosis, granulocytes/hyalinocytes ratio, DNA strand breaks and micronuclei frequency were measured in haemocytes. To better summarize the biological significance of obtained results and to facilitate the seasonal comparison of mussels susceptibility, the overall biomarkers results have been elaborated through a widely validated quantitative hazard model (Sediqualsoft): this model applies weighted criteria based on the toxicological relevance of each analyzed biomarker and on the magnitude of observed variations to summarize a synthetic cellular hazard index (Piva et al., 2011; Benedetti et al., 2012).

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The obtained results are intended to expand our knowledge on the impacts of synergistic effects of multiple stressors on cellular and tissue health status, and to relate them with the physiology of mussels as a response to seasonal change.

2.Materials and Methods

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2.1 Animal collection and experimental design

Mussels, M. galloprovincialis (6.0 ± 0.5 cm shell length), were obtained in January 2015 from a shellfish farm in an unpolluted area of Central Adriatic Sea (Regoli et al., 2014) and maintained for 7 days in aquaria with aerated artificial seawater (ASW; Instant Ocean®) at local environmental conditions of salinity (37 practical salinity units), temperature (10°C) and pH (8.20).

After acclimation, mussels were randomly assigned and exposed to one of the following treatments, each containing 36 organisms in 20 L tanks: 1) control condition (CTRL), at environmental temperature of 10°C, normocapnia with pH=8.20/ $pCO_2=\sim400 \mu atm$; 2) Cd exposure (Cd), 10°C, pH=8.20/ pCO₂= \sim 400 μ atm and 20 μ g/L Cd; 3) acidification (A), 10°C, hypercapnia with pH=7.40/ $pCO_2=\sim3000 \mu atm$; 4) warming (W), 15°C and pH=8.20/ $pCO_2=\sim400 \mu atm$; 5) acidification + Cd (A-Cd), 10° C, pH=7.40/ pCO₂=~3000 μ atm and 20 μ g/L Cd; 6) warming + Cd (W-Cd), 15° C, pH=8.20/pCO₂=~400 μ atm and 20 μ g/L Cd; 7) acidification + warming (A-W), 15° C and pH=7.40/ $pCO_2=\sim3000~\mu$ atm; 8) acidification + warming + Cd (A-W-Cd), 15°C, pH=7.40/ $pCO_2 = 3000 \mu$ atm and 20 μ g/L Cd. Despite the effects related to climate changes are expected to occur over the course of decades, organisms were exposed to the tested environmental conditions of Cd, pH and temperature without gradual acclimation, thus simulating sudden changes as those more easily occurring in coastal or estuarine areas. The same experimental design had been previously applied on mussels sampled during the summer with the exception of temperature values, respectively at 20 and 25 °C for the control and warming treatments (Nardi et al., 2017). Cd dose is representative for a polluted but environmentally realistic condition in coastal waters, typically ranging from less than 1 up to hundreds μ g/L in highly polluted areas (Neff, 2002; Bakary et al., 2015). Lowered pH was adapted from scenario RCP 8.5 and IPCC WGII AR5 (IPCC, 2014; Wong et al., 2014) reporting a mean pH value of 7.7 for open oceans, but predicting more pronounced reductions in coastal areas. The experimental pH was reached by adding to each treatment ASW (pH=8.2) small and defined amounts of CO₂-saturated ASW, obtained by bubbling pure CO₂ in ASW for at least 24h, until reaching the target pH within a couple of hours (Nardi *et al.*, 2017). Salinity and pH were measured daily, while total alkalinity (A_T) was measured twice per week according to Dickson *et al.* (2007), and used for calculating seawater carbonate parameters (pCO_2 , and saturation state (Ω) for calcite and aragonite) in CO2SYS (Pierrott *et al.*, 2006) using barometric pressure values (full seawater chemistry is provided in Table 1). For calculations, we used NBS scale for seawater pH, carbonate constants from Millero (2010), KSO_4^- constant from Dickson *et al.* (2007), and concentrations of silicate and phosphate from Instant Ocean® seawater (0.21 μ mol/kg and 0.05 μ mol/kg, respectively). Water in each treatment was changed every other day using water at the same pH and temperature to avoid fluctuations of these parameters during the exposure period. Mussels fed 12 hours prior the water change with a commercial mixture of zooplankton for filter-feeding organisms (Zooplanktos-S 50-300 μ m, Brightwell Aquatics, Elysburg, PA).

After four weeks, animals were sampled for chemical and biological analyses. Specifically gills and digestive glands were excised from the 36 specimens for each treatment, pooled in 12 samples, each constituted by the tissues of 3 individuals, rapidly frozen in liquid nitrogen and maintained at -80°C: these 12 pooled samples were shared for analyses of Cd and biomarker responses, to guarantee a n value = 5 for replicates analysed for each parameter in each treatment. Haemolymph was also withdrawn from the adductor muscle of 5 specimens for each treatment and immediately used for measurement of immunity parameters and genotoxic damage.

2.2 Cd determination

Concentration of Cd in digestive gland and gills of mussels was analyzed according to previously described methods (Regoli *et al.*, 2005). For each treatment, tissues were dried at 60°C overnight and digested in a microwave system (Mars V, CEM). After digestion samples were analyzed by atomic absorption spectrophotometry with graphite furnace atomization and Zeeman effect. Quality assurance and control were assessed by processing blank samples and reference standard material (Mussel Tissue Standard Reference Material SRM NIST-2977, National Institute of Standards and Technology Gaithersburg, MD, USA), which always resulted within the 95%

confidence interval of certified values. Data are expressed as μ g/g dry weight (mean value \pm standard deviation, n = 5), limits of quantification and detection were respectively 0.002 and 0.001 μ g/g.

2.3 Biomarkers responses

Standardized methods, described in detail in Supplementary Material 1 (SM1), were used for biomarkers analyses. Digestive gland and gills were analyzed for metallothioneins (MTs), antioxidant enzymes and scavenger (catalase, glutathione S-transferases, glutathione peroxidases, glutathione reductase, total glutathione), total oxyradical scavenging capacity toward peroxyl radicals (TOSC ROO•) and hydroxyl radicals (TOSC HO•), malondialdehyde content (MDA). Lipofuscin and neutral lipids were histochemically determined on 8 μ m thick cryostat sections of digestive glands. Haemocytes were immediately processed for immune-related alterations, such as lysosomal membrane stability (neutral red retention time, NRRT), phagocytosis activity and granulocytes versus hyalinocytes ratio. Also genotoxic effects were evaluated in haemocytes in terms of percentage of DNA integrity (Comet assay) and micronuclei frequency (MN).

2.7 Statistical analyses

Cd accumulation and biological responses results were submitted to analysis of variance (One-way ANOVA), after checking the normal distribution (Shapiro-Wilk test) and homogeneity of variances (Levene's Test). The null hypothesis tested was that no significant difference existed between different treatments, level of significance was set to p < 0.05, and post-hoc tests (Student–Newman–Keuls, SNK), were used to compare group of means. All statistical analyses were performed using RStudio (version 0.99.491).

The overall significance of biomarkers results was summarized in a cellular hazard index elaborated through a previously developed quantitative model which applies weighted criteria to discriminate different endpoints and the magnitude of effects (Sediqualsoft, Piva et al., 2011). Despite whole calculations and assumptions have been fully given elsewhere (Piva et al., 2011; Benedetti et al., 2012), the general rationale of the model is to assign a weight to various biomarkers depending on their toxicological relevance and to compare variations of responses toward a threshold specific

for each biomarker, which considers the different responsiveness among various species and tissues, and the possibility of biphasic responses. The calculated Hazard Quotient (HQ) does not include the contribution of biomarkers with variations lower or equal to their threshold, and it averages or adds the effects respectively for those biomarkers with variations up to 2-fold or more than 2-fold greater than the specific threshold (Piva *et al.*, 2011; Benedetti *et al.*, 2012, 2014; Regoli *et al.*, 2014; Avio *et al.*, 2015). The model finally assigns the elaborated HQ to one of five classes of hazard, from Absent to Severe (Piva *et al.*, 2011).

3. Results

A significant increase of Cd concentrations was observed in digestive gland and gills of mussels exposed to this metal (Fig.1a, b), and further enhanced in both the tissues by co-exposure at higher temperature (Cd and A-Cd *versus* W-Cd and A-W-Cd) (Fig. 1a, b).

Metallothioneins in the digestive gland were not significantly induced in organisms exposed to Cd alone, while increasing in those co-exposed to the metal at reduced pH and/or higher temperature (A-Cd, W-Cd and A-W-Cd) (Fig. 1c). In gills, metallothioneins were significantly enhanced only in organisms exposed to Cd at higher temperature and control pH (W-Cd) (Fig.1d).

Antioxidant defenses showed tissue-specific effects of the investigated factors. In the digestive gland, no statistically significant variations were observed in various treatments for catalase, GSTs, Se-dependent GPx and TGSH (Fig. 2a, b, c, f). On the other hand, Cd caused a marked induction of total GPx, particularly evident at control pH (compare CTRL vs. Cd and W vs. W-Cd); at reduced pH, Cd did not further modulate the increase of total GPx caused by acidification itself (A vs. A-Cd, and A-W vs. A-W-Cd) (Fig. 2d). Higher temperature was the only treatment to significantly reduce GR activity in digestive gland (Fig. 2e). Total oxyradical scavenging capacity toward peroxyl radical (TOSC ROO•), levels of malondialdehyde and lipofuscin were generally enhanced by treatments involving Cd alone and/or in combination with other stressors, but the statistical significance was obtained only for TOSC and lipofuscin in mussels co-exposed to Cd, acidification and higher

temperature (A-W-Cd) (Fig. 2g, i, j). Conversely, neutral lipids content was reduced in all Cd-exposed organisms at reduced pH and/or higher temperature (A-Cd, W-Cd and A-W-Cd) compared to organisms exposed to Cd alone (Fig. 2k).

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In gills, significant variations of antioxidant enzymes were observed for glutathione Stransferases, selenium-dependent and total glutathione peroxidases. GSTs were inhibited in mussels exposed to Cd alone, Cd at higher temperature (W-Cd) and higher temperature at reduced pH (A-W) (Fig. 3b). Opposite effects were observed on Se-dependent GPx which were increased by Cd and temperature as individual stressors (Cd and W) and by the combination of these factors with acidification (A-Cd, A-W, A-W-Cd) (Fig. 3c). Enhanced values were also observed for total GPx activity, particularly for treatments involving higher temperature and/or acidification (Fig. 3d). Limited variations were measured in the total oxyradical scavenging capacity toward hydroxyl radical (TOSC HO•) with slightly higher values in organisms exposed to Cd or higher temperature (Fig. 3h). Lysosomal membrane stability in haemocytes significantly decreased in all experimental treatments, with major effects in organisms co-exposed to all multiple stressors (A-W-Cd) (Fig. 4a). Phagocytosis rate (Fig. 4b) was reduced in mussels treated at higher temperature (alone or in combination with Cd), and also granulocytes/hyalinocytes ratio (Fig. 4c) was significantly reduced in organisms co-exposed to Cd at higher temperature (W-Cd and A-W-Cd). No significant variations were observed in terms of DNA strand breaks (Fig. 4d) while micronuclei frequency was generally enhanced in all organisms exposed to Cd alone or with higher temperature (Fig. 4e).

Biomarkers responses observed in each experimental condition were summarized in a single hazard index through the application of weighted criteria (Fig. 5). The elaborated class of hazard revealed that digestive gland was generally more sensitive to Cd dosed alone and in combination with A or W, when the assigned class of hazard was "Moderate", compared to "Slight" in all the other treatments (Fig. 5). In gills, higher values of HQ were typically measured compared to digestive gland, and variations of biomarkers were generally reflected by a "Moderate" class of hazard (Fig.

4. Discussion

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This study provided new evidence that climate change can affect accumulation and responsiveness to metals in marine organisms.

The obtained results highlighted an increased Cd accumulation in mussels co-exposed to the metal at higher temperature during the winter period, contrasting with our previous study in which this effect was not observed during the summer (Nardi et al., 2017). The winter environmental temperature was 10°C and the increased value selected for this study (15°C) was still below the upper thermal limit of M. galloprovincialis (Anestis et al., 2007; Gazeau et al., 2014). On the other hand, the environmental and warmer values used for summer exposures were 20° and 25°C (Nardi et al., 2017). In this respect, only during the colder period the increase of temperature would have enhanced metabolism rate and consequent accumulation of metals, confirming that the effects of climate-related stressors are closely related to thermal tolerance of organisms (Ioannou et al., 2009; Pörtner, 2010), as confirmed in mussels exposed to thermal stress (21°C). The effect of temperature on enhanced Cd accumulation was noticed both in digestive gland and gills, suggesting a whole organism, physiological response, rather than a tissue-specific pathway. The different seasonal effects of temperature on Cd accumulation suggests that future warming scenarios should take into account this variability in temperate marine organisms, as M. galloprovincialis, which experience wide natural fluctuations of this environmental parameter. The metal uptake was not further modulated by reduction of pH. Similar results were previously observed in mussels exposed during the summer and in scallops from Antarctic region (A. colbecki) (Benedetti et al., 2016; Nardi et al., 2017), while a slight influence of lowered pH on Cd accumulation occurred in the smooth Mediterranean scallop Flexopecten glaber (Nardi et al., 2018). Overall these results confirm the more limited influence of acidification compared to temperature on the bioaccumulation of Cd, despite a certain influence of species-specific characteristics.

Metallothioneins induction is typically associated to the increase of tissue metal concentrations, but a similar parallelism was not always observed in our study. In both digestive gland and gills, MTs

were not induced in mussels exposed to Cd alone, suggesting that basal levels of these proteins were able to compensate for the increased metal content. The lack of correlations between Cd bioaccumulation and MTs induction observed during co-exposures supports the modulation on protein synthesis by oxidative pressure, confirming the role of prooxidant mechanisms caused by synergistic effects of multiple stressors, as also previously observed in mussels exposed during the summer (Viarengo, 2000; Regoli and Giuliani, 2014; Nardi *et al.*, 2017).

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The oxidative challenge was assessed through the analysis of an extensive array of biomarkers which included antioxidant defenses, total oxyradical scavenging capacity and oxidative damages. In digestive gland, total GPx appeared as responsive enzymes, particularly toward Cd and acidification. However, the more relevant results in terms of varied susceptibility to oxidative stress were observed in mussels co-exposed to Cd, acidification and higher temperature. In these treatments, the significant increase of total oxyradical scavenging capacity toward peroxyl radicals and of lipofuscin content, confirm the importance of oxidative pathways in modulating the responsiveness of marine organisms to multiple stressors, as also suggested for *Hediste diversicolor* co-exposed to Hg and pH reduction (Freitas et al., 2017). Neutral lipids content tended to increase in digestive gland after exposure to Cd alone and to decrease during co-exposures to multiple stressors, suggesting a greater use of reserve materials under increasing cellular stress, as already hypothesized for M. galloprovincialis challenged by trace metals and organic pollutants (Regoli, 1992; Koukouzika and Dimitriadis, 2008). As for MTs, also variations of oxidative parameters in digestive gland did not correlate with Cd content, and thus confirmed the occurrence of synergistic cellular effects between multiple stressors. Compared to actual results, mussels previously exposed during the summer showed a greater sensitivity of antioxidants in digestive gland toward temperature rise (alone or in combination with Cd), further indicating that pathways of response and susceptibility to changes of environmental stressors can vary between seasons.

Different and more frequent oxidative variations were observed in gills. Cd, higher temperature and acidification (alone or in combination) were generally responsible for a certain reduction of GSTs

and the induction of Se-dependent and Se-independent GPx, confirming the involvement of these antioxidant pathways also reported for mussels exposed during the summer (Nardi *et al.*, 2017). The results obtained in gills at higher temperature, conversely to digestive gland, highlight that metabolic and physiological functions of tissues may be responsible for their different responsiveness and sensitivity toward the investigated stressors. The increase of winter temperature might exert oxidative challenge in the gills due to enhanced filtration rates, thus explaining the raise of antioxidant protection (Fields *et al.*, 2012). The higher enzymatic activities caused by individual stressors were not synergistically enhanced during co-exposures, and were not typically paralleled by considerable variations of the total oxyradical scavenging capacity; the concomitant lack of changes in malondialdehyde content indicates that reported variations of antioxidant system were efficient in preventing an oxidative impairment in gills: once again, winter results are partly in contrast with those obtained in summer experiments when the total oxyradical scavenging capacity was decreased toward hydroxyl radicals (Nardi *et al.*, 2017). The generally higher efficiency of antioxidant protection observed during the winter reinforce the evidence that responsiveness of mussels to climate changes is influenced by seasonal-specific features.

Lysosomal membranes are well known targets of oxyradical toxicity and their destabilization is a typical effect of chemical, environmental or biological factors. In our study, lysosomal impairment was a reliable biomarker toward the investigated stressors, further evidencing the occurrence of synergistic effects during co-exposures to multiple factors. At the same time, temperature, alone or in combination with Cd lowered phagocytosis rate, probably due to a reduction of granulocytes, which are the haemocytes responsible of this activity (Gorbi *et al.*, 2013). The generalized impairment of lysosomal membranes stability and phagocytosis rate confirm the sensitivity of haemocytes in marine invertebrates and their utility as early warning signal of environmental disturbance (Beesly *et al.*, 2008; Matozzo *et al.*, 2012; Sureda et al., 2013; Nardi *et al.*, 2017). Sensitivity of haemocytes to Cd, both alone and in combination with temperature was supported also by the increase in micronuclei frequency which, considering the absence of changes in DNA strand breaks, might be modulated by

changes in mitotic frequency, rather than a direct genotoxic mechanism, as previously hypothesized in similar experiments (Nardi *et al.*, 2017; Benedetti *et al.*, 2016).

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The biological significance of the observed biomarker responses was better summarized by the elaboration of specific hazard indices based on weighted criteria, which consider both the magnitude of variations and the relative toxicological importance of investigated parameters (Piva et al., 2011; Benedetti et al., 2012). Biomarkers have been widely used for early warning detection of environmental disturbance, and various integrative methods and health indices have been recently developed to improve their use in ecological risk assessment (Beliaeff and Burgeot, 2002; Piva et al., 2011; Broeg et al., 2005; Dagnino et al., 2008; Benedetti et al., 2012; Marigómez et al., 2013). The elaboration applied in the present study is part of a more articulated Weight Of Evidence model (Sediqualsoft) which integrates multiple typologies of data including sediment and seawater chemistry, bioaccumulation, biomarker, ecotoxicological bioassays and benthic communities: integrative rules and assumptions have been detailed in previous papers and already validated in several risk assessment studies (Benedetti et al., 2012, 2014; Regoli et al., 2014; Avio et al., 2015; Bebianno et al., 2015). The hazard indices derived from biomarker results highlighted that, toward individual stressors, digestive gland is more sensitive to variations of Cd (HQ Moderate) compared to acidification or warming alone (HQ Slight): despite different combinations of multiple stressors were shown to highlight synergistic effects on various biological responses and mechanisms of action, they did not determine an overall increase of the assigned class of cellular.

Gills typically exhibited a Moderate hazard index suggesting that the more direct contact with seawater, and the lower basal metabolic activity could render these tissues more susceptible to fluctuations of environmental factors. Once again, the variations observed for various biomarker responses after contemporary exposure to multiple stressors did not change the elaborated class of cellular hazard indices compared to individual factors:

Overall, the sensitivity of winter mussels is partly in contrast with results obtained in summer season, when organisms exhibited more clear synergistic effects of Cd dosed in association to

warming and/or acidification, highlighting the importance of seasonal-specific sensitivity of temperate organisms toward multiple stressors. Our data suggest also the need to still investigate tissue-specific sensitivity, mechanisms of action related to physiological functions, and seasonal-dependent effects to better understand tolerance to climate changes and consequences of multiple stressors in marine organisms. Finally, the application of weighted criteria to elaborate hazard indices was confirmed as a fundamental procedure to summarize the biological significance of large datasets of complex biomarker data, thus facilitating prediction and comparisons between the effects of individual and multiple stressors.

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Table 1 - Summary of water chemistry parameters during experimental exposure. S (salinity), T (temperature), pHNBS (pH calibrated with National Bureau of Standard scale), AT (total alkalinity), pCO2 (partial pressure of CO2), Ω c and Ω a (saturation state of respectively calcite and aragonite). Data are presented as means \pm standard deviations.

Treatment	measured parameters				calculated parameters			
rreatment	S	T (°C)	pH _{NBS}	A _τ (μmol/kg)	<i>p</i> CO2 (μatm)	Ωc	Ωa	
CTRL	37 ± 0.5	9.95 ± 0.11	8.18 ± 0.03	3283.2 ± 88.8	386.9 ± 26.2	6.2 ± 0.5	3.9 ± 0.3	
Cd	37 ± 0.5	9.97 ± 0.06	8.16 ± 0.03	3334.2 ± 102.6	411.8 ± 35.8	6.1 ± 0.3	3.9 ± 0.2	
Α	37 ± 0.5	10.54 ± 0.08	7.40 ± 0.05	3364.1 ± 112.9	2882.2 ± 363.8	1.3 ± 0.2	0.8 ± 0.1	
W	37 ± 0.5	14.95 ± 0.12	8.17 ± 0.03	3378.4 ± 121.2	416.8 ± 34.5	7.2 ± 0.5	4.7 ± 0.4	
A-Cd	37 ± 0.5	10.04 ± 0.15	7.39 ± 0.02	3360.6 ± 36.8	2860.9 ± 207.2	1.2 ± 0.1	0.8 ± 0.1	
W-Cd	37 ± 0.5	15.02 ± 0.11	8.17 ± 0.03	3350.5 ± 164.9	403.5 ± 53.1	7.3 ± 0.5	4.7 ± 0.3	
A-W	37 ± 0.5	14.98 ± 0.05	7.39 ± 0.04	3354.5 ± 80.1	2916.3 ± 288.7	1.5 ± 0.2	1.0 ± 0.1	
A-W-Cd	37 ± 0.5	14.92 ± 0.06	7.39 ± 0.02	3326.5 ± 67.1	2886.4 ± 174.1	1.5 ± 0.1	1.0 ± 0.1	

625 Figure 1 – Cd concentrations (μ g/g dry weight) and level of metallothioneins (nmol/mg of proteins) in digestive 626 gland (a and c) and gills (b and d) of mussels exposed to various treatments. Data are given as mean 627 values ± standard deviations (n=5). Different letters indicate significant differences between group of means 628 (ANOVA and SNK post-hoc). CTRL= Control; Cd= Cadmium; A= acidification; W= warming; A-Cd= 629 acidification + Cd; W-Cd= warming + Cd; A-W= acidification + warming; A-W-Cd= acidification + warming + 630 Cd. 631 Figure 2 – Antioxidant defenses and oxidative stress biomarkers in digestive gland of mussels exposed to various 632 treatments. CAT: catalase (µmol/mg proteins) (a), GSTs: glutathione S-transferases (nmol/mg proteins) (b), Se-Dep. GPx: 633 Se-dependent glutathione peroxidases (nmol/mg proteins) (c) total GPx: sum of Se-dependent and Se-independent 634 glutathione peroxidases (nmol/mg proteins) (d), GR: glutathione reductase (nmol/mg proteins) (e), TGSH: total 635 glutathione (µmol/g wet weight) (f), TOSC ROO: total oxyradical scavenging capacity toward peroxyl radical (TOSC 636 units/mg proteins) (g), TOSC HO: total oxyradical scavenging capacity toward hydroxyl radical (TOSC units/mg 637 proteins) (h), MDA: levels of malondialdehyde (mmol/g wet weight) (i), lipofuscin (intensity/µm²) (j) and neutral lipids 638 (intensity/ μ m²) (k). Data are given as mean values ± standard deviations (n=5). Different letters indicate significant 639 differences between group of means (ANOVA and SNK post-hoc). CTRL= Control; Cd= Cadmium; A= acidification; 640 W= warming; A-Cd= acidification + Cd; W-Cd= warming + Cd; A-W= acidification + warming; A-W-Cd= acidification 641 + warming + Cd. 642 Figure 3 - Antioxidant defenses and oxidative stress biomarkers in gills of mussels exposed to various treatments. CAT: 643 catalase (µmol/mg proteins) (a), GSTs: glutathione S-transferases (nmol/mg proteins) (b), Se-Dep. GPx: Se-dependent 644 glutathione peroxidases (nmol/mg proteins) (c) total GPx: sum of Se-dependent and Se-independent glutathione 645 peroxidases (nmol/mg proteins) (d), GR: glutathione reductase (nmol/mg proteins) (e), TGSH: total glutathione (µmol/g 646 wet weight) (f), TOSC ROO: total oxyradical scavenging capacity toward peroxyl radical (TOSC units/mg proteins) (g), 647 TOSC HO•: total oxyradical scavenging capacity toward hydroxyl radical (TOSC units/mg proteins) (h), MDA: levels 648 of malondialdehyde (mmol/g wet weight) (i). Data are given as mean values ± standard deviations (n=5). Different letters 649 indicate significant differences between group of means (ANOVA and SNK post-hoc). CTRL= Control; Cd= Cadmium; 650 A= acidification; W= warming; A-Cd= acidification + Cd; W-Cd= warming + Cd; A-W= acidification + warming; A-W-651 Cd= acidification + warming + Cd. 652 Figure 4 - Lysosomal membrane stability (a), phagocytosis rate (b), granulocytes/hyalinocytes ratio (c), DNA damage 653 (d) and frequency of micronuclei (e) in haemocytes of mussels exposed to various treatments. Data are given as mean 654

values ± standard deviation (n=5). Different letters indicate significant differences between group of means (ANOVA

and SNK *post-hoc*). CTRL= Control; Cd= Cadmium; A= acidification; W= warming; A-Cd= acidification + Cd; W-Cd=
warming + Cd; A-W= acidification + warming; A-W-Cd= acidification + warming + Cd.
Figure 5 - Weight of Evidence (WOE) classification of biomarkers data for the whole dataset of analyzed
parameters in digestive gland and gills for each exposure condition. The quantitative hazard quotients (HQ) and the
assigned class of hazard are given.

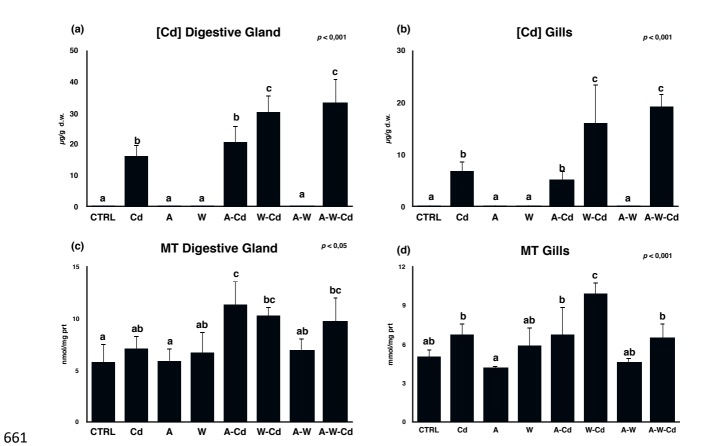
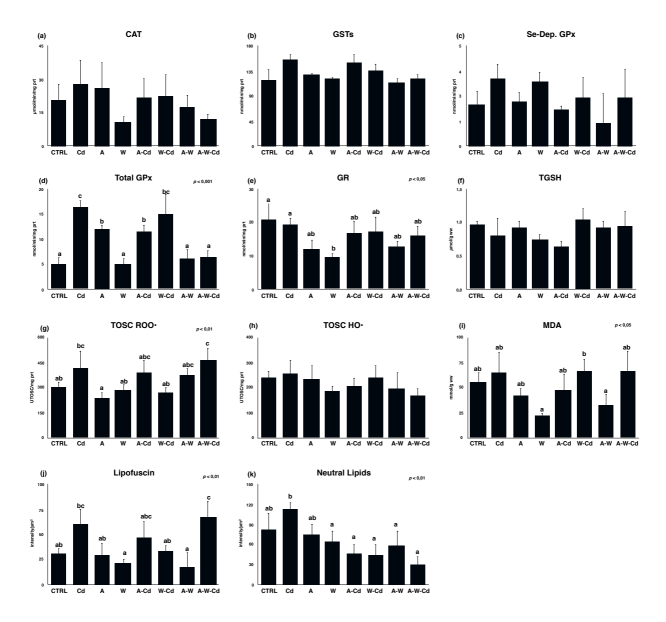
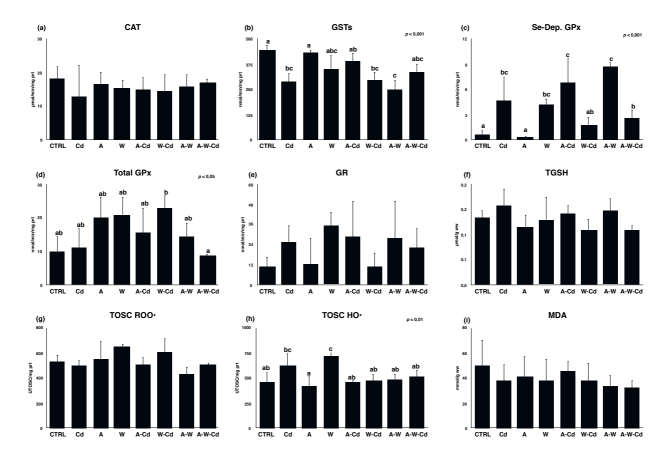


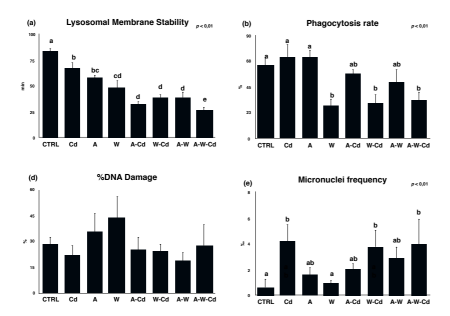
Figure 1.



665 Figure 2.



667 Figure 3.



(c)

Granulocytes/Hyalinocytes

p < 0,01

669 Figure 4.

		Digestive gland		Gills			
Experimental Treatment	Hazard Quotient (HQ)	Class of Hazard	Level	Hazard Quotient (HQ)	Class of Hazard	Level	
Cd	9.87	MODERATE		13.82	MODERATE		
А	4.60	SLIGHT		7.48	MODERATE		
w	2.31	SLIGHT		31.16	MODERATE		
A-Cd	6.66	MODERATE		39.55	MODERATE		
W-Cd	8.57	MODERATE		9.66	MODERATE		
A-W	0	SLIGHT		50.84	MODERATE		
A-W-Cd	2.86	SLIGHT		12.47	MODERATE		