Characterization of Pacific oyster (Crassostrea gigas) proteomic response to

natural environmental differences

- 3 Running head: Oyster proteomic response to environment
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Abstract

- 14 Global climate change is rapidly altering coastal marine ecosystems important for food
- production. A comprehensive understanding of how organisms will respond to these
- 16 complex environmental changes can come only from observing and studying species
- within their natural environment. To this end, the effects of environmental drivers pH,
- dissolved oxygen content, salinity, and temperature on Crassostrea gigas physiology
- were evaluated in an outplant experiment. Sibling juvenile oysters were outplanted to
- 20 eelgrass and unvegetated habitat at five different estuarine sites within the Acidification
- 21 Nearshore Monitoring Network in Washington State, USA to evaluate how regional
- 22 environmental drivers influenced molecular physiology. We tested effects environmental
- 23 conditions at outplant sites and habitat to determine if macrophyte presence and diurnal
- cycling buffered pH conditions and changed the oysters' expressed proteome. A novel,
- 21 Systing Bunstou pri conditions and sharinged the System expressed protectine. At here
- 25 two-step, gel-free proteomic approach was used to identify differences in protein
- abundance in *C. gigas* ctenidia tissue after a 29 day outplant by 1) identifying proteins in
- 27 a data independent acquisition survey step and 2) comparing relative quantities of
- 28 targeted environmental response proteins using selected reaction monitoring. While
- 29 there was no difference in protein abundance detected between habitats or within Puget
- 30 Sound, C. gigas outplanted at Willapa Bay had significantly higher abundances of
- antioxidant enzymes and carbohydrate metabolism proteins. Environmental factors at
- 32 Willapa Bay, such as higher average temperature, could have driven this protein
- 33 abundance pattern. These findings generate a suite of new hypotheses for lab and field
- 34 experiments to compare the effects of regional conditions on physiological responses of
- 35 marine invertebrates.

- 37 Keywords: proteomics, oysters, in situ measurements, estuarine systems, antioxidant
- 38 enzymes, carbohydrate metabolism

Introduction

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Global climate change will influence estuarine dynamics and impact the organisms that inhabit these environments. Estuaries are already variable across spatial and temporal scales in terms of phytoplankton production (Pennock and Sharp 1986), nutrient availability, (Paerl et al. 2014), heavy metal contamination (Liu et al. 2015), salinity (Banas et al. 2004), and carbonate chemistry (Pelletier et al. 2018; Feely et al. 2010; Ruesink et al. 2015; Baumann and Smith 2018). Since climate change will affect these parameters, it is important to consider how estuarine organisms will respond. Proteomics, or the study of protein abundance and expression, can be used to shed light on physiological changes on a molecular level. Proteins direct all major cellular functions, thus examining protein abundance provides direct evidence of an organism's physiological response to the estuarine environment (Tomanek 2014). The proteome is dynamic, as it must rapidly respond to perturbation, providing mechanistic information that standard gene expression and mRNA quantification studies cannot (Veldhoen, Ikonomou, and Helbing 2012; Flores-Nunes, Gomes, et al. 2015). As a result of the proteome's dynamic nature, proteins analyzed at the time of collection represent an organism's response to the environment in near real-time. Discoverybased proteomic methods can unbiasedly elucidate responses to environmental drivers (Flores-Nunes, Gomes, et al. 2015). Several studies have connected protein abundances with changes in laboratory-simulated environmental conditions, identifying key proteins and mechanisms involved in specific environmental responses (Timmins-Schiffman et al. 2014; Tomanek 2014; Papakostas et al. 2012; Meng et al. 2017; Parker et al. 2011; Dineshram et al. 2016). While these studies provide insight into organismal

adaptation and physiology, laboratory studies alone cannot fully encapsulate the effects of multiple environmental drivers within an ecosystem context (Riebesell and Gattuso 2014).

Although challenging, *in situ* field studies provide a necessary biological realism when considering variable environments (Cornwall and Hurd 2016; Slattery et al. 2012). Such experiments can be leveraged to study the effects of multiple environmental drivers on organismal physiology and to incorporate realistic variability, as opposed to examining the effect of a single stressor on an organism (Riebesell and Gattuso 2014). Through transcriptomics Chapman et al. (2011) demonstrated the power of an *in situ* experimental design for examining the impacts of regional environmental conditions on Eastern oyster (*Crassostrea virginica*) physiology. Transcript signatures from *C. virginica* sampled from various locations in southeastern United States revealed that temperature, pH, salinity, dissolved oxygen and pollutant load at each location impacted gene expression. Furthermore, they were able to disentangle the interactions of these environmental factors on gene expression. RNA and protein abundances can be influenced by several environmental factors, and *in situ* studies can determine which drivers will be more important to consider for organismal physiology.

Marine invertebrates have proven to be informative bioindicators in proteomic studies to examine the effects of *in situ* conditions on organismal physiological responses to environmental change. When marine invertebrates have been exposed to varying environmental conditions, proteomics has demonstrated changes in cellular defense, immune responses, and genome function (Veldhoen, Ikonomou, and Helbing 2012). Changes in protein abundance in bivalves like blue mussels (*Mytilus edulis spp.*)

and the Pacific oyster (*Crassostrea gigas*) have been used to develop biomarkers for environmental contaminants (Beyer et al. 2017; Slattery et al. 2012). Proteomic responses to natural environmental drivers have also been evaluated in bivalves. For example, shotgun proteomic analysis of *M. edulis* ctenidia from Baltic Sea microcosms revealed that low salinity conditions impact abundance of cytoskeleton proteins and signalling and intracellular membrane trafficking pathways (Campos et al. 2016). With a growing wealth of genomic information for these species, and the importance of this information for commercial aquaculture, understanding how these species fare under different environmental conditions is critical.

Pacific oyster (*Crassostrea gigas*) rearing in estuarine environments in Washington State, USA provides a perfect system to examine the effect of *in situ* environmental conditions on the expressed proteome. *C. gigas* are extensively farmed in two different estuarine systems that show substantial regional variation: Puget Sound and Willapa Bay. Puget Sound is a complex estuarine system with interconnected subbasins, each with different freshwater inputs, residence times, and stratification levels (Feely et al. 2010; Pelletier et al. 2018; Bianucci et al. 2018). Willapa Bay is a large shallow estuary on the Pacific coast that exchanges approximately half its water volume with the Pacific Ocean at each tide (Banas et al. 2004, 2007). Seasonality and location within Puget Sound dictates temperature, dissolved oxygen, salinity, and pH conditions, while Willapa Bay conditions are influenced by diurnal fluctuations and proximity to either the ocean or rivers draining into the bay (Pelletier et al. 2018; Feely et al. 2010; Banas et al. 2007; Ruesink et al. 2015). Both Puget Sound and Willapa Bay also host eelgrass beds (*Zostera spp.*) that affect environmental conditions, such as oxygen

concentrations, on diurnal time scales and may provide refuge from ocean acidification or rapid change in temperature through photosynthetic activity. Understanding how different aquaculture grow-out locations and habitats will affect the oyster's ability to persist through environmental change is crucial for the industry and the ecosystem.

The purpose of this study was to use proteomic techniques to uncover the impacts of environmental drivers on Pacific oyster physiological outcomes in estuarine environments in Washington State, USA. Naturally existing environmental variation was harnessed by outplanting *C. gigas* in different locations within Puget Sound and Willapa Bay, and habitat effects were taken into consideration by placing oysters in eelgrass and unvegetated habitats. Gel-free proteomic methods were used to examine the effects of outplant conditions on relative quantities of all expressed proteins in a series of *in situ* experiments in order to identify differentially abundant proteins. We predicted that differences in environmental drivers at each outplant location and within outplant habitats would yield unique protein abundance patterns. Oysters at outplant locations with higher temperatures, lower dissolved oxygen content, lower salinity, or lower pH may have higher abundances of proteins related to environmental response. Eelgrass beds were expected to ameliorate stressful conditions, resulting in lower abundances of environmental stress response proteins than oysters in unvegetated habitats.

Methods

Shellfish deployment

Sibling *C. gigas* (average shell length = 27.2 cm) were outplanted for 29 days starting June 19, 2016 at five locations: Case Inlet (CI), Fidalgo Bay (FB), Port Gamble

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Bay (PG), Skokomish River Delta (SK), and Willapa Bay (WB) in Washington State, USA (Table 1; Figure S1). These sites were selected for differences in environmental parameters, as well as for the presence of unvegetated areas and eelgrass beds within each site. All sites were part of the Acidification Nearshore Monitoring Network, a network of sensors placed in various Washington locations to monitor marine chemistry (ANeMoNe; Washington Department of Natural Resources). At each site, fifteen juvenile oysters were placed in bags directly onto the substrate at a tidal height of -1.5 MLLW, both inside and outside of eelgrass beds (n=15 per habitat type), for a total of thirty outplanted oysters per site. Oysters were housed in exclusion cages to prevent predation. Custom-built Durafet-based sensors (Honeywell) were used to monitor pH. Commercially-available MiniDOT loggers (Precision Measurement Engineering) were used to measure dissolved oxygen, and Odyssey loggers were used to measure conductivity. All sensors recorded temperature measurements, and all sensors logged at 10-minute intervals across the outplant period, with the exception of SK, where sensors were installed 21 days into the outplant period. Juvenile oysters remained at each site for a 29-day exposure period. Oyster shell length was measured before and after the outplant period. Because the ctenidia is the primary site where oysters interact with the environment, ctenidia samples were dissected at the end of the outplant and held on dry ice until storage at -80°C (Beyer et al. 2017; Meng et al. 2017; Meistertzheim et al. 2007). Environmental data was treated as follows. Conductivity observations were removed when less than 0, which occurs when the instrument is dry. Remaining observations were converted to salinity measurements using the swSCTp function in

the *oce* package in R (Kelley and Richards 2018; R Core Team 2018), with temperature at 25°C and pressure at 10 dbar. For dissolved oxygen, pH, and salinity datasets, data were removed when collected by probes 1) during low tide or 2) when tidal depth was less than one foot to remove readings where the probes may have been exposed. These values were retained for temperature datasets. Outliers were screened using the Tukey method for temperature, dissolved oxygen, pH, and salinity datasets (Hoaglin, Iglewicz, and Tukey 1986). Uniform outplant tidal heights were checked using *prop.test* in R (R Core Team 2018). R Scripts are available in the supplementary Github repository (S2).

Protein Discovery

To identify select protein targets for characterization across locations and environmental conditions, a subset of tissue samples (2 from each site) were analyzed with data independent acquisition (DIA) mass spectrometry analysis (Table S3).

Protein Quantification

Tissue samples were homogenized in a solution of 50 mM NH₄HCO₃ with 6M Urea (500μl). Tissues were then sonicated 3 times (Fisher Scientific Sonic Dismembrator Model 100) for 10 seconds each and cooled between sonications in a bath of ethanol and dry ice. Protein quantities were measured with the ThermoScientific BCA Protein Assay Kit microplate assay with a limited quantity of sonicated sample (11 μL). The protein concentration was measured via spectroscopy at 540 nm in a Labsystems (Waltham, MA) Multiskan MCC/340 and accompanying Ascent Software

Version 2.6. Protein concentrations were calculated based on a standard curve with

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BSA (Pierce) per manufacturer's instructions. **Protein Digestion** Protein digestion followed the protocol outlined in Timmins-Schiffman et al. (2013). To each sample of 30 µg protein, 1.5 M Tris pH 8.8 buffer (6.6 µL) and 200 mM TCEP (2.5 µL) were added. After solvent additions, each sample's pH was verified to be basic (pH ≥ 8), and placed on a 37°C heating block for one hour. Iodoacetamide (200 mM, 20 µL) was then added to each sample to maximize digestion enzyme access to protein cleavage sites. Samples were covered with aluminum foil to incubate in the dark for 1 hour at room temperature. Afterwards, diothiolthreitol (200 mM, 20 µL) was added and samples were incubated at room temperature for one hour. Lysyl Endopeptidase (Wako Chemicals) was then added to each sample in a 1 µg enzyme:30 µg oyster protein ratio, followed by one hour of incubation at room temperature. Urea was diluted with NH₄HCO₃ (25 mM, 800 μL) and HPLC grade methanol (200 μL). Trypsin (Promega) was added to each sample in a 1 µg trypsin: 30 µg oyster protein ratio for overnight digestion at room temperature. Peptide Isolation After overnight incubation, samples were evaporated to near dryness at 4°C with a speedvac (CentriVap ® Refrigerated Centrifugal Concentrator Model 7310021). Samples were then reconstituted in 100 µL of a 5% Acetonitrile and 0.1%

Trifluoroacetic Acid (Solvent B) to isolate peptides. If samples were not at pH ≤ 2, 10-20 µL aliquots of 10% Formic Acid were added until this pH was achieved.

Before desalting peptide samples, Macrospin C18 columns (The Nest Group) were prepared by adding 200 µL of a 60% Acetonitrile with 0.1% Trifluoroacetic Acid (Solvent A). The columns were spun for three minutes at 2000 rpm, and flow-through liquid from the column was discarded. The spinning and discarding process was completed a total of four times. To wash columns, 200 µL of Solvent B was added to each column. The columns were once again spun for three minutes at 2000 rpm and liquid was discarded afterwards; the solvent addition, spinning, and discarding process was completed a total of three times.

To bind peptides to the columns, digested peptides were added to prepared columns, then the columns were spun at 3000 rpm for three minutes. The filtrate was pipetted back onto the column and spun again at 3000 rpm for three minutes. Solvent B (200 μ L) was added to each column three separate times, then the column was spun for three minutes at 3000 rpm to wash salts off the column.

Peptides were eluted with two additions of 100 μ L Solvent A to each column. Columns were spun at 3000 rpm for three minutes and the peptide fraction (filtrate) was reserved. Samples were placed in a speed vacuum at 4°C until they were nearly dry (approximately two hours) to dry peptides. Peptides were reconstituted with 60 μ L of 3% Acetonitrile + 0.1% Formic Acid, and stored at -80°C.

Internal Standard Addition

Peptide Retention Time Calibration (PRTC; Pierce) is a mix of known peptides in known quantities used as an internal standard to ensure consistency of peptides detected and measured throughout a mass spectrometry run. The stock solution of PRTC was diluted to 0.2 pmol/μL PRTC using 3% Acetonitrile with 0.1% Formic Acid. In a clean centrifuge tube, 6 μg of oyster protein and 0.376 pmol of PRTC were mixed together. Sample volume was brought up to 15 μL using a 3% acetonitrile and 0.1% formic acid solution. A quality control solution was also prepared (1 μL PRTC + BSA:3 μL 3% Acetonitrile and 0.1% Formic Acid solution).

Data Independent Acquisition Mass Spectrometry

Peptides were analyzed on an Orbitrap Fusion Lumos mass spectrometer (Thermo Scientific) for Data Independent Acquisition Mass Spectrometry (DIA). DIA analyses were completed as a comprehensive, non-random analytical method for detecting all peptide ions present within a sample. The detection of peptides was then leveraged to develop a targeted proteomics assay for quantification (see Selected Reaction Monitoring Assay). A 30 cm analytical column and 3 cm pre-column were packed in-house with 3 μm C18 beads (Dr. Maisch). Samples were run in a randomized order. A blank injection followed each sample, with procedural blanks run at the very end. Every injection was 3 μL, which included 1 μg of oyster protein and 0.0752 pmol of PRTC. Peptides were analyzed in MS1 over the *m/z* range of 450-950 with 12 *m/z* wide windows with 5 *m/z* overlaps (Egertson et al. 2013). MS1 resolution was 60,000 and AGC target was 400.000 with a three second cycle time. The MS2 loop count was set to

20 and MS2 data was collected with a resolution of 15,000 on charge state of 2 with an AGC target of 50,000. No dynamic exclusion was used (Table S4).

Peptide-Centric Proteomic Analyses

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Unknown peptide spectra from mass spectrometry samples were matched with known peptides using Peptide-Centric Analysis in the PECAN software (Ting et al. 2015). Raw mass spectrometry files were converted to mzML files, then demultiplexed using the MSConvert command line interface (S5; Chambers et al. 2012). The C. gigas proteome was digested with in silico tryptic digest using Protein Digestion Simulator (S6). All known peptides from the mzML files were identified in an comparison to the digested C. gigas proteome (S6). The PECAN-generated spectral library (.blib) file was used to detect peptides of interest in raw DIA files in Skyline (MacLean et al. 2010). Skyline identified peptides using chromatogram peak picking, where ions that elute at the same time and mass are detected as a peptide (S7). All PRTC peptides and approximately 100 different oyster proteins and their peptide transitions were manually checked for retention time and peak area ratio consistency to determine a Skyline auto peak picker error rate (24.3% ± 25%, range: 0% to 100%). Proteins had to satisfy 4 criteria to be considered appropriate targets for the study. 1) To select proteins for a targeted MS assay, protein data was first evaluated in Skyline to ensure there was no missing data for any peptide or sample. 2) Peaks with significant interference were also not considered. 3) Proteins needed at least two

peptides with three transitions per peptide to qualify as a potential target for

downstream assays. 4) Proteins with annotated functions related to oxidative stress, hypoxia, heat shock, immune resistance, shell formation growth and cellular maintenance were considered for targets to address hypotheses of response to in situ environmental differences and eelgrass vs. bare habitat. From this list, fifteen proteins (41 peptides and 123 transitions total), with various environmental response and general maintenance functions, were selected as targets (Table 2).

Selected Reaction Monitoring Assay

Following the protein discovery phase (DIA), proteins were isolated as described above from an additional five randomly selected samples per site and habitat combination (for a total of 5 oysters per group) and analyzed with Selected Reaction Monitoring (SRM). Samples were prepared as described for DIA, except tissue samples were homogenized in 100 µL, and peptide samples were evaporated at 25°C after peptide isolation.

Proteins of interest identified from the DIA analysis were used as targets in a SRM assay following the workflow and informatic pipeline of Timmins-Schiffman et al. 2016; 2017. Target peptide transitions were monitored using SRM on a Thermo TSQ Vantage. SRM data were collected during a gradient of 2-60% acetonitrile over 40 minutes. All samples were run in technical duplicates in a randomized order with a 1 µg oyster peptide and 0.0752 pmol PRTC injection. A quality control injection and blank injection were run after every five sample injections, and PRTC peptides were monitored throughout the experiment.

Target Peptide Specificity

To ensure SRM assay specificity to oyster peptides of interest, oyster peptides were diluted in a background matrix of similar complexity (Pacific geoduck — *Panopea generosa* — peptides), then analyzed using the oyster SRM assay. An oyster-specific SRM target would decrease in abundance with a decreasing abundance of oyster peptides in a mixture. Non-specific peptides — more likely to be found in background matrix of similar complexity — or peptides susceptible to interference would not correlate with oyster peptide abundance, and therefore, would be uninformative. Five *C. gigas* samples used for SRM were randomly selected and pooled in equal quantities. A ten-sample oyster:geoduck dilution series was prepared and run using the same methods as other SRM samples (Table S8).

Target Analysis

Raw SRM files, a background C. gigas proteome, and the PECAN spectral library file from DIA were used to create a Skyline document (S9, S10). Correct transition peaks were selected based on predicted retention times from DIA results by comparing the relative retention times between identical PRTC peptides in the DIA and SRM datasets ($R^2 = 0.99431$). Based on peptide specificity analyses, heat shock protein 70 B2 and one constituent peptide of glucose-6-phosphate 1-dehydrogenase were removed from analyses (S11).

Coefficients of variation were calculated between technical replicates for each peptide transition. Any sample missing data for more than 50% of peptide transitions was deemed poor quality for downstream analyses and excluded. Abundance data was normalized using total ion current (TIC) values from the mass spectrometer.

Consistency between technical replicates was verified in remaining samples using a non-metric multidimensional scaling plot (NMDS) with TIC-normalized data and a euclidean dissimilarity matrix, and technical replicates were averaged after verification with the NMDS. Analysis of Variance and post-hoc Tukey's Honest Significant Difference tests were used to determine significant differences in peptide abundance among sites for average abundances. The Benjamini-Hochberg method with a false discovery rate (FDR) of 10% was used to correct for multiple comparisons.

Results

Conditions at Outplant Locations

Over the course of the outplant, Willapa Bay (WB) had the highest average temperatures (Table 3, Figure S12). There were no large differences in mean pH between locations (Table 3). Oysters at Case Inlet spent more time at low tide than those at the other four sites (χ 2 = 25.29, df = 4, p = 4.408e-05).

Data Independent Acquisition Mass Spectrometry

Out of 39,816 predicted proteins in the *C. gigas* FASTA proteome, 9,047 proteins were detected in *C. gigas* across five sites and two habitats using DIA (Skyline auto peak picker error rate $24.3\% \pm 25\%$, range: 0% to 100%). Proteins detected included, but were not limited to, those annotated from processes such as responses to hypoxia and oxidative stress, removal of superoxide radicals, protein folding, muscle organ development and negative regulation of apoptosis.

Selected Reaction Monitoring Assay

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Differential abundance of protein targets was evaluated at the peptide level after combining technical replicates (Figure S16, Figure S17). Proteins were considered differentially abundant if at least one monitored peptide was differentially abundant. There was no significant difference in SRM peptides between unvegetated and eelgrass habitats across sites (Figure S18). Abundance data from both habitats were pooled for downstream analyses comparing protein abundances among sites. Peroxiredoxin-5 (PRX), thioredoxin reductase 3 (TrxR), catalase (EC 1.11.1.6) (CAT), NAD(P) transhydrogenase (NADPt), glucose-6-phosphate 1-dehydrogenase (G6PD), protein phosphatase 1B (PP1), and glycogen phosphorylase (GPH) had differentially abundant peptides between Willapa Bay and the other four locations (Table S19; ANOVA and Tukey's Honest Significant Difference, FDR = 0.1). These proteins are either involved in oxidative stress response or related to carbohydrate metabolism (Table 4). All differentially abundant proteins were detected at higher levels in the WB oysters than in oysters from the other four PS locations (CI, FB, PG, and SR), regardless of protein function (Figure 1). No differences in protein abundance were

detected among the PS sites (Table S19; ANOVA and Tukey's Honest Significant Difference).

Discussion

Among sibling Pacific oysters placed at five sites in Washington State, we found higher abundances of antioxidant enzymes and carbohydrate metabolism proteins in

oysters outplanted at Willapa Bay (WB). Significant differences in protein abundances were detected between WB and all four PS sites; no differences were observed within PS. Outplanted individuals at WB had higher abundances of antioxidant enzymes and proteins involved in carbohydrate metabolism. Increased antioxidant activity and changes in carbohydrate metabolism have been consistently documented as responses to extreme temperatures, hyposalinity, low pH and air exposure (Tomanek 2014; Zhang et al. 2015). Higher average temperatures may have driven differential protein abundances (Table 3, Figures S12).

To our knowledge, this is the first time combined use of DIA and SRM analytical methods have been applied in an ecological context. Previously, both gel-based (Campos et al. 2016; Tomanek 2014; Zhang et al. 2015; Parker et al. 2011) and gel-free (Timmins-Schiffman et al. 2014; Timmins-Schiffman et al. 2017; Campos et al. 2016; Dineshram et al. 2016; Müller et al. 2018) proteomic methods have been used in lab settings to evaluate the effects of environmental conditions on organismal physiology, with gel-based methods having a limited dynamic range. Similar gel-free methods has also been used to examine physiology of other non-model marine species (Kültz et al. 2015; Plumel et al. 2013). Our methods allowed for increased acquisition of proteomic data for a non-model organism. We detected over 9,000 proteins — far more than previous studies have detected. The application of a novel, two step, gel-free proteomic approach broadened the scope of proteins detected in the DIA phase, thus revealing more possibilities for SRM target design. The SRM assay allowed for increased detection when protein abundance was relatively low.

Antioxidant enzymes

Higher antioxidant enzyme abundances can be a direct response to an increase in reactive oxygen species (ROS) that arise from changes in oxygen conditions (Limón-Pacheco and Gonsebatt 2009; Zhang et al. 2015). During electron transport, oysters can produce ROS that induce oxidative stress if not neutralized. (Abele et al. 2007; Limón-Pacheco and Gonsebatt 2009). Peroxiredoxin-5 (PRX), thioredoxin reductase (TrxR), and catalase (CAT) scavenge these ROS and degrade them before they cause cellular harm (Tomanek 2014; Sussarellu et al. 2012; Limón-Pacheco and Gonsebatt 2009; Flores-Nunes, Mattos, et al. 2015; Sussarellu et al. 2010), while NAD(P) transhydrogenase (NADPt) maintains the cellular redox state. Higher abundances of antioxidant enzymes in WB oysters suggest the need for ROS scavenging to acclimatize to local conditions.

The shallow bathymetry of WB may have also contributed to elevated ROS and the increased need for antioxidant enzymes. At high tide, shallow waters would warm much faster, leading to the observed higher temperatures at WB (Table 1). Warmer waters at low tide may prompt oysters to spend more times with their shells closed, producing higher levels of ROS that would need to be neutralized. Lack of oxygen and higher pCO₂ while the oyster is closed would lead to lower hemolymph pH, causing higher levels of ROS. Oysters would respond through increased abundance of antioxidant enzymes (PRX, CAT, TrxR, and NADPt).

ROS are produced in response to many environmental changes, thus biomarkers of ROS scavenging are difficult to link to a single environmental difference in a variable and complex setting. In particular, reduction of ROS species was found to be a common

response to both increased temperatures and aerial exposure in another experiment with *C. gigas* (Zhang et al. 2015). Because ROS mediation is a conserved response to several stressors, it is possible that environmental parameters we did not measure (e.g., contaminants, microbiota abundance, trace metals) could explain the observed variation in antioxidant enzyme abundance. Future work at these locations should take these variables into account.

Carbohydrate metabolism

Higher abundance of carbohydrate metabolism proteins at WB — protein phosphatase 1B (PP1), glucose-6-phosphate 1-dehydrogenase (G6PD), and glycogen phosphorylase (GPH) — indicate increased glycogen catabolism at this location (Table 3). Increased glycogen catabolism (PP1 and GPH) has been reported with respect to environmental drivers (Fuhrmann et al. 2018; Papakostas et al. 2012; Bacca et al. 2005; Serrano et al. 1993; David et al. 2005). For example, enhanced GPH activity was found in *M. galloprovincialis* responding to thermal stress to provide more energy during high temperature acclimation (Serrano et al. 1993). *C. gigas* outplanted at WB could be using the same process to acclimatize to higher average temperatures. Warmer WB temperatures could have also sped up basal metabolism, necessitating igher abundances of PP1 and GPH to catabolize glycogen stores at a faster rate (Wu et al. 1998; She et al. 2015; Bacca et al. 2005). Increasing the amount of energy available to bivalves by catabolizing glycogen could explain higher abundance of PP1 and GPH in WB oysters.

Increased glycogen catabolism at WB may also be explained by decreased rates of food capture due to less time spent filter feeding or less food availability at this

location. With closed shells, *C. gigas* are unable to filter-feed and would need to catabolize glycogen for energy, facilitated by increased abundances of PP1 and GPH. The WB oysters also had lower total fatty acid content when compared to oysters outplanted at PS locations, indicative of less access to food (less food availability or less time with valves open for filtration) or poorer quality food (Lowe et al. *in prep*). Whether due to more time spent with their shells closed or decreased caloric intake, the WB oysters had to rely on energy stores of glycogen more than oysters throughout Puget Sound.

Higher G6PD abundance in WB oysters indicate that oysters at WB were still metabolically active. G6PD catalyzes the oxidative portion of the pentose phosphate pathway, and products from this pathway are often precursors for nucleic and aromatic amino acids ((L., n.d.)). Additionally, G6PD activity generates NADPH and can indirectly ameliorate effects of ROS ((L., n.d.)), as demonstrated in *C. gigas*: after exposure to linear alklybenzenes and other pollutants, *C. gigas* ctenidia expressed higher G6PD activity to regulate the redox environment (Flores-Nunes, Mattos, et al. 2015). Increased abundance of G6PD in WB not only could have maintained transcription and translation processes, but also levels of cellular and metabolic activity by indirectly dealing with ROS. Increased G6PD abundance, therefore, provides a link between carbohydrate metabolism and oxidative stress response in oysters outplanted at Willapa Bay.

Proteomic Responses in Puget Sound and Between Habitats

Due to the observed differences across environmental parameters between Willapa Bay and Puget Sound locations, similar abundances for proteins involved in various environmental responses may be evidence of physiological plasticity. One

particular protein that we expected to be differentially abundant was heat shock 70 kDA protein 12A (HSP 12A), since Willapa Bay had higher average temperatures (Table 3). Higher abundances of heat shock proteins (HSPs) are generally induced when organisms are exposed to thermal stress (Hamdoun, Cheney, and Cherr 2003; Zhang et al. 2015; Meistertzheim et al. 2007). In our experiment, average temperatures *C. gigas* experienced were lower than the threshold to induce elevated HSP expression (Figure 1; Table 2) (Hamdoun, Cheney, and Cherr 2003). While HSP12A is involved in a general response to environmental drivers, differentially expressed antioxidant enzymes and carbohydrate metabolism proteins could be more indicative of molecular processes required for acclimatization.

The lack of differential abundance for protein targets — both among Puget

Sound sites and between unvegetated and eelgrass habitats — was unexpected. These similar proteomic profiles could be due to three factors. First, it is possible that a different suite of environmental response proteins in the SRM assay could have yielded a different view of acclimatization to the various outplant sites; however, the targets we chose have proven to yield insight into a range of environmental responses in previous studies. Second, outplant duration could have been too short to capture varied physiological response within Puget Sound, or oysters could have also acclimatized to conditions in their outplant locations Finally, is that it is possible that the proteomic response was not different because the environmental conditions that would elicit up- or down-regulation of monitored proteins were similar across these five locations (Table 3). While eelgrass beds are touted as a potential refuge for organisms from the negative effects of environmental drivers like pH, they can also drive more extreme carbonate

chemistry conditions (Pacella et al. 2018). It is possible that the refuge effects and extreme effects cancelled each other out over the course of our outplant, effectively making the eelgrass beds just like the unvegetated sites for outplanted oysters. If this is the case, it would explain why there was no detected difference in protein abundance between unvegetated and eelgrass habitats.

Conclusion

The differential protein abundance observed between sibling oysters placed for 29 days in Willapa Bay or Puget Sound indicates that environmental factors at Willapa Bay lend themselves to increased antioxidant enzyme and carbohydrate metabolism protein abundance. Because the observed proteomic responses can be triggered by a variety of drivers, from a field trial it is difficult to discern if high temperatures or other environmental factors at WB led to these responses. However, the observed responses generate a suite of new hypotheses for lab and field experiments comparing the effects of environmental conditions on physiological responses of marine invertebrates. This is the first to link regional environmental conditions to physiological responses in Puget Sound, and one of the first to compare responses between Puget Sound and Willapa Bay. Linking environmental drivers to other performance metrics, such as growth and fatty acid composition, is crucial for contextualizing these protein abundance findings (Lowe et al. *in prep*).

Understanding the difference between these two estuaries is important for the persistence of oyster reefs and aquaculture in the face of climate change. Since aquaculture species are outplanted in the same location for many months to years, the

lack of differential protein abundance among Puget Sound outplants could mean that environmental response should not be heavily considered when selecting grow-out locations in this region. Future experiments with a longer outplant duration and multiple sampling points may provide a clearer connection between environmental drivers, protein expression, and performance, as well as assist with developing a practical assay for elucidating stress responses in the field.

Figures and Tables

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Table 1. Latitude and longitude of *C. gigas* outplants. Oysters were placed at five locations sites: Case Inlet (CI), Fidalgo Bay (FB), Port Gamble Bay (PG), Skokomish River Delta (SK), and Willapa Bay (WB).

Location	Latitude	Longitude
CI	47.3579367	-122.7957627
FB	48.481691	-122.58353
PG	47.842676	-122.583832
SK	47.35523	-123.1572
WB	46.4944789	-124.0261356

Table 2. Proteins used as targets for a Selected Reaction Monitoring Assay (SRM).

Targets were identified based on differential abundance and stress-related annotations.

At least two peptides and six transitions were included in the assy for each protein. The

protein Catalase had two isoforms under separate proteome IDs —

CHOYP_CATA.1.3|m.11120 and CHOYP_CATA.3.3|m.21642 — and target peptides

were chosen from both IDs. A total of four peptides and twelve associated transitions

were used as SRM targets.

Protein	Proteome ID	Number of Peptides	Number of Transitions
3-ketoacyl-CoA thiolase	CHOYP_ACAA2. 1.1 m.30666	3	9

Peroxiredoxin-5	CHOYP_BRAFLD RAFT_119799.1. 1 m.23765	2	6
Thioredoxin reductase 3	CHOYP_BRAFLD RAFT_122807.1. 1 m.3729	3	9
Protein phosphatase 1B	CHOYP_BRAFLD RAFT_275870.1. 1 m.12895	3	9
Carbonic anyhdrase 2	CHOYP_CAH2.1. 1 m.42306	3	9
Catalase 1	CHOYP_CATA.1. 3 m.11120	3	9
Catalase 2	CHOYP_CATA.3. 3 m.21642	1	3
Glucose-6-phosphate 1-dehydrogenase	CHOYP_G6PD.2. 2 m.46923	3	9
Heat shock 70 kDa protein	CHOYP_HS12A. 25.33 m.60352	2	6

Heat shock protein 70	CHOYP_HSP74.	2	6
NAD(P) transhydrogenase	CHOYP_LOC100 633041.1.1 m.354 28	2	6
Glycogen phosphorylase	CHOYP_LOC100 883864.1.1 m.417 91	3	9
Multidrug resistance- associated protein	CHOYP_MRP1.5. 10 m.34368	2	6
Protein disulfide- isomerase	CHOYP_PDIA1.1 .1 m.5297	3	9
Protein disulfide- isomerase	CHOYP_PDIA3.1 .1 m.60223	3	9
Puromysin-sensitive amirase	CHOYP_PSA.1.1 m.27259	3	9

Table 3. Environmental conditions (mean ± standard deviation) at outplant locations (Case Inlet (CI), Fidalgo Bay (FB), Port Gamble Bay (PG), Skokomish River Delta (SK), Willapa Bay (WB).

Metric	CI	FB	PG	SK	WB
Mean	16.1 ± 1.7	14.8 ± 1.8	15.1 ± 2.9	15.1± 2.2	18.0 ± 1.3
Temperature					
(°C)					
Mean Salinity	24.47 ±	30.20 ±	23.25 ± 1.56	13.43 ± 1.04	27.28 ±
(PSU)	1.35	0.38			0.73
Mean	8.27 ± 1.91	9.63 ± 4.59	11.70 ± 3.70	9.99 ± 4.58	8.76 ± 1.78
Dissolved					
Oxygen					
Content					
(mg/L)					
Mean pH	7.63 ± 0.19	7.54 ± 0.23	7.33 ± 0.25	7.37 ± 0.24	7.56 ± 0.18

Table 4. Functions of differentially abundant proteins

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Protein	Function	Citations
Peroxiredoxin-5 (PRX)	Scavenges ROS located in	Tomanek 2014; Sussarellu
	mitochondria using	et al. 2012
	cysteine residuals to	
	reduce substrates	
Thioredoxin reductase	Thiol-specific antioxidant	Limón-Pacheco and

(TrxR)	reducing agent	Gonsebatt 2009	
Catalase (CAT)	Degrades hydrogen	Flores-Nunes, Mattos, et	
	peroxide into water and	al. 2015; Sussarellu et al.	
	oxygen	2012	
NAD(P) transhydrogenase	Involved in maintenance of	Sussarellu et al. 2010,	
(NADPt)	cellular redox state	2012	
Glucose-6-phosphate 1-	Catalyzes first step in the	Livingstone., 1982	
dehydrogenase (G6PD)	pentose phosphate		
	pathway and controls rate		
	of pathway's oxidative		
	portion		
Protein phosphatase 1B	Regulation of glygcogen	Wu et al. 1998	
(PP1)	metabolism by		
	dephosphorylating		
	glycogen synthase		
Glycogen phosphorylase	Key glycogen catabolism	David et al. 2005; Oyster	
(GPH)	enzyme. Removes glucose	and Gigas 1993; She et al.	
	from glycogen molecules	2015	
	to aid in glycogen		
	breakdown		

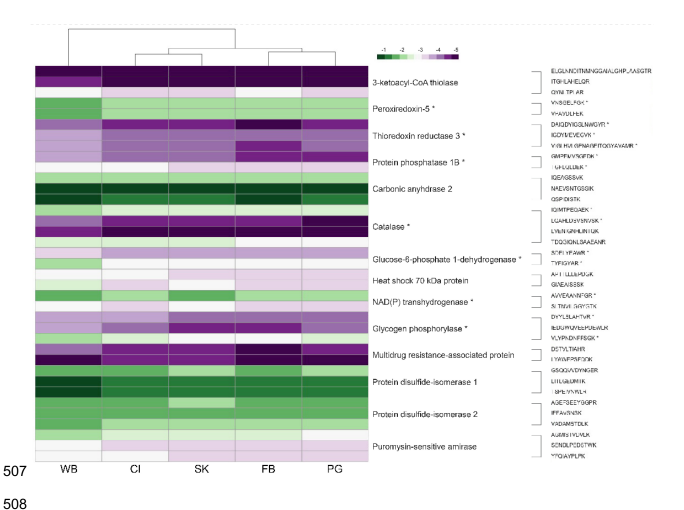


Figure 1. Average protein abundance by constituent peptides across experimental sites from SRM. Peptide abundance data at each site was averaged, then log transformed. Proteins were considered differentially abundant if at least one constituent peptide was significantly different (indicated by an asterisk). There were no significant differences in protein abundance among the Puget Sound locations, or between unvegetated and eelgrass habitats.

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