Transcriptome of the Eastern Oyster Crassostrea Virginica in Response to Bacterial Challenge

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TRANSCRIPTOME OF THE EASTERN OYSTER CRASSOSTREA VIRGINICA IN RESPONSE TO BACTERIAL CHALLENGE

BY

IAN MCDOWELL

A THESIS SUBMITTED IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF

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ABSTRACT

The Eastern oyster *Crassostrea virginica*, an ecologically and economically important estuarine organism, suffers mortalities as high as 90-100% in affected areas due to Roseovarius Oyster Disease (ROD), caused by the bacterial pathogen, *Roseovarius crassostreae*. Advanced genotypic breeding techniques necessitate information regarding markers and genes associated with disease resistance. As yet, the host-pathogen interaction between *C. virginica* and *R. crassostreae* is poorly understood at the molecular level. The identification of potential genes and pathways responsible for an effective host defense response in the Eastern oyster to *R. crassostreae* is important not only to provide a basis for enhanced breeding techniques, but also to enhance understanding of innate immunity in a broader, evolutionary sense. The present study proposed to uncover not only genes and general processes potentially involved in disease resistance to ROD in the Eastern oyster, but also diversified gene families. To that end the present study entailed a disease challenge exposing ROD-resistant and ROD-susceptible families of oysters to *R. crassostreae*, high-throughput cDNA sequencing of samples from several timepoints throughout the disease challenge, assembly of sequence data into a reference transcriptome, analysis of the transcriptome through differential gene expression and gene family similarity clustering, and single nucleotide polymorphism (SNP) detection to identify candidate gene markers. Oyster resistance to *R. crassostreae* was found to involve extracellular matrix
remodeling, cell adhesion, inflammation, metabolism, and other processes. Several gene families identified as putatively diversified and important in the oyster host defense response were enumerated and described, including serine proteases, serine protease inhibitors, c-type lectins, C1q domain-containing proteins, fibrinogen domain-containing proteins, scavenger receptors with class B SRCR domains, interferon-induced protein 44 (IFI44) family proteins, and GTPase of the immunity associated protein (GIMAP) family proteins. Further, similarity clustering of proteins and translated transcripts from diverse invertebrates suggested that GIMAP proteins are expanded in molluscs and IFI44 proteins are expanded in bivalves.
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INTRODUCTION

The Eastern oyster *Crassostrea virginica* is an ecologically and economically important estuarine organism cultured from Louisiana, USA to New Brunswick, Canada (Kennedy et al. 1996). Oyster production is an important sector of United States agriculture and the Eastern oyster was estimated in 2010 to have a farm gate value of $90.4 million in the United States (NMFS 2010). Several oyster diseases, both protozoan and bacterial, have expanded in range and increased in severity during the latter half of the twentieth century, often causing staggering losses (Cook et al. 1998; Ford and Chintala 1996; Ford and Smolowitz 2007). Roseovarius Oyster Disease (ROD), caused by the bacterium *Roseovarius crassostreae*, went unreported before 1988 and presently affects oysters from the Long Island Sound north to Maine (Bricelj et al. 1992; Ford and Borrero 2001; Boettcher et al. 2005; Markey and Gomez-Chiarri 2009). ROD can cause mortalities as high as 90-100% in affected areas (Bricelj et al. 1992; Ford and Borrero 2001; Boettcher et al. 2005). ROD can cause mortality events that last a few weeks, often coinciding with or following closely upon peak summer temperatures. Gross clinical signs include uneven shell margins, soft tissue emaciation, and conchiolin depositions on the inner shell surfaces (Bricelj et al. 1992; Barber et al. 1998; Boettcher et al. 1999; Ford and Borrero 2001).

The host-pathogen interaction between *C. virginica* and *R. crassostreae* is poorly understood. It is known that *R. crassostreae* colonizes the oyster's
inner shell surface before lesions develop in the epithelial mantle (Boardman et al. 2008). Shell colonization may enable *R. crassostreae* to evade cell-mediated killing by hemocytes (Boardman et al. 2008), which are the active cells of the immune system present in the hemolymph and involved in triggering and sustaining both cell-mediated and humoral host defense, wound and shell repair, and other processes (Ford and Tripp 1996). Colonization likely stimulates the oyster to deposit conchiolin and it has been suggested that smaller oysters succumb to ROD because they lack adequate metabolic resources to fuel deposition, leading to emaciation (Bricelj et al. 1992; Ford and Borrero 2001; Boardman et al. 2008). *R. crassostreae* may produce a toxin with ciliostatic activity (Boettcher et al. 2000) and extracellular products from *R. crassostreae* have a cytotoxic effect on oyster hemocytes that cannot be solely attributed to lipopolysaccharide (LPS), a component of the membrane of gram-negative bacteria (Gomez-Leon et al. 2008).

At least two oyster lines with resistance to ROD are currently available, including the Frank M. Flowers (FMF) line and the University of Maine Flowers Select (UMFS) line, the latter of which contains germline materials from the former (Barber et al. 1998, Davis and Barber 1999; Lewis 2001). While traditional breeding practices have led to the production of ROD-resistant oysters, the genetic basis of resistance is presently unknown. Advanced genotypic breeding techniques, which necessitate information regarding markers and genes associated with disease resistance, have multiple
advantages over the current, traditional phenotypic breeding techniques including increased accuracy in estimating breeding values; amenability to disease resistance as a breeding trait, which only becomes apparent at death and may wane under traditional breeding regimes without consistent disease pressure; and enablement of simultaneous culture of mixed pedigrees whereby environmental effects can be minimized, which have been shown to be of great effect in oyster culture and may seriously confound main effects in traditional family-based selection (Langdon et al. 2003; Guo et al. 2008; Massault et al. 2008; Cancela et al. 2010).

The identification of potential genes and pathways responsible for an effective host defense response in the Eastern oyster to *R. crassostreae* is important not only to provide a basis for enhanced breeding techniques, but also to enhance understanding of innate immunity in a broader, evolutionary sense. Deep sequencing of the Eastern oyster transcriptome in response to bacterial challenge is a valuable and interesting contribution because the Eastern oyster is a member of Lophotrochozoa, a superphylum that has been poorly represented among genomic and transcriptomic datasets until the recent release and publication of the Pacific oyster, *Crassostrea gigas*, genome (Zhang et al. 2012). Research conducted on the innate immunity of model invertebrates with sequenced genomes has focused on deuterostomes like the purple sea urchin *Strongylocentrotus purpuratus*, the tunicate *Ciona intestinalis*, and the amphioxus *Branchiostoma floridae*, while research into the
The innate immunity of protostomes has disproportionately focused on members of Pancrustacea and, more specifically, Arthropoda, including *Drosophila* spp. (Azumi et al. 2003; Kim and Kim 2005; Hibino et al. 2006; Huang et al. 2011). Invertebrate hosts lack the classical adaptive immune mechanism of receptor diversification through RAG-1- and RAG-2-mediated somatic recombination and gene conversion, yet they successfully combat widely varied types of microbes and parasites, which have comparatively high rates of mutation and rapid generation times (Flajnik and Du Pasquier 2004; Du Pasquier 2005). To mount effective and flexible defense responses to diverse pathogens, invertebrate hosts have developed diversified repertoires of receptors, regulators, and/or effectors (Messier-Solek et al. 2010). Though the precise role, relative importance, and presence/absence of specific genes and gene families relevant to host defense differs among the invertebrates in the taxa referred to above, several general strategies of diversification have been described including allelic diversity/SNPs/indels, alternative splicing, and gene family expansion (Ghosh et al. 2010; Messier-Solek et al. 2010). As an example of the latter, the genome of sea urchin *S. purpuratus* has undergone extensive expansions of Toll-like receptors (TLRs), NACHT- and leucine-rich repeat-containing (NLR) proteins and multi-domain scavenger receptors cysteine-rich (SRs), all of which directly or indirectly bind to pathogen associated molecular patterns (PAMPs) (Hibino et al. 2006). The *Dscam* gene, as studied in several arthropods, illustrates the strategy of diversification.
through alternative splicing. The Dscam intron-exon architecture enables a possible production of tens of thousands of protein isoforms that function in homophilic binding and, likely, heterophilic, bacterial binding (Watson et al. 2005). Fibrinogen-related proteins (FREPs) identified in arthropods and molluscs have been shown to participate in agglutination and phagocytosis and have attained diversity through the multiple strategies of gene family expansion, alternative splicing, allelic diversity, and even somatic recombination (Leonard et al. 2001; Zhang et al. 2004; Ghosh et al. 2010).

While high-throughput, digital transcriptomic studies of host defense have been performed in molluscs other than the Eastern oyster, including the Pacific oyster C. gigas, the mussel Mytilus edulis, the Manila clam Ruditapes philipinarum, and others (de Lorgeril et al. 2011; Brulle et al. 2012; Philipp et al. 2012), disease challenge transcriptomic studies specific to Eastern oyster host defense have used medium-throughput approaches including expressed sequences tag (EST) analysis and microarrays (e.g. Jenny et al. 2002; Wang et al. 2010). The present study proposed to uncover genes, gene families, and general processes potentially responsible for disease resistance to ROD in the Eastern oyster. Sequences of cDNA from ROD-resistant and susceptible families of oysters exposed to R. crassostreae were assembled into a reference transcriptome. Differential gene expression and gene family analyses were used to identify genes, gene families, and general processes involved in oyster immunity, and single nucleotide polymorphism (SNP) were
identified in candidate genes. The present study found that genes involved in extracellular matrix (ECM) restructuring, cell adhesion, inflammation, metabolism, catecholamine signaling, and several other processes distinguished resistance from susceptibility to Roseovarius Oyster Disease. The present study also identified several large gene families important in Eastern oyster host defense including two families poorly characterized in invertebrates, the GTPase of the immunity associated protein (GIMAP) family and the interferon-induced protein 44 (IFI44) family, which appear to be expanded in molluscs and bivalves, respectively.
METHODS

Bacterial Challenge of Eastern Oysters

Juvenile Eastern oysters from 2 families with known differential susceptibility to ROD as determined in a previous study were provided by the Rutgers University shellfish hatchery (Guo and Gomez-Chiarri, in preparation). F3L oysters were F3 generation progeny derived from a single pair mating of a female oyster from Rutgers NEH (Northeastern High-survival) line and a male oyster from Louisiana (LA). GX09 (henceforth GX) oysters were progeny of two full-sib families of F3 generation derived from three full-sib families whose parents contained germline material from the following stocks/lines: Rutgers NEH and DBH (Delaware Bay High-survival line), LA, University of Maine Flower’s Select (UMFS), and Frank M. Flower’s (FMF). Oysters from the F3L families had an admixture of four possible haplotypes, while GX oysters had an admixture of eight possible haplotypes. Two hundred seventy-three juvenile oysters (shell height, 10-15 mm) each from F3L and GX were divided into two replicate tanks with filtered sterile seawater (FSSW) for bacterial challenge (designated F3L and GX). Three groups of additional oysters (2 x 50 GX and 1 x 50 F3L) were kept in separate tanks as unchallenged controls (designated CGX and CF3L, respectively). Oysters were acclimated during a period of 2 weeks from conditions at origin to experimental conditions (salinity 28-30‰, temperature 15-19°C). Oysters in the challenge tanks were exposed
by bath to *R. crassostreae*, strain CV919-312<sup>T</sup> (Boettcher et al. 2005) at a final tank concentration of 7.5 × 10<sup>6</sup> colony forming units (CFU) ml<sup>−1</sup> (day 0 of challenge), while oysters in control tanks were not exposed. Oysters were fed Instant Algae (Reed Mariculture) every other day and water was partially changed (50%) weekly. Oysters were monitored weekly for 93 days for mortalities and for the presence of clinical signs of ROD (uneven valves and conchiolin deposits in shells of dead oysters). Infection by *R. crassostreae* was confirmed by PCR (Maloy et al. 2005).

**Sample collection, cDNA preparation, and sequencing**

Oyster whole body tissue was collected from 5 randomly sampled oysters each from CGX, GX, and F3L at days 1, 5, 15, and 30 following challenge and stored in Qiagen RNAlater® RNA Stabilization Reagent until time of RNA isolation. All RNA molecules >200 nucleotides were purified using Qiagen RNAeasy Mini Kit. Samples were checked for RNA purity using the Nanodrop 8000 spectrophotometer and random samples were checked using the Agilent 2100 Bioanalyzer. Total RNA from whole body tissue from 5 oysters per time per treatment was pooled for a total of 12 experimental samples (3 treatments x 4 time points). Pools of RNA samples for each of the experimental groups were selectively enriched for poly-A containing mRNA using the Illumina mRNA-Seq-8 Sample Prep Kit which involved poly-A capture by poly-T oligo-attached magnetic beads, fragmentation, cDNA
synthesis, adapter ligation, and purification and enrichment by PCR. The resultant cDNA libraries were sequenced on the Illumina GAIIx platform (one sample/5 individuals per lane, 1 lane per sample for a total of 12 lanes) (Genome Quebec, Canada).

Read processing and de novo assembly

Raw sequencing reads of 108 bp were pooled then processed and filtered for contamination of mitochondrial and ribosomal sequences by mapping to all Crassostrea spp. rRNA and mtDNA in NCBI Genbank database, and were filtered for vector sequences by mapping to Univec using bowtie2 (Langmead and Salzberg 2012; ftp://ftp.ncbi.nih.gov/pub/univec). The 5’-end of reads was trimmed to reduce GC-bias (Hansen et al. 2010). Using the btrim software package, Illumina adapters were trimmed, low-complexity artifacts were removed, and reads were further trimmed using adaptive quality trimming (Kong 2011). Reads less than 20 bp in length at this stage were discarded. Processed transcriptome reads were assembled using Trinity (release 20111126) with default options (Grabherr et al. 2011). Only those assembled contigs ≥ 200 bp were retained.

Contamination Removal

Transcriptome contigs were compared to the RefSeq protein database (Sayers et al. 2012). A custom python script, created by L. Dong (Brown...
University), was used to parse BLAST output and identify possible contaminants. Contigs that had all top blast hits (a maximum of 10) with associated e-value ≤ 1e-06 to proteins from Archaeabacteria, Bacteria, or Protozoa were discarded. Additional mitochondrial and ribosomal contaminants were identified and discarded during prot4EST in silico translation (see below) and through text searching of BLAST results. Phage integrase sequences were identified and discarded by comparing the transcriptome to Pfam_A using Pfam scan (version 1.3) and HMMER (version 3.0) with hits retained where e-value ≤ 1e-05 (Finn et al. 2011). DNA transposons were identified and discarded using RepeatMasker (Smit et al. 2010).

**Differential Gene Expression**

Reads from individual treatments were aligned to the reference transcriptome using bowtie with parameters, "-v 3 --a --best --strata," such that 3 mismatches were allowed per read to account for the high rate of polymorphism in oysters, while only reporting the alignment with the least number of mismatches for each “stratum” (Langmead et al. 2009). Transcript abundances in reads per kilobase per million reads mapped (RPKM) were estimated using RSEM (RNA-Seq by Expectation Maximization) through the Trinity plug-in, run_RSEM.pl. (Mortazavi et al. 2008; Grabherr et al. 2011; Li and Dewey 2011). To reduce bias from differential sequencing depth across
lanes, the trimmed mean of M values (TMM) method was used to calculate normalization factors for each lane by which read abundances were multiplied and adjusted (Robinson and Oshlack 2010). Read abundances in the control (non-challenged CGX) samples from days 15 and 30 were considered jointly as a control pool to increase the accuracy of estimating the biological variability between samples and increase power in identifying DEGs. Read abundances from control samples (CGX 1 and 5 d) were not included in this pool because of a mortality event that occurred between days 1 and 7 for CGX (Fig. 1). Read abundances for contigs in each of the samples were compared to read abundances in the control pool (CGX 15 and 30 d). Only those contigs with at least 1 count-per-million in at least 2 samples were tested for differential expression. Differential gene expression (DGE) testing was performed using edgeR, which assumes a negative binomial distribution and uses an empirical Bayes estimation and exact test to identify differentially expressed genes (DEGs) (Robinson et al. 2010). Significance values yielded by hypergeometric test were adjusted using the False Discovery Rate (FDR) correction and a contig was considered differentially expressed (DE) if it had an FDR-adjusted p-value ≤ 0.05 (Benjamini and Hochberg 1995; Robinson et al. 2010). DGE testing was performed on all contigs (“gene isoform” testing) and on read abundances summed for all contigs within each Trinity component (“gene model” testing) (Grabherr et al. 2011). Only those DEGs that were significant at the level of “gene isoforms” and belonged to
components that were significant at the level of “gene models” were considered differentially expressed. While the relationship of Trinity contig to Trinity component works on the level of graph space and is not precisely biological (Grabherr et al. 2011), the relationship as it was applied only added a stringent filter to the pool of DEGs and did not contribute new DEGs.

Heatmap Analysis

For all genes differentially expressed in any one of the treatments GX-d1, GX-d5, F3L-d1, F3L-d5, F3L-d15, and F3L-d30, the log$_2$-transformed RPKM for each gene in each of eight treatment-days (including control, CGX-d15 and CGX-d30) were Z-score centered by subtracting then dividing by the mean log$_2$-transformed RPKM by gene. Genes were hierarchically clustered using Euclidean distance and complete linkage of the Z-score-transformed gene expression. Treatment-days were clustered using the complete linkage Euclidean distance of the Spearman correlation of the Z-score-transformed gene expression. Clustering and visualization were performed using the fastcluster and gplots packages, respectively, in the R programming environment (Bolker et al. 2010).

Annotation and Functional Enrichment

Transcriptome contigs were compared to the NCBI protein non-redundant database and Uniref100 using BLASTX (Altschul et al. 1997). Hits
were retained with e-value ≤ 1e-6. For all subsequent methods herein employed, the annotation to NR was used because a greater percentage of the transcriptome could be annotated using this database. Gene Ontology (GO) terms were mapped to best BLASTX hits for each contig using the Blast2GO pipeline (version 2.3.5) (Conesa et al. 2005). Functional enrichment for each treatment comparison (GX_early, F3L_early, and F3L_late, where early includes days 1 and 5 after challenge, and late days 15 and 30) was performed by comparing the numbers of gene ontology (GO) terms associated with annotations of DEGs to the numbers of terms associated with all transcripts not DE using the R package topGO, which also accounts for the hierarchical topology of the GO graph (Alexa et al. 2006). The topGO “elim” algorithm eliminates significant child nodes from tests of significance for parent nodes, as significant child nodes may otherwise confer significance to their parent nodes. Fisher’s exact test was used to determine significance of enrichment of each GO term, with Bonferroni-adjusted p-values ≤ 0.05 taken as significant. Significantly functionally enriched GO terms were visualized in semantic space using SimRel functional similarity measure (Schlicker et al. 2006), REViGO online visualization tool (Supek et al. 2011), and custom R scripts.

Protein Family Identification and Test for Enrichment
Transcriptome contigs were compared to Pfam_A using PfamScan (version 1.3) and HMMER (version 3.0) and only hits with an e-value ≤ 1e-5 were retained (Eddy 1998; Finn et al. 2011; Pfam_A downloaded July 7, 2012). It is possible for a contig to have multiple hits to the same Pfam profile hidden Markov model (HMM). These hits were made non-redundant by retaining only the most significant contig-to-HMM hit. DEGs for each treatment comparison were tested for enrichment of both Pfam families and Pfam domains. Pearson’s chi-squared test was performed to test for the enrichment of each unique HMM accession. If the expected count for any cell in the 2x2 contingency table was 5 or fewer, Fisher’s exact test was performed instead. Significance p-values were adjusted for the number of independent enrichment tests by the False Discovery Rate correction method and p-values ≤ 0.05 were considered significant (Benjamini and Hochberg 1995).

**Peptide Similarity Clustering**

Prot4EST, a perl script that integrates several programs and approaches to find the best open reading frame, including, in order of priority, BLASTX similarity, codon usage bias, and longest open reading frame (Wasmuth and Blaxter 2004), was used to translate transcriptome contigs into a set of putative peptide sequences. For codon usage bias, all full-length Bivalvia Uniref100 proteins were used as the training set. Translated transcripts were then clustered into groups by similarity using TribeMCL. A
custom perl script provided by I. Misner (University of Rhode Island) was used to prepare the translated transcriptome for all-versus-all BLASTP and to execute this similarity search. Hit, or edges, with e-value $\leq 1e^{-5}$ and hit identity $\geq 20\%$ were retained as significant. Negative log10-transformed e-values above 99 were set to 99 so that subsequent edge weights for MCL clustering were not skewed (Chen et al. 2007). The Markov cluster (MCL) algorithm set on the default inflation index, 1.5, was then used to dissolve less reliable edges that may result from sequence similarity errors or from shared protein domains across families (Enright 2002). MCL returns connected components or clusters of contigs based on protein sequence similarity, which can be defined as a set of nodes such that any two nodes in the same cluster is connected by a path of edges and any two nodes in different clusters is not connected by any path of edges.

Two similarity graphs were constructed: a *C. virginica*-only graph, containing translated contigs from our *de novo* assembly only, and a multi-species sequence similarity graph, constructed using translated *C. virginica* transcripts along with sequences from evolutionarily diverse species including the basal eumetazoan *Nematostella vectensis*; the deuterostome *Strongylocentrotus purpuratus*; the basal chordates *Branchiostoma floridae*, *Saccoglossus kowalevskii*, and *Ciona intestinalis*; the basal vertebrate *Petromyzon marinus*; the pancrustaceans *Drosophila melanogaster* and *Daphnia pulex*; and several molluscs including the gastropods *Lottia gigantea*,
Aplysia californica, Biomphalaria glabrata, and Lymnaea stagnalis, and a bivalve species, the Pacific oyster Crassostrea gigas. Sequences were downloaded from a variety of sources in the form of ESTs and protein sequence (Appendix A, Table S1). ESTs were translated to peptide using the prot4EST pipeline. Full-length Gastropoda Uniref100 proteins were used as the training set for determination of the codon usage bias for the three gastropod species, while full-length Bivalvia Uniref100 proteins were used as the training set for C. gigas ESTs.

Protein sequences and translated ESTs from all organisms were concatenated, formatted, and filtered using the first two steps of the OrthoMCL pipeline, orthomclAdjustFasta and orthomclFilterFasta (Li et al. 2003; Fischer et al. 2011). Proteins were compared in a parallel run of all-versus-all BLASTP, retaining hits with e-value $\leq 1e^{-10}$ (Altschul et al. 1997). The orthomcl utility orthomclBlastParser was used to calculate the percent identity of each hit. The list of significant hits, or edges, was then made non-redundant and with a maximum negative log10-transformed e-value of 99 using custom python scripts. The resultant network file was filtered using MCL set on the default inflation index of 1.5 (Enright et al. 2002).

Enrichment of DEGs among TribeMCL clusters

Each C. virginica-only TribeMCL cluster was independently tested for enrichment of the 3 DEG treatment comparison groups (GX_early, F3L_early,
and F3L_late). Pearson’s chi-squared or Fisher’s test was performed to test for the enrichment of each group of DEGs for each TribeMCL cluster as described above. Pragmatic rules were adopted for annotating clusters. Enriched TribeMCL clusters were annotated with a protein name if > 50% of the contigs had identical or nearly identical BLASTX best hits and the remainder of contigs had no significant hits, or if ≥ 80% of the contigs had identical or nearly identical BLASTX best hits and the remainder of contigs had dissimilar best BLASTX hits (which may be the case in the sharing of domains, e.g., chymotrypsin and neurotrypsin). Enriched TribeMCL clusters were putatively annotated (noted with asterisks) if < 50% half of the contigs had identical or nearly identical BLASTX best hits and the remainder of contigs had no significant hits or had a small number of dissimilar but non-repeating hits (frequency equal to one). Multiple names were used for the annotation if a number of non-identical hits of comparable frequency composed a majority of the cluster (e.g., “hemicentin/rhamnospondin/thrombospondin*”). All other TribeMCL clusters were named “unknown.”

**Selection of Putative Diversified Families**

Gene diversification implies a gene family expansion comparative to other lineages (e.g. by gene duplication) and/or gene diversity significantly greater than that which is present in the genomic sequence alone (e.g. by alternative splicing). To identify “diversified” families with host defense
relevance, working definitions were adopted for the terms: gene family, expansion, and host defense relevance. A gene family was defined as a group of transcripts with similarity between one another (membership in the same TribeMCL cluster) and/or similarity to the same gene family based on BLAST matches (e.g. two transcripts with BLAST similarity to C1qDC proteins). Gene families were considered to be expanded if: 1) families consisted of at least 50 non-redundant transcripts (in order to find the largest families and to significantly reduce the number of gene families under consideration); and 2) the number of non-redundant transcripts in that gene family in *C. virginica* exceeded the number of non-redundant transcripts/proteins in that gene family in at least half of the other 14 organisms considered. The putative diversified gene families were regarded as relevant to the oyster host defense to bacterial challenge if they contained a significant portion of DEGs as determined by Pfam and/or TribeMCL-DEG enrichment.

Certain diversified gene families were selected for further analyses based on the following criteria: 1) high rank of significance in the enrichment of DEGs belonging to the gene family in Pfam and/or TribeMCL-DEG enrichment sets (preferably both); 2) large sizes of TribeMCL cluster(s) composing the gene family; and 3) previously known importance in immunity, based on literature searches. Those clusters containing proteins with repetitive domains (i.e. leucine-rich-repeats, LRR, or epidermal-growth factor, EGF, domains)
were not included in further analyses, due to the fact that the presence of multiple repeats frustrated similarity clustering. When a diversified family was found that met several of the conditions stated above, a term search was conducted to find relevant hits among the BLAST results (e.g. terms like "serine protease," "trypsin," etc. were used to find serine proteases). The contig list in each TribeMCL cluster was expanded to include all contigs in a TribeMCL cluster if and only if more than or equal to half of the contigs in that cluster had BLASTX best hits to the group of interest and the remainder of contigs had no significant or conflicting hits, whereas the contig list was contracted if less than half of the contigs in that cluster had BLASTX best hits to the group of interest and the remainder of contigs had hits to dissimilar proteins, in which case, all contigs in that cluster were excluded from that group of interest. The contig list was neither contracted nor expanded but remained unchanged in the cases of a TribeMCL cluster in which more than or equal to half of the contigs in that cluster had BLASTX best hits to the group of interest but the remainder of contigs had significant hits to dissimilar proteins or if less than half of the contigs in that cluster had BLASTX best hits to the group of interest and the remainder of contigs had no significant hits. Those contigs (with hits to the gene family of interest) that were not contained in the TribeMCL graph were also retained.

Translated transcripts for each of the select diversified families of interest were reduced to non-redundant sets using CD-HIT, on settings “-G 0 –
aS 0.50 –c 0.95,” which reduced the sets of translated transcripts by 95% redundancy for an alignment length at least 50% of the smaller translated transcript (Weizhong and Godzik 2006). The nucleotide sequence of the non-redundant transcripts for each group of interest was mapped to the *C. gigas* genome using BLAT (Kent 2002; oyster.v9.fa, http://gigadb.org/pacific_oyster/, Zhang et al. 2012). BLAT hits were scored as #Matches + #Repmatches - #Mismatches - #Query Gap Count - #Target Gap Count and scores \( \geq 30 \) were retained. The number of unique *C. gigas* loci was determined by summing the number of non-overlapping regions of the *C. gigas* genome regions to which queries aligned, with each BLAT target region padded by an additional 200 bp upstream and downstream to account to some extent for the possibility of truncated mappings. The estimation of the number of genome loci for each diversified family was separate from the estimation of the number of genes for that family in the *C. gigas* genome. For the remainder of the diversified families of interest, nucleotide sequences for the non-redundant transcripts and protein sequences from *C. gigas* GLEAN gene models were reciprocally compared by BLASTX and TBLASTN, respectively (oyster.v9.glean.rename.gff.pep.gz, http://gigadb.org/pacific_oyster/, Zhang et al. 2012). The number of genes in each family present in the *C. gigas* genome was estimated as the number of peptides that had reciprocal hits to the non-redundant *C. virginica* transcript sequences with e-value \( \leq 1e-05 \) and that had
best BLASTP hits to NCBI’s NR database correspondent to the family of interest.

**SNP detection and polymorphism in candidate oyster host defense genes**

For single nucleotide polymorphism (SNP) detection, processed reads were aligned to the transcriptome using Bowtie2 on “--very-sensitive” setting. PCR duplicates were removed with the Picard Tools utility MarkDuplicates (http://picard.sourceforge.net/). SNPs were called using samtools mpileup with “-g” flag to generate genotype likelihoods (Li et al. 2009). The output was filtered using vcfutils varFilter with minimum mapping quality of 25, minimum read depth of 25, and maximum read depth of 200 (Q 25 -d 25 -D 200). In the R programming environment, SNPs were further filtered for minor allele frequency ≥10% and minor allele count ≥ 5. All transcriptome contigs were aligned to their protein translations using estwise (version 2.2.0) to find the reading frame and start and stop sites (http://www.ebi.ac.uk/Wise2/). Custom R scripts utilizing the R packages, ShortRead and SeqinR, were used to determine whether SNPs were synonymous or non-synonymous based on the reading frame, start and stop sites (Charif et al. 2007; Morgan et al. 2009). For each contig containing SNPs within the determined coding region, a multi-sequence file was generated consisting of the original nucleotide contig sequence, and one nucleotide sequence per SNP consisting of the original
nucleotide sequence modified to reflect the base composition of the SNP.

Each multi-sequence file in categories of interest like DEG groups (GX_early, F3L_early, F3L_late) and diversified gene families, was run through the codeml program in the PAML package (version 4.1b) to obtain the rate of non-synonymous mutations (dN), the rate of synonymous mutations (dS) and the ratio of these two rates (dN/dS) (Yang 2007). The M0 model was assumed of a single $\omega$ for all lineages, initial K and $\omega$ values of 1.0 and 0.5, respectively, and the F3x4 codon frequency model (Goldman and Yang 1994). The final K and $\omega$ values were estimated by codeml (Yang 2007).

Comparison of putative diversified families between diverse taxa

The diversified gene families analyzed in depth using the C. virginica-only TribeMCL graph were tracked and enumerated in the multi-species TribeMCL graph. For each diversified family, the set of nodes first consisting of the known C. virginica nodes was reiteratively expanded with members from other taxa, adding neighbor nodes until the set failed to grow. The final subgraph was reduced by examination of the C. virginica annotations and the C. gigas annotations (downloaded from http://public-contigbrowser.sigenae.org:9090/ Crassostrea_gigas/index.html). Multi-species families were retained if at least half of the contigs had BLASTX hits to the gene family of interest and the remainder of contigs had no significant hits, otherwise, the cluster was excluded. The final set of proteins for each
diversified family of interest were subsequently reduced to non-redundant sets, independently for each family, using CD-HIT, on settings “-G 0 –aS 0.50 –c 0.95,” which reduced the sets of proteins by 95% redundancy for a alignment length at least 50% of the smaller protein (Li and Godzik 2006). The number of non-redundant proteins was enumerated for each species for each diversified gene family. In a few select cases, proteins from different species were so similar as to cluster together, in which case, proteins were counted for both species. For example, two serine proteases from *D. melanogaster* that reduced to one sequence by CD-HIT, would have been counted as one serine protease, while one serine protease each from *D. melanogaster* and *D. pulex* that reduced to one sequence by CD-HIT would have been counted as one serine protease for each species. In addition to an estimation of the number of non-redundant proteins for each species for each diversified family, the number of gene models was estimated for each species for each diversified family, according to the formula:

\[ NPX \times (NGM + NP) = NGMX \]

where \( NPX \) = Number of non-redundant proteins in diversified family X, species Y

\( NGM \) = Number of total gene models in species Y

\( NP \) = Number of total non-redundant proteins used for similarity clustering in species Y

\( NGMX \) = Number of gene models in diversified family X, species Y
Estimations of the total number of gene models for each species (NP, above) were taken from various sources including the published literature and websites of genome sequencing centers (Appendix A, Table S2). Estimates for total number of non-redundant proteins and number of gene models for *L. stagnalis* are not reproduced here because the estimates were unreasonably low.

**Phylogeny of diversified families**

For a select few diversified groups of interest, the non-redundant protein sequences from the multi-species TribeMCL graph were aligned using the E-INS-I option of MAFFT (version 6, Katoh and Toh, 2008). Multiple sequence alignments were viewed in JalView (version 2.6.1) and manually curated to extract blocks of well-aligned sequence (Waterhouse et al. 2009). The models of sequence divergence that best fit the multiple sequence alignments were found using ProtTest (version 3, Abascal et al. 2005). FastTree2 was used to generate a phylogenetic tree (Price et al. 2010). For both IFI44 and GIMAP families, the WAG model was used to model sequence divergence (Whelan and Goldman 2001), and a hybrid of CAT (Stamatakis 2006) and gamma approximations were used to account for rate variation across sites (Price et al. 2010). Trees were viewed as circular cladograms in Dendroscope (version 3.2.2, Huson et al. 2007).
RESULTS

**Oyster survival in response to bacterial challenge**

Oysters from the F3L family experienced a consistent and high rate of mortality after challenge with the bacterial pathogen *R. crassostreae*, reaching over 90% cumulative mortality by the end of the 93-day period (Fig. 1). The survival curve of the challenged F3L oysters was significantly different from all other groups (log-rank survival, \( p < 0.01 \)). Pearson’s chi-squared test with Bonferroni corrections was used to compare oyster survival in the 3 groups used for sequencing, CGX, GX, and F3L, at day 28 (the closest time point at which mortality was tallied before collection of samples for RNA isolation at day 30), and at day 93 (the final time point of the bacterial challenge). F3L had a significantly higher cumulative mortality than GX at day 28 (\( p < 0.01 \)), and a significantly higher cumulative mortality than GX and CGX at day 93 (\( p < 0.01 \)). No significant differences in mortality were observed between unchallenged control oysters (CF3L and CGX) and oysters from the resistant family challenged with *R. crassostreae* at days 28 and 93 after challenge. Oysters from the control ROD-resistant family (CGX) suffered a mortality event between days 1 and 7 (20% cumulative percent mortality by day 7), with an additional 10% mortality for the 86 days following that event (Fig. 1).

**Oyster transcriptome assembly**
From a total of 4.1x10^8 Illumina GAIIx-sequenced cDNA reads of 108 bp, after filtering and trimming, the final set consisted of 3.8x10^6 reads of 94 ± 5 bp (Appendix B. Fig. S1). After the Trinity assembly of 374,029 contigs was further filtered for contaminants, 356,237 contigs remained with a mean length of 440 bp and an N50 of 487 bp (Table 1). A BLASTX search to the NCBI NR protein database led to annotation of 19.8% of the transcriptome. Of the total transcriptome, 22,934 contigs (16.3%) were at least 1 Kb in length.

**Differential gene expression in oysters in response to bacterial challenge**

Differential gene expression analyses of samples GX-1d, GX-5d, F3L-1d, F3L-5d, F3L-15d, and/or F3L-30d yielded a total 6,296 differentially expressed genes (DEGs), or 1.8% of the total de novo-assembled transcriptome. When samples were clustered by gene expression patterns, two major clusters separating F3L and GX/CGX treatments were evident. Furthermore, within each of these clusters, the following subclusters were detected: F3L 1 and 5d (F3L_early); F3L 15 and 30d (F3L_late); GX 1 and 5d (GX_early); and CGX 15 and 30d (control) (Fig. 2). This pattern of treatment-day clustering (GX_early, F3L_early, and F3L_late) was used in further analyses. The focus of the analysis was placed on GX_early DEGs according to the rationale that genes involved in disease resistance would likely be expressed in resistant GX oysters at early time points after exposure to the pathogen.
The more dramatic response to bacterial challenge of F3L compared to GX in terms of cumulative mortality was reflected in a similar differential response in terms of gene expression (Fig. 3). Of the 356,237 total transcripts tested for differential expression, 6,097 transcripts were differentially expressed in F3L_early and/or F3L_late, compared to only 552 transcripts differentially expressed in GX_early. F3L DEGs were described at the gene level where overlap was found with GX_early DEGs and were further described at the scale of Gene Ontology functional enrichment, Pfam enrichment, and TribeMCL-DEG enrichment (Tables 2 - 5). A greater share of GX_early DEGs was shared with F3L_early and/or F3L_late DEGs (64%) than was unique to GX_early (36%) (Fig. 3).

DEGs shared by GX_early and F3L treatments should include (among others) genes associated with host defense and supporting functions (Tables 4 & 5). DEGs unique to GX_early should include genes contributing to disease resistance in the GX family (Tables 2 & 3). The most highly differentially expressed, annotated, up-regulated DEGs unique to GX_early (potentially involved in disease resistance) included, among others, transcripts that annotated to 2 scavenger receptor cysteine-rich proteins, 2 fibropellin ia proteins, 2 inhibitor of apoptosis (IAP) proteins, cytochrome p450, interleukin 17d, a fibrinolytic enzyme, and a disintegrin and metalloprotease with thrombospondin motifs 8 (ADAMTS8) (Table 2). The most highly differentially expressed, annotated, down-regulated DEGs unique to GX_early included
transcripts that annotated to the development-related protein rapunzel, 2 collagen proteins, tenascin xb, 2 cubilin proteins, inhibitor of apoptosis (IAP), a c-type lectin, a melatonin receptor (Table 3). The most highly differentially expressed, annotated, up-regulated DEGs shared between GX_early and F3L_early and/or F3L_late (potentially involved in responses to bacterial infection) included transcripts that annotated to 2 serine protease inhibitors, 2 dopamine beta-hydroxylases, 2 fatty acid synthases, 2 sulfatases, cytochrome p450, a C1q domain-containing (C1qDC) protein, and heat shock protein 60 (HSP60) (Table 4). The most highly differentially expressed, annotated, down-regulated DEGs shared between GX_early and F3L_early and/or F3L_late included transcripts that annotated to a C1qDC protein, two monocarboxylate transporters, multiple epidermal growth factor 11 (MEGF 11), deleted in malignant brain tumors 1 (DMBT1), and sushi-repeat-containing x-linked 2 (Table 4). Because different contigs sometimes shared the similar annotations (suggesting that they could potentially be members of a gene family, alternatively spliced forms, or parts of the same transcript that were not assembled together), annotations were manually compared to find truly unique transcripts among GX_early unique DEGs. Sixteen non-redundant transcript annotations were found in the GX_early sample, including arginase I, arginase II, rho gtpase, and cubilin (down-regulated at day 5); and unc-5, furin, and interleukin 17d (up-regulated at day 1) (Appendix A. Table S3).
The relative ratio of up-regulated and down-regulated DEGs differed between GX and F3L, with a significantly greater number of up-regulated transcripts in GX_early DEGs (Pearson’s chi-squared analysis, $p < 0.01$) and a significantly greater number of down-regulated transcripts in F3L DEGs ($p < 0.01$).

**Gene Ontology categories enriched among oyster DEGs upon bacterial challenge**

As there were fewer DEGs in GX_early compared to the F3L groups, so there were comparatively fewer enriched GO terms (Fig. 4). The most significantly enriched biological process GO term among GX_early up-regulated DEGs was “protein folding”, corresponding to 3 DEGs annotated to HSP60 that were up-regulated only at day 1. “Protein folding” was also enriched among F3L_early DEGs, also corresponding to HSP60 transcripts, some of which were down-regulated and some up-regulated. The enrichment in “defense response” among GX_early DEGs corresponded to transcripts that annotated to interleukin 17 (IL17), which were strongly up-regulated at day 1. Terms closely allied to defense response were found enriched among F3L_early DEGs including “defense response to bacterium” and “response to molecule of bacterial origin”, corresponding to transcripts that annotated to angiotensin-converting enzyme, defensin, and immune-responsive gene 1, the latter of which was also up-regulated in GX at day 1. The other biological
process terms enriched among up-regulated GX_early DEGs were the related terms “programmed cell death” and “apoptotic process”, corresponding to transcripts that annotated to IAP transcripts. Though related cell death terms were not found to be enriched in F3L treatments, 3 IAP transcripts were differentially expressed in F3L, 2 of which were down-regulated at all time points. The most significantly enriched F3L early biological process term, found among up-regulated DEGs, was “cholesterol transport”, corresponding to epididymal secretory protein E1. Other transport terms enriched among F3L_early DEGs include “hexose transport” (up-regulated) and “amino acid transport”-related and “carboxylic acid transport” (down-regulated). Various development-related terms, including the closely related terms “blood vessel morphogenesis,” “angiogenesis,” and “vascular development” as well as “fin development” were enriched among F3L_early down-regulated DEGs while “fin development” and neuron-related development terms were enriched among F3L_late down-regulated DEGs. Several terms related to carboxylic acid metabolism were uniquely enriched among F3L_late up-regulated DEGs, corresponding to various decarboxylases. Also uniquely enriched among F3L_late up-regulated DEGs was “oxidation-reduction process”, corresponding to several cytochrome p450 transcripts.

With respect to enriched molecular function GO terms, commonalities across treatments included “enzyme inhibitor activity” among GX_early and F3L_late up-regulated DEGs (Fig. 4). The closely related terms
“endopeptidase inhibitor/regulator activity” were enriched among F3L_early up-regulated DEGs, while “peptidase inhibitor activity” was enriched among F3L_late up-regulated DEGs and was the most significantly enriched molecular function term among F3L_late DEGs. Also enriched among F3L_early up-regulated DEGs was the highly significant term “monooxygenase activity”, corresponding to several cytochrome p450 and dopamine beta hydroxylase transcripts, while among F3L_late up-regulated DEGs the term “oxidoreductase activity” was enriched, also corresponding to several cytochrome p450 and other transcripts. There were several “hydrolase”-related terms enriched among F3L_early DEGs including “hydrolase activity, hydrolyzing O-glycosyl compounds”, corresponding to CHIT3 protein among others.

The GO Cellular Component hierarchical superstructure is not as ramified as are Biological Process and Molecular Function, and accordingly, the enriched terms found here were fewer. “Extracellular region” was the only cellular component GO term enriched among GX_early up-regulated DEGs, and was also enriched among F3L_early up-regulated and F3L_late down-regulated DEGs (Fig. 4). The transcripts that annotated to this term were diverse and included tissue inhibitors of metalloproteinase (TIMPs), HSP60, fibropellin, and others. Interestingly, all enriched cellular component terms were related to the membrane or extracellular matrix.
**Pfam protein families and domains enriched among oyster DEGs upon bacterial challenge**

Of 356,237 contigs, 21,446 contigs (6%) had significant hits to 2,367 different Pfam families. The most significantly enriched Pfam family among GX_early DEGs, which was also enriched among F3L_early DEGs, was NAD_binding_5, a family of myo-inositol-1-phosphate synthases involved in signal transduction and the mobilization of calcium (Table 5; Berridge 1984). Several proteins in the thrombospondin_1 (TSP_1) family, including 7 hemicentin-like transcripts, were differentially expressed in all treatments, with one of these up-regulated at days 1 and 5 in GX. Transcripts matching the AIG1 family included many annotating as GIMAP proteins; nearly all GIMAP DEGs were down-regulated in F3L (data not shown). The remainder of the Pfam families enriched among GX_early DEGs were enriched solely in that class of DEGs including IL17, HSP20, bZIP_2 (a family of transcription factors including creb-binding protein; Schumacher et al. 2000), patched (involved in developmental signaling; Ingham et al. 1998), and sulfatase. Families enriched among DEGs from both F3L_early and F3L_late and not GX_early included An_peroxidase (response to oxidative stress and bactericidal defense; Zamocky et al. 2008), mannose-6-phosphate receptors (Man-6-P_recep, involved in biogenesis of lysosomes; Griffiths et al. 1998), Dynamin_N (necessary for endocytosis; McClure and Robinson 1996), and ApoL (role in lipid transport, linked to innate immunity in humans; Perez-Morga...
et al. 2005; Vanhollebeke et al. 2006). Slightly greater than half (18) of the 39 Pfam families enriched among F3L_early DEGs were uniquely enriched among that class of DEGs. Among Pfam families uniquely enriched among F3L_early DEGs, some have obvious relevance to host defense including tissue inhibitors of metalloproteinase (TIMP), Pacifastin_1 (serine protease inhibitor; Simonet et al. 2003), and Von Willebrand Factor (VWF) type D domain-containing proteins (involved in blood clotting and cell adhesion; Jorieux et al. 1998). Other Pfam families uniquely enriched among F3L_early DEGs included several types of transporters including SSF, Sugar_tr, SNF, and AA_permease_2. Fewer than half (8) of the 21 Pfam families enriched among F3L_late DEGs were uniquely enriched among that class of DEGs, including ovomucin-binding proteins (VOMI), proteins that contain the C-terminal domain of Chitobiase/beta-hexosaminidase (CHB_HEX_C_1, which degrade chitin), and Mucin2_WxxW, (help form an insoluble extracellular matrix that protects epithelial linings; http://pfam.sanger.ac.uk/; Johansson et al. 2011).

Of 356,237 contigs, 23,220 contigs (6.5%) had significant hits to 1,591 different Pfam domains. The most significantly enriched Pfam domain among F3L_early and second most significantly enriched among F3L_late DEGs, was C1q, which was also enriched among GX_early DEGs (Table 6). Only three other domains were enriched among all classes of DEGs, namely, SRCR, Kunitz_BPTI, and HYR. Only two domains were enriched among F3L_early
DEGs and GX_early DEGs, including Trypsin (protease involved in digestion and innate immunity; Rawlings and Barrett 1994; Ross et al. 2003). Most of the domains that were enriched among GX_early DEGs were unique to that class of DEGs and included baculovirus inhibitor of apoptosis repeat (BIR) protein (involved in the inhibition of apoptosis; Silke and Vaux 2001); Complement Clr-like EGF (cEGF)-like (involved in blood coagulation; Wouters et al. 2005); and CUB (involved in a variety of processes including inflammation, angiogenesis, and endocytosis; Blanc et al. 2007). Ten Pfam domains were enriched only among F3L_early and F3L_late DEGs including Fibrinogen_C; Stichodactyla toxin (ShK, present in proteins that block voltage-gated potassium channels and found in antimicrobial proteins in C. elegans; Tudor et al. 1996; Troemel et al. 2006); and Copper type II ascorbate-dependent monooxygenase (Cu2_monooxygen; present in dopamine beta-hydroxylases and associated with immunity through the regulation of catecholamine biosynthesis; Flieri et al. 2009). Twenty-eight of the 44 Pfam domains enriched among F3L_early DEGs were uniquely enriched among that class of DEGs and included Von Willebrand Factor A (VWA, present in mostly extracellular proteins involved in cell migration, cell adhesion, and other processes; Colombatti et al. 1991; Colombatti et al. 1993) and Kazal_1 (present in serine protease inhibitors; Rawlings et al. 1994). Only 4 of 18 Pfam domains enriched among F3L_late DEGs were uniquely enriched among that class of DEGs and included Cupin_8, found here in transcripts that annotated...
as jumonji domain-containing proteins (involved in histone demethylation; Klose et al. 2006) and phosphatidylserine receptors (involved in apoptotic cell clearance; Li et al. 2003).

**Sequence similarity clusters enriched for oyster DEGs upon bacterial challenge**

Of 356,237 contigs, 82,498 contigs (23%) were clustered by TribeMCL similarity clustering into 18,873 clusters, 187 of which were enriched for F3L_early, F3L_late, and/or GX_early DEGs (Fisher’s exact test or Pearson’s chi squared test, \( p < 0.01 \)). The top six clusters most significantly enriched for GX_early DEGs were also enriched for F3L_early DEGs and sometimes for F3L_late DEGs. Generally, the clusters most significantly enriched for F3L_early DEGs were also enriched for F3L_late DEGs and *vice versa*.

Several putative diversified gene families were selected for further analyses based on the criteria listed in the methods. These included serine proteases (SPs), serine protease inhibitors (SPIs), fibrinogen domain-containing proteins/ fibrinogen-related proteins (FREDs/FREPs), C1qDC proteins, c-type lectin domain-containing (CTLDC) proteins, deleted in malignant brain tumors 1 (DMBT1) and scavenger receptor cysteine-rich type 12, interferon-induced protein 44 (IFI44), and GTPase of the immunity associated protein family (GIMAP) proteins (Table 9).
Characteristics of selected putative diversified gene families in oysters in terms size, polymorphism, and response to bacterial challenge

C1qDC was among the largest gene families differentially expressed in *C. virginica* in response to bacterial challenge, with 391 non-redundant annotated transcripts, 323 of which mapped to 149 genomic loci (non-overlapping mappings ±200bp) in the *C. gigas* genome (Table 9). The estimated number of C1qDC transcripts in the *C. virginica* transcriptome is consistent with the recent estimate made by Zhang et al. (2012) of 321 C1qDC genes in the *C. gigas* genome. The widest disparity between the number of family members identified in the *de novo* transcriptome for *C. virginica* and the numbers estimated from the *C. gigas* genome was observed for the GIMAP family (Table 9). Of a total of 173 non-redundant GIMAP *C. virginica* transcripts, 158 transcripts mapped to 33 *C. gigas* loci. Only 19 genes were found in the *C. gigas* set of gene models. Certain *C. gigas* loci had a great many mapped GIMAP transcripts, including one locus of 11 transcript hits, two loci of 12 transcript hits, one locus of 17 transcript hits, and one locus of 25 transcript hits (data not shown).

The average rate of polymorphism for the entire transcriptome, based on our stringent thresholds, was 272 ± 299 bp/SNP (mean ± SD). Forty-nine non-synonymous SNPs and 109 non-synonymous SNPs were identified in transcripts uniquely differentially expressed in GX_early and in both GX_early and F3L, respectively (Appendix A, Table S4). There was a wide range of
variation (from tens to hundreds) in the number of non-synonymous SNPs that were identified in each of the selected diversified gene families (Appendix A, Table S4). Mean dN/dS must be interpreted with caution given the limited numbers of individuals in each sample (limited SNP density) and the necessity of adopting certain assumptions (initial \( \omega/K \), codon freq. model, etc.). While many individual transcripts had dN/dS values > 1, limited SNP density led to a consideration of dN/dS by gene family rather than by sequence or site. Serine proteases had the lowest mean dN/dS, by at least an order of 2, suggesting that this group is under stronger functional constraint than other groups. The highest mean dN/dS belonged to C1qDC proteins and IFI44.

In terms of differential expression within the diversified families of interest, nearly all differentially expressed SPs were up-regulated at GX day 5 or F3L day 5; nearly all SPIs were up-regulated at GX day 1 and 5 or F3L day 1 and 5; all FREDs/FREPs were down-regulated in F3L at one or more time points and no FREDs/FREPs were differentially expressed in GX; nearly all GIMAP DEGs were down-regulated in F3L at one or more time points and a few were down-regulated in GX day 1, while one transcript was up-regulated in GX day 5 (Fig. 6); nearly all IFI44 DEGs were down-regulated in F3L at one or more time points, while one transcript was up-regulated at F3L day 5, and one was up-regulated in GX at day 5 (Fig. 7). For the remainder of the diversified families, C1qDC, CTLDC, and DMBT1/SRCR type 12, a consistent pattern of differential expression could not be observed (data not shown). For
the relatively few transcripts in the select diversified families that were differentially expressed in both GX_early and F3L_early (19 transcripts total), the direction of differential expression, that is, up- or down-regulation relative to control, tended to match between GX_early and F3L_early, except for 2 CTLDC transcripts and 1 DMBT1/SR type 12 transcript (data not shown).

**Comparison of size and phylogeny of selected diversified gene families in oysters and other organisms**

Certain patterns could be seen in the size of selected diversified gene families across taxa and across the diversified groups (Fig. 5). First, the number of proteins and the number of gene models for each diversified family varied widely, especially for species for which ESTs or poorly-curated gene models were used. FREPs/FREDs and CTLDC proteins, known to be highly expanded in molluscs (Ghosh et al. 2010; Gerdol et al. 2011), appeared to have the greatest number of members. Compared to the number of C1qDC proteins present in molluscan species, few are present in non-molluscan species. Scavenger receptors with class B SRCR domains, well studied in *S. purpuratus*, showed tens of proteins/gene models in the bivalve species considered here. The warm colors signifying a high column z-score for IFI44 for bivalve species and for GIMAP proteins for molluscan species indicates that IFI44 is greatly expanded in bivalves and GIMAP is greatly expanded in molluscs. No IFI44 transcripts were found in the gastropod species considered
here and few were found in other non-bivalve invertebrate species.

Phylogenetic analysis of GIMAP sequences showed 3 major clusters (Fig. 8 and 9). One major cluster of sequences contained, in addition to many C. *virginica* sequences, comparable numbers of sequences from the other bivalves species, including *P. fucata* and *C. gigas*, and a comparable number of sequences from one gastropod, *L. gigantea*, along with one sequence each from the hemichordate, *S. kowalevskii*, the amphioxus, *B. floridai*, and the basal deuterostome, *N. vectensis*. The second major cluster contained many sequences from all mollusc species, *S. kowalevskii*, and *B. floridai*. The third major cluster contained the majority of *C. virginica* sequences (>100), about 10 *C. gigas* sequences, 1 *P. fucata* sequence, and 1 *B. floridai* sequence. Because the 150+ non-redundant GIMAP *C. virginica* transcripts mapped to only 33 *C. gigas* loci, I conclude that some of the diversification (expansion in protein number relative to other organisms) derives from mechanisms different from gene duplication like polymorphism and/or alternative splicing.

Phylogenetic analysis of IFI44 sequences showed two main clusters, one of which was composed solely of bivalve sequences (159 sequences) and the other of which was composed of 18 sequences from 8 diverse organisms including *C. virginica* (Fig. 10 and 11). The main bivalve-only group could be further subdivided into subgroups that had variable numbers of sequences from both *C. virginica* and *C. gigas*. 
DISCUSSION

The present study provides a rich view of the processes and genes that constitute the oyster host defense responses to bacterial challenge. Extracellular matrix (ECM) restructuring, cell adhesion, inflammation, metabolism, catecholamine signaling, and several other processes are key in distinguishing resistance from susceptibility to Roseovarius Oyster Disease. The present study also identified 8 putative diversified gene families important in the host defenses of Eastern oysters, including two families, GTPase of the immunity associated protein (GIMAP) and interferon-induced protein 44 (IFI44) families, which appear to be expanded in molluscs and bivalves, respectively.

Oysters from the resistant family did not show clinical signs of infection and suffered mortalities comparable to non-challenged oysters, suggesting that these oysters were able to eliminate the pathogen rapidly. Prior to a discussion of differential gene expression and gene family analysis results, it should be noted that only 20% of the transcriptome could be annotated by BLAST and only 30% of the transcriptome could be described by the combination of BLAST similarity, Pfam, and TribeMCL clustering. Conclusions should be regarded with caution and viewed as foundation for future study. With that in mind, comparison of the patterns of gene expression between resistant and susceptible oysters suggest that resistance to R. crassostreae may involve a targeted hemocytic response followed by tight control of inflammatory processes and detoxification.
ROD-resistant juvenile oysters responded to the bacterial pathogen *R. crassostreae* mainly by up/down-regulating the expression of transcripts coding for proteins that modify the extracellular matrix, proteins that bind self or non-self ligands, stress proteins, and receptors and/or proteins involved in signaling. The interaction between *R. crassostreae* and its oyster host occurs primarily in the extracellular matrix based on DEG annotations and enriched GO terms (Fig. 4). The unique up-regulation in resistant oysters of several subtilisin-like pro-protein convertases (PPC) (Table 2) suggests the involvement of neuroendocrine signaling and/or host defense-relevant protein processing. PPCs are involved in the processing of von Willebrand Factor, matrix metalloproteinases, and antimicrobial peptides in invertebrates, having multiple downstream effects on cell migration, differentiation, inflammation control, and the restructuring of the ECM (Sato et al. 1996; Parks et al. 2004). The importance of ECM proteolysis and restructuring in the response of resistant oysters to bacterial challenge is corroborated by the up-regulation of ADAMTS8 (Table 2), a matrix metalloproteinase that negatively regulates proliferation and participates in ECM proteolysis (Apte 2004, Feinberg and Weiss 2009), the up-regulation of a fibrinolytic enzyme (Table 2), and the down-regulation of tenascin-xb (Table 3), a glycoprotein involved in wound healing and matrix maturation with anti-adhesive properties (Egging et al. 2007). The multiple transcripts that annotate as tenascin in oysters further dispel the notion that tenascins are unique to chordates (Tucker et al. 2006),
as tenascin-like transcripts have also been found in the transcriptome of the Antarctic bivalve *Laternula elliptica* (Clark et al. 2010). Other transcripts involved in ECM restructuring and cell adhesion identified here as distinguishing resistance and susceptibility include those coding for hemicentin and fibropellin-ia. Hemicentin increases cell adhesion and re-shapes areas of cell contact in *C. elegans* (Vogel and Hedgecock 2001). Fibropellin-ia increases cell adhesion in sea urchin embryos (Burke et al. 1998). By sequence similarity, fibropellin transcripts separated into a cluster enriched for susceptible DEGs and another cluster enriched for resistant DEGs (Table 7). Cell adhesion has long been known to be important in invertebrate innate immunity in general (Johansson 1999) and in oyster immunity (Gueguen et al. 2003). I hypothesize that the differential response of resistant oysters in respect to ECM restructuring and cell adhesion molecules enabled a more effective hemocytic response in these oysters, facilitating cell migration to the extrapallial cavity (the space between the oyster mantle tissue and the shell, where ROD is known to have its primary effects, Paillard et al. 1996; Boardman et al. 2008), likely followed by aggregation, phagocytosis, and apoptosis (Terahara et al. 2005; Anderson et al. 2011; de Lorgeril et al. 2011). Phagocytosis of the bacterial pathogen *Vibrio tapetis* by carpet shell clam (*Ruditapes decussatus*) hemocytes has been shown to have an important role in the resistance of this clam species to Brown Ring Disease (BRD), a disease with many similarities in pathology to ROD (Allam et al. 2001; Allam and Ford...
I hypothesize that the cell adhesion transcripts seen in the present study may play a role in the defense capabilities of hemocytes in resistant oysters.

The early resistant response also involved the pro-inflammatory mediator, interleukin 17 (IL17), and the nitric oxide modulator, arginase. Previous research shows that injection of heat-killed gram-positive and gram-negative bacteria into *C. gigas* oysters produces a rapid and transient up-regulation in IL17 transcript abundance in hemocytes, suggesting that IL17 is an important mediator of the pro-inflammatory response in oysters (Roberts et al. 2008). My results support an important role for IL17 in the immune response of oysters against bacterial infection and a potential role in disease resistance to ROD. Here, while IL17 was uniquely up-regulated in resistant oysters, arginase was uniquely down-regulated (Table 3). Arginases have been shown in macrophages to modulate the production of nitric oxide (Chang et al. 1998), which is an immune effector in the Eastern oyster (Villamil et al. 2007). Using microarray technology, a transcript annotating as arginase was shown to increase rapidly after 6 h of heat stress in *C. gigas* (Lang et al. 2009). The down-regulation of arginase in resistant oysters on day 5 may signalize a down-regulation of the inflammation and stress response following a successful defense response.

Genes and processes activated in susceptible oysters in response to bacterial challenge and absent or present to a much lesser degree in resistant
oysters provide potential information on the molecular basis for disease susceptibility or are signs of an unsuccessful defense response. Many transcripts involved in metabolic processes (e.g. carbohydrate metabolism) were differentially expressed in susceptible oysters at both early and late time points following bacterial challenge, but not in resistant oysters as illustrated by enriched molecular function GO terms (Fig. 4). A decrease in energy metabolic enzyme activity and a down-regulation of genes related to energy metabolism have been shown to coincide with mortality events in the Eastern oyster (Genard et al. 2011; 2012).

The up-regulation of multiple dopamine beta-hydroxylase (DBH) transcripts early in both resistant and susceptible oysters (Table 4) suggests the role of catecholamine signaling in host defenses against bacteria. However, both DBH and Dopa decarboxylase (DDC) families/clusters, as well as several transcripts containing the tyrosinase domain were uniquely enriched for susceptible DEGs (Table 7 & 8) suggesting that catecholamine signaling and/or melanization were greater in susceptible oysters. DBH and DDC produce/modify catecholamines, which have been shown to modulate both the immune and stress response in the scallop Chlamys farreri (Chen et al. 2008; Zhou et al. 2011a; Zhou et al. 2011b). Crassostrea gigas hemocytes have been shown to respond to neuroendocrine signaling with changes in gene expression (Bricelj et al. 1992; Lacosta et al. 2001). It can be imagined that hemocytes at the site of ROD infection and injury are both signaling and
responding to neuroendocrine signals to coordinate a response. DDC, along with enzymes with phenoloxidase activity like tyrosinases, also participates in melanization, a process whose products can kill bacteria (Kan et al. 2008, Sideri et al. 2008) and which may be responsible for the pigmentation characteristic of the conchiolin depositions characteristic of ROD (ref). Both resistant and susceptible oysters appear to use catecholamine signaling to coordinate a response but they may have a differential response in terms of melanization, as supported by the up-regulation of tyrosinase-like transcripts in susceptible oysters. It is not known, however, whether the differentially expressed tyrosinases in the present study had phenoloxidase activity. While tyrosinase has been shown to have phenoloxidase and antibacterial activities in several bivalves including the scallop Chlamys farreri (Zhou et al. 2012), Manila clams, and Sydney rock oysters, laccase and catecholase but not tyrosinase were shown to be responsible for phenoloxidase activity in C. gigas or C. virginica (Luna-Acosta et al. 2011 and references therein). A BLAST search for reciprocal hits between the tyrosinase transcripts in the present transcriptome and the C. gigas translated GLEAN gene models revealed 24 tyrosinase-like peptides therein, providing evidence that C. gigas, too, has tyrosinase proteins.

The host defense response to bacterial challenge in resistant and susceptible oysters shared some commonalities, including the involvement of catecholamine signaling (discussed above), detoxification, and apoptosis.
Transcripts that were highly up-regulated in both resistant and susceptible oysters included transcripts annotating as glutathione s-transferase, cytochrome p450, and heat shock protein 60 (HSP60), which are involved in preventing oxidative damage (Table 4). Glutathione s-transferase is an antioxidant and is up-regulated in hemocytes of oysters challenged with a pathogenic Vibrio sp. (de Lorgeril et al. 2011). Although cytochrome p450s have been best studied in detoxification of xenobiotics in bivalves (Snyder 2000), they have also been implicated in the host defense response of the flat oyster Ostrea edulis to the parasite Bonamia ostreae (Morga et al. 2011) and the clam Ruditapes philippinarum to Vibrio tapetis (Brulle et al. 2012). HSP60 is involved in xenobiotic detoxification and the stress response in oysters (Ivanina et al. 2008). Generally, detoxification was intensified in susceptible oysters as related detoxification terms like “monooxygenase activity” and “oxidoreductase activity” were functionally enriched among the susceptible but not the resistant DEGs (Fig. 4). Detoxification-related transcripts are highly up-regulated in C. virginica prior to mass mortality events (Genard et al. 2012). While an early response of detoxification/stress transcripts may contribute to resistance, a persistent and more generalized response (that is, a greater number of up-regulated stress transcripts) may signalize imminent mortality. Another process involved in both the resistant and susceptible response was apoptosis. Inhibitor or apoptosis (IAP) transcripts were found to be both up- and down-regulated among resistant oyster DEGs and largely down-regulated
among susceptible oyster DEGs. IAP proteins are associated with molluscan immunity (Sokolova 2009), participating in the defense response of clams to BRD (Donaghy et al. 2009; Brulle et al. 2012), so they likely play a role in the oyster response to ROD.

Transcripts uniquely expressed in susceptible oysters (F3L) at both early and late time points may reveal genes linked to stress or mortality, since this family suffered consistent levels of mortality throughout the challenge. The most significantly enriched Pfam family among late susceptible oyster DEGs was ankyrin Ank_2 (Table 6). The ankyrin domain serves as a mediator of protein-protein interactions in proteins with a wide variety of functions (Bennett and Baines 2001). Little is known about the function of ankyrin transcripts, which includes ankyrin unc-44, nacht and ankyrin domain containing, and other ankyrin repeat-containing transcripts, in oysters. Ankyrin repeat-containing proteins have been shown to be differentially expressed in C. gigas as a result of heat shock, exposure to Vibrio, and between resistant and susceptible C. gigas to summer mortality (Lang et al. 2009; de Lorgeril et al. 2011; Fleury et al. 2012). Further research is necessary to elucidate the role of these proteins in oyster immunity.

Studies in multiple marine invertebrates suggest that diversified groups of receptors, regulators, and/or effectors enable these organisms to meet the challenge of counteracting pathogens and parasites with relatively short generation times and high mutation rates, without the adaptability of the
adaptive immune system (Messier-Solek et al. 2010). Two gene family analysis approaches, Pfam annotation and TribeMCL clustering, followed by enrichment testing for DEGs enabled the identification of putative “diversified” gene families. Together the two approaches covered some 96K contigs (27% of transcriptome) at the protein/gene family level. Pfam annotated 13K contigs not clustered by TribeMCL and TribeMCL clustered 50K contigs not annotated by Pfam and both approaches covered 38K contigs not annotated using BLAST. When used to describe differential expression at the gene family and domain level (in the case of Pfam domains) both the Pfam and TribeMCL approaches largely complimented one another and helped to distinguish the resistant from susceptible responses. The Pfam approach was more specific and automatically included annotation, yet it described a smaller portion of the transcriptome and was of no use for previously unannotated or shallowly annotated gene families. While TribeMCL clusters sometimes contained false positives (likely because of domain sharing), these were filtered when examining specific families. Because TribeMCL clustering was unbiased and more sensitive, it was used as the primary technique for the identification of enumeration of the members of 8 potentially diversified gene families of immune relevance. Some candidate diversified gene families, like C1q domain-containing proteins spanned multiple TribeMCL clusters.

The abundance and diversity of serine protease (SP) and serine protease inhibitor (SPI) transcripts, combined with the observed patterns of
differential expression, suggest a role of these gene families as diversified effectors of oyster host defense to bacterial challenge. The role of SPIs in oyster host defense has long been recognized, although previous studies mainly focused on the role of these molecules in the interplay between the Eastern oyster and the protozoan parasite *Perkinsus marinus*. The hemolymph of *Crassostrea* spp. contains effective protease inhibitors and *C. gigas*, naturally resistant to *P. marinus*, has significantly greater inhibitory activity than *C. virginica*, which has been interpreted as suggestive of the role of protease inhibitors in host defense and resistance (Faisal et al. 1998; Jenny et al. 2002). The list of known oyster SPs and SPIs has grown with each EST analysis (Gueguen et al. 2003; Roberts et al. 2008) and several SPIs have been biochemically characterized (Xue et al. 2006; Xue et al. 2009; La Peyre et al. 2010). Polymorphism in the promoter of an Eastern oyster SPI has been associated with disease resistance to *P. marinus* (Yu et al. 2011). Proteases from *Perkinsus* sp. inhibit phagocytosis of *Vibrio tapetis* in clams (Ordas et al. 1999) and the virulent effects of *V. tapetis* on clam hemocytes are consistent with the effects of bacterial proteases (Borrego et al. 1996; Allam and Ford 2006). I hypothesize that SPIs can neutralize *R. crassostreae* proteases. Less work has been conducted on the role of SPs in oyster host defense, yet SPs may take part in both digestion and host defense, as in *Drosophila* spp. (Ross et al. 2003). Here, none of the *C. virginica* SPs annotated as clip domain SPs, which are important in insect immunity. However, because the clip domain is
N-terminal to the chymotrypsin domain, true clip domain SP transcripts in the transcriptome may have had their clip domains truncated. A clip domain SP has been found in the scallop *C. farreri* (Zhu et al. 2008) and the pearl oyster *P. fucata* (Zhang et al. 2009). I speculate that at least some of the SPs identified here are truly clip domain SPs. More work will need to be done to definitively describe the domain architecture of Eastern oyster SPs and their role in the host defense response.

Another putative diversified gene family herein identified as abundant and differentially expressed in response to bacterial challenge was the fibrinogen domain-containing gene family. The patterns of expression of members of the FBG domain-containing family in oysters in response to bacterial challenge differed between resistant and susceptible oysters, suggesting a potential role in disease resistance/susceptibility. Invertebrate fibrinogen domain-containing proteins— all of which have a C-terminal fibrinogen (FBG) domain—include fibrinogen-related proteins (FREPs), which contain one or two N-terminal IgSF domains; fibrinogen-related molecules (FREMs), which contain epidermal growth factor-like repeats; and fibrinogen-related domain-containing (FREDs) which includes all FBG domain-containing proteins that do not fit into the other aforementioned groups. These proteins function in pathogen recognition, agglutination, and parasite resistance (Hanington and Zhang 2011). FBG domain-containing proteins, particularly FREPs, have been studied in depth in the gastropod *B. glabrata*, perhaps
partially because of the surprising finding that FREP genes can diversify somatically (Zhang et al. 2004). FREPs have been shown in *B. glabrata* to contribute to resistance against the parasite *Schistosoma mansoni* (Hertel et al. 2005). A FREP in the bay scallop *Argopecten irradians* has been shown to have agglutinating activity against chicken and human erythrocytes and bacteria and to increase in expression following challenge by gram-negative bacteria (Zhang et al. 2009). No IgSF domains were found in the identified FBG domain-containing transcripts in the Eastern oyster, while one potential FREM-like transcript with five EGF-like repeats was identified. This does not mean that FREPs are absent from the oyster transcriptome—as 190 FREPs were recently found in the *C. gigas* genome (Zhang et al. 2012)—but underscores the 3’-bias of cDNA sequencing following poly-A capture and the limitation of annotation with transcripts that are rarely full-length.

My results also suggest a role of the gene families DMBT1/SR type 12, C1qDC, and CTLDC in the oyster host defense response. Scavenger receptors with class B SRCR domains have undergone expansions in *S. purpuratus* and show differential expression following challenge by fungi and bacteria (Pancer 2000; Hibino et al. 2006). Scavenger receptors have not undergone comparable expansion/diversification in *D. melanogaster*, *C. intestinalis*, nor in *C. elegans* (Hibino et al. 2006). Scavenger receptor diversity in lophotrochozoans has yet to be adequately addressed. These proteins are differentially expressed in oyster species in response to summer
mortality (Huvet et al. 2004; Fleury et al. 2010) and hypoxia (David et al. 2005). Recently, a SR protein has been characterized in the scallop C. farreri that is up-regulated significantly by exposure to PAMPs like LPS, peptidoglycan and β-glucan and can bind LPS and peptidoglycan (Liu et al. 2011). The discovery here of tens to hundreds of non-redundant sequences in each of several oyster species and the differential expression of many SR sequences in response to bacterial challenge in C. virginica suggests that SR proteins may play a role in oysters comparable to that in C. farreri and with diversity, as a gene family, that may approach that of S. purpuratus. Several oyster transcripts also annotated to class A and F SRs, the latter of which consist mostly of annotations to cell death abnormality-1, which has been linked in C. elegans to the unfolded protein response, apoptotic cell debris engulfment, and resistance to at least one species of bacteria (Lamitia and Cherry, 2008; Haskins et al. 2008). While apparently very abundant in the transcriptome, class F SRs contain epidermal growth factor-like (egf-like) domains present in serial repetitions that frustrate similarity clustering by TribeMCL. Enrichment for DEGs was a prerequisite for consideration of a gene family as relevant to immunity. Adulteration of class F SR clusters with other multiple egf-like domain-containing proteins thwarted enrichment testing and might have possibly precluded definitive enumeration of class F SRs, which illustrates the limitations of similarity clustering, especially when using transcripts assembled from short reads in the absence of a reference genome.
The most abundant putative diversified family studied was the C1qDC gene family. C1qDC proteins can participate in self and non-self binding and function in a variety of processes such as agglutination, cell adhesion, inflammation, and clearance of apoptotic bodies (Kishore et al. 2004). The high sequence variability of C1qDC transcripts in C. virginica has also been shown in transcripts in another bivalve, the mussel *M. galloprovincialis* (Gerdol et al. 2011). C1qDC transcripts are up-regulated in *M. galloprovincialis* following gram-positive and gram-negative bacterial challenge and show highest tissue-specific expression in hemocytes (Gerdol et al. 2011). The role of C1qDC proteins as pattern recognition receptors (PRRs) has been solidified by a demonstration of the ability of a recombinant C1qDC protein from the scallop *Argopecten irradians* to bind PAMPs from diverse pathogens including gram-negative and gram-positive bacteria and fungi (Kong et al. 2010). Results of the present study suggest that the Eastern oyster expresses a great number of C1qDC proteins and that they play a role in host defense against gram-negative bacteria. While > 100 non-redundant translated transcripts/proteins were found by network similarity clustering in all three oyster species examined, only a handful of transcripts were found in gastropod species (Fig. 5). These observations are consistent with the hypothesis that C1qDC genes likely expanded in bivalves, independent from the expansion in the chordate lineage (Gerdol et al. 2011).
Another abundant putative diversified gene family consisted of CTL domain-containing (CTLDC) transcripts. CTLDC proteins are extracellular proteins that contain conserved carbohydrate recognition CTL or CRD domains and function in processes as diverse as cell adhesion, endocytosis, activation of antimicrobials, and pathogen recognition and agglutination (Weise et al. 2006). In *C. elegans*, a total of 278 diversified CTLDC genes have been identified, some of which show differential expression upon pathogen challenge (Schulenburg et al. 2008). Recently a CTLDC from *C. farreri* was shown to act as a PRR, binding LPS and β-glucan, and as an opsonin, enhancing the phagocytic capabilities of *C. farreri* hemocytes (Yang et al. 2011). CTLDC proteins may play a role as diversified PRRs and/or activators of the host defense response of the Eastern oyster to gram-negative bacteria.

The above three groups of proteins, DMBT1/SR type 12, C1qDC, and CTLDC were demonstrably diversified relative to other taxa, yet showed differential expression patterns in oysters in response to bacterial challenge that were not easily interpretable. For each group, while some transcripts were up-regulated following challenge, other transcripts were down-regulated. Several explanations for this apparent lack of consistency include dynamic expression regimes, as demonstrated in sea urchin SRs (Pancer 2000), complex/compensatory regulation as demonstrated in mussel C1qDC transcripts (Gerdol et al. 2011), variability of function within gene family, high
polymorphic rate, basal expression variability, which is high in many C. gigas genes (Rosa et al. 2011), and the intrinsic difficulty of mapping reads to a large gene family in a de novo-assembled transcriptome.

Interestingly, I have identified two families that have not been previously identified as diversified in other studies. To my knowledge, the present study is the first to identify GIMAP genes in invertebrates as diversified mediators of the invertebrate immune response. Previously, similarity searches for GIMAP genes within the genomes of fission yeast Saccharomyces pombe and brewer’s yeast S. cerevisiae, C. elegans, and D. melanogaster turned up no hits, and consequently, it was concluded that GIMAP genes are present only in vertebrates and angiosperms (Filén and Lahesmaa 2010). While no GIMAP proteins were found in the proteomes of two protostomes herein studied, D. melanogaster and D. pulex, GIMAP sequences where plentiful in molluscs (Fig. 5). My study advances the importance of including lophotrochozoans in genomic surveys for genes of interest, now enabled by the recent release of the Pacific oyster genome (Zhang et al. 2012). Phylogenetic analysis of GIMAP sequences supports the possibility of several gene expansion events (with an expanded set of GIMAP sequences from basal chordates and molluscs grouping together and expanded sets of GIMAP sequences unique to bivalves/molluscs) likely combined with diversification through alternative mechanisms (e.g. alternative splicing/INDELs/allelic variation/somatic diversification) (Fig. 9). In vertebrates, GIMAP proteins have been best
characterized in their role as regulators of apoptosis (Nitta et al. 2007), though it has been shown that GIMAP family members show differential expression patterns across tissue types and may serve varying functions at different times and in different tissues (Wang and Li 2009). Exposure of human monocytes to LPS induces the down-regulation of 28 genes by >4 fold, four of which are GIMAP proteins (Dower et al. 2008). It has been suggested that the down-regulation of GIMAP proteins in humans may serve to promote the survival of monocytes by negatively regulating apoptosis (Dower et al. 2008). It may be that GIMAP proteins fulfill a parallel role in oyster hemocytes though further work will be needed to define this role.

Another gene family proposed as a novel diversified mediator of the oyster immune response is the IFI44 gene family. IFI44, inducible by interferon-α, is implicated in antiviral host defense (Kitamura et al. 1994) and shows antiproliferative activity, possibly by contributing to cell cycle arrest (Hallen et al. 2007). IFI44 transcripts are up-regulated in C. gigas in response to challenge with the virus OsHV-1 (Renault et al. 2011) and a pathogenic Vibrio (de Lorgeril et al. 2011). Phylogenetic analysis of IFI44 sequences from diverse taxa supports the hypothesis of a bivalve-only gene expansion(s) likely combined with diversification through alternative mechanisms (e.g. alternative splicing/INDELs/allelic variation/somatic diversification) (Fig. 11). Further expression profiling on a finer time scale and in a variety of conditions may
help to determine why the IFI44 gene family has become diversified in bivalves and what specific challenges induce IFI44 expression.

Many unannotated gene families were certainly involved in the defense and/or stress response. Eighty-six of the 187 TribeMCL clusters that were found to be enriched for DEGs could not be annotated. The inability to annotate a great portion of the transcriptome (in this case 80% without BLAST similarity) remains a challenge in describing the oyster host defense response. I show here that similarity clustering does offer the means of transferring annotations. The majority of clusters that were herein annotated included one or more transcripts that could not be annotated by BLAST alone, yet whose identity could be inferred from its neighbors. Moreover, the very process of reducing a set of transcripts to a smaller set of connected components has the promise to focus efforts and resources in the effort to characterize genes and gene families that presently cannot be annotated by similarity search to the public databases. While these clusters could not be used to describe the oyster host defense response here, TribeMCL or an analogous clustering technique could be used to facilitate annotation in future studies. By considering transcripts at the level of TribeMCL clusters, sequence similarity motifs may be extracted to aid eventual characterization.

While much work remains to be done in characterizing the present Eastern oyster transcriptome and describing the oyster host defense response to bacterial challenge, the present study has made great advances to these
ends. When ROD-resistant and ROD-susceptible oysters were exposed *R. crassostreae* and gene expression was compared throughout the challenge by high-throughput cDNA sequencing, several processes emerged as key to resistance including ECM remodeling, cell adhesion, and inflammation. The present study has generated a pool of candidate disease resistance genes for advanced genotypic selection regimes. Additionally, several gene families were identified as putatively diversified and of immune relevance in the Eastern oyster, two of which, IFI44 and GIMAP families, are of especial interest as expansions were found to be specific to bivalves and molluscs, respectively. Transcript translation and similarity clustering followed by gene family analysis should prove useful in describing the transcriptomes of other invertebrates in response to immune and/or stress challenge as an unbiased means of identifying putatively diversified groups of host receptors, regulators, and effectors.
TABLES

Table 1. Assembly metrics for transcriptome assembly

| Metric                           | Value       |
|----------------------------------|-------------|
| Number of contigs                | 356237      |
| Total span (bp)                  | 156920694   |
| Number of contigs (> 1Kb)        | 22934       |
| Max Contig Length (bp)           | 16256       |
| Mean Contig Length (bp)          | 440         |
| N50 (bp)                         | 487         |
| Number of contigs with BLAST hits* | 70621   |
| % of contigs with BLAST hits*    | 19.8        |

*Contigs compared to NCBI's non-redundant protein database using BLASTX, significant hits retained with e-value ≤ 1e-06.
Table 2. Top 50 most highly differentially expressed genes (DEGs) unique to GX_early up-regulated relative to control pool (CGX_late)

| Contig          | Reg. | Reg. | logFC | log CPM | p-value | Best blastx hit to nr db | Hit accession | Hit e-value |
|-----------------|------|------|-------|---------|---------|--------------------------|---------------|-------------|
| comp17501_c0_seq1 | up   | -    | 6.1   | 4.55    | 2.01E-14| -                        | -             | -           |
| comp5950_c0_seq1 | up   | up   | 5.21  | 3.76    | 3.51E-12| -                        | -             | -           |
| comp4755_c0_seq2 | up   | -    | 5.13  | 2.95    | 1.03E-10| -                        | -             | -           |
| comp2875_c0_seq4 | up   | up   | 4.41  | 2.89    | 2.50E-09| -                        | -             | -           |
| comp1023_c0_seq2 | up   | up   | 4.28  | 3.85    | 5.46E-09| scavenger receptor cysteine-rich | ACT53266      | 1.30E-15    |
| comp24124_c0_seq1 | -   | up   | 4.49  | 2.59    | 9.31E-09| ched related family member (ptr-19) | XP_002734100 | 1.39E-138   |
| comp12059_c0_seq1 | up   | up   | 4.18  | 4.67    | 1.46E-08| -                        | -             | -           |
| comp1165_c0_seq2 | -   | up   | 3.99  | 4.75    | 1.48E-08| -                        | -             | -           |
| comp2015_c0_seq24 | up  | -    | 3.95  | 2.67    | 8.73E-08| inhibitor of apoptosis    | AEB54800      | 4.91E-09    |
| comp1023_c0_seq5 | up   | up   | 3.95  | 3.23    | 9.03E-08| -                        | -             | -           |
| comp2870_c0_seq2 | up   | up   | 3.79  | 4.29    | 1.07E-07| -                        | -             | -           |
| comp31762_c0_seq1 | up  | up   | 3.73  | 2.96    | 1.20E-07| -                        | -             | -           |
| comp657_c0_seq3  | -    | up   | 3.57  | 3.74    | 2.18E-07| -                        | -             | -           |
| comp3858_c0_seq5 | up   | -    | 3.63  | 2.82    | 6.54E-07| isoleucyl-tRNA synthetase | NP_001090690  | 2.18E-59    |
| comp9303_c0_seq3 | -    | up   | 3.67  | 1.49    | 8.37E-07| -                        | -             | -           |
| comp3628_c0_seq2 | up   | -    | 3.62  | 3.14    | 8.53E-07| -                        | -             | -           |
| comp7475_c2_seq3 | up   | -    | 3.66  | 1.75    | 8.71E-07| -                        | -             | -           |
| comp18756_c0_seq2 | up  | -    | 3.83  | 2.06    | 9.27E-07| sushi repeat-containing   | XP_002664481  | 2.19E-22    |
| comp20853_c1_seq3 | -   | up   | 3.57  | 2.16    | 1.19E-06| af397902_1egf-like       | XP_002601693  | 1.28E-35    |
| comp6834_c0_seq1 | -    | up   | 3.75  | 2       | 1.42E-06| Protein                  | XP_002592396  | 8.35E-12    |
| comp6161_c0_seq5 | up   | -    | 3.49  | 3.24    | 1.57E-06| type 2 proinsulin processing endopeptidase | 2206277A      | 2.33E-42    |
| comp1023_c0_seq1 | up   | up   | 3.53  | 3.08    | 1.67E-06| -                        | -             | -           |
| GenBank ID   | Description                          | Log2 Fold Change | p-Value  | FPKM  | Description                          | Log2 Fold Change | p-Value  | FPKM  |
|-------------|--------------------------------------|------------------|----------|-------|--------------------------------------|------------------|----------|-------|
| comp21651_c0_seq1 | up -                                | 3.58             | 2.33     | 2.23E-06 | -                                   | -                | -        | -     |
| comp4755_c0_seq1  | up -                                | 3.52             | 2.56     | 2.40E-06 | -                                   | -                | -        | -     |
| comp1316_c0_seq1   | up -                                | 3.26             | 4.6      | 2.79E-06 | -                                   | -                | -        | -     |
| comp1157_c0_seq3   | up up                               | 3.34             | 2.72     | 3.28E-06 | -                                   | -                | -        | -     |
| comp4626_c0_seq4   | up -                                | 3.34             | 2.55     | 3.32E-06 | alpha-ketoglutarate-dependent hypophosphite dioxygenase-like | XP_002944900     | 1.40E-10 |
| comp18756_c0_seq3  | up -                                | 3.48             | 2.61     | 4.03E-06 | fibropellin ia                       | XP_002601363     | 2.21E-13 |
| comp24671_c0_seq1  | - up                                | 3.57             | 1.99     | 4.38E-06 | -                                   | -                | -        | -     |
| comp5314_c0_seq1   | up up                               | 3.36             | 3.57     | 4.85E-06 | -                                   | -                | -        | -     |
| comp1788_c0_seq4   | - up                                | 3.26             | 2.21     | 5.26E-06 | fibrinolytic enzyme                  | CAA64472         | 5.10E-12 |
| comp22438_c0_seq1  | - up                                | 3.26             | 2.81     | 5.49E-06 | cytochrome p450 family 4            | ACM16804         | 4.94E-106|
| comp2015_c0_seq17  | up -                                | 3.23             | 3.14     | 6.25E-06 | -                                   | -                | -        | -     |
| comp7137_c0_seq2   | - up                                | 3.26             | 3.02     | 7.06E-06 | organic solute transporter subunit alpha | XP_002732822     | 4.93E-20 |
| comp6713_c0_seq2   | up up                               | 3.02             | 2.8      | 8.59E-06 | -                                   | -                | -        | -     |
| comp1506_c0_seq4   | up up                               | 3.08             | 5.47     | 1.01E-05 | a disintegrin and metalloproteina se with thrombospondin motifs 8 | XP_002940685     | 2.89E-12 |
| comp6837_c0_seq1   | up -                                | 3.15             | 4.45     | 1.16E-05 | interleukin 17d                      | A9XE49           | 1.05E-56 |
| comp18756_c0_seq5  | up -                                | 3.27             | 3.01     | 1.18E-05 | fibropellin ia                       | XP_002599260     | 2.50E-27 |
| comp3628_c0_seq3   | up -                                | 3.34             | 2.03     | 1.18E-05 | -                                   | -                | -        | -     |
| comp14520_c0_seq2  | up -                                | 3.21             | 2.31     | 1.26E-05 | -                                   | -                | -        | -     |
| comp274_c0_seq1    | - up                                | 3             | 6.82     | 1.29E-05 | -                                   | -                | -        | -     |
| comp4755_c0_seq3   | up -                                | 3.1              | 4.57     | 1.34E-05 | hypothetical protein                 | ACU33972         | 3.51E-35 |
| comp15440_c0_seq1  | up -                                | 3.18             | 1.8      | 1.34E-05 | inhibitor of apoptosis               | XP_002426441     | 1.01E-13 |
| comp11365_c0_seq2  | - up                                | 2.93             | 2.83     | 1.36E-05 | -                                   | -                | -        | -     |
| comp664_c0_seq5    | up up                               | 3.26             | 2.38     | 1.43E-05 | Protein                             | XP_001642030     | 4.44E-10 |

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The top fifty most differentially expressed and up-regulated genes unique to GX days 1 and 5 (genes not differentially expressed in any F3L treatment) are shown, ranked by false discovery rate-adjusted p-value. Magnitude of differential expression is expressed as log-10 fold change over control pool (logFC) and abundance is expressed as log-10 counts per million (logCPM). Regulation, or “Reg.”, (up- or down-regulated) is shown for each contig. Hyphen (-) indicates that the contig is not differentially expressed at that timepoint (Reg. columns) or that the contig does not have BLASTX hit to the NCBI non-redundant protein database with associated e-value $\leq 1e^{-6}$ (annotation columns). Where contigs are differentially expressed in both timepoints, logFC, logCPM, and p-value correspond to the timepoint in which the contig was most highly differentially expressed and that timepoint is indicated in the regulation columns in bold.

| contig          | Reg. | logFC | logCPM | p-value | BLASTX hit     |
|-----------------|------|-------|--------|---------|----------------|
| comp28783_c0_seq1 | -    | 3.1   | 2.08   | 1.52E-05 | BRAFLDRAFT_129258 |
| comp6161_c0_seq11 | up   | 3.1   | 3.16   | 1.59E-05 | XP_002612219   |
| comp9269_c0_seq3 | up   | 3.1   | 2.3    | 1.76E-05 | AAA49718       |
| comp1023_c0_seq3 | up   | 3.07  | 3.85   | 1.98E-05 | scavenger receptor cysteine-rich |
| comp22224_c0_seq1 | -    | 3.07  | 2.07   | 2.01E-05 | XP_001622238   |
Table 3. Differentially expressed genes (DEGs) unique to GX_early down-regulated relative to control pool

| Contig          | Reg. GX-1d | Reg. GX-5d | logFC | log CPM | p-value | Best blastx hit to nr db | Hit accession | Hit e-value |
|-----------------|------------|------------|-------|---------|---------|--------------------------|---------------|-------------|
| comp24428_c0_seq1 | down       | down       | -8.64 | 1.68    | 6.67E-07 | rapunzel 5              | NP_001103594  | 2.10E-10    |
| comp1572_c0_seq4 | -          | down       | -8.67 | 1.53    | 1.31E-06 | -                        | -             | -           |
| comp5722_c1_seq2 | -          | down       | -4.84 | 2.94    | 1.91E-06 | collagen alpha           | XP_001512734  | 2.87E-32    |
| comp14853_c0_seq2 | down      | down       | -8.29 | 1.28    | 3.49E-06 | -                        | -             | -           |
| comp1572_c0_seq3 | -          | down       | -4.34 | 4.24    | 5.84E-06 | -                        | -             | -           |
| comp24428_c0_seq1 | -          | down       | -8.09 | 1.08    | 7.32E-06 | cubilin                 | XP_00273392   | 0.00E+0      |
| comp869_c0_seq4 | down       | -          | -8.02 | 0.95    | 1.25E-05 | -                        | -             | -           |
| comp1572_c0_seq9 | -          | down       | -4.64 | 1.93    | 2.13E-05 | -                        | -             | -           |
| comp11408_c0_seq2 | -          | down       | -7.89 | 0.77    | 2.30E-05 | -                        | -             | -           |
| comp1285_c1_seq8 | -          | down       | -3.9  | 3.84    | 2.38E-05 | arginase type i-like     | AEB70965      | 5.87E-28    |
| comp3240_c1_seq2 | -          | down       | -4.27 | 1.99    | 4.15E-05 | -                        | -             | -           |
| comp25746_c0_seq4 | -          | down       | -7.6  | 0.55    | 5.20E-05 | tenascin xb              | XP_002741293  | 4.52E-38    |
| comp19167_c0_seq1 | -          | down       | -4.79 | 1.39    | 5.95E-05 | -                        | -             | -           |
| comp34093_c0_seq1 | -          | down       | -7.54 | 0.48    | 6.63E-05 | -                        | -             | -           |
| comp22172_c0_seq3 | down      | -          | -7.57 | 0.44    | 7.22E-05 | -                        | -             | -           |
| comp810_c1_seq1  | -          | down       | -3.4  | 6.65    | 8.89E-05 | heat shock protein 22    | ACU83231      | 2.79E-28    |
| comp1285_c1_seq3 | -          | down       | -3.54 | 3.50    | 9.56E-05 | arginase ii              | XP_002130834  | 6.53E-12    |
| comp39520_c0_seq1 | -          | down       | -4.31 | 1.47    | 1.30E-04 | polyprote in             | XP_0027 40782 | 0.00E+0      |
| comp10161_c0_seq1 | down       | -          | -3.47 | 3.97    | 1.41E-04 | -                        | -             | -           |
| comp9135_c0_seq1  | -          | down       | -3.54 | 3.17    | 1.59E-04 | -                        | -             | -           |
| comp1190_c0_seq7  | down       | -          | -3.44 | 3.60    | 2.04E-04 | -                        | -             | -           |
| comp38620_c0_seq1 | -          | down       | -7.26 | 0.16    | 2.14E-04 | -                        | -             | -           |
| comp2015_c0_seq13 | down | - | -7.19 | 0.12 | 2.29E-04 | inhibitor of apoptosis | AEB5479 9 | 1.28E-41 |
| comp43138_c0_seq1 | down | - | -7.22 | 0.12 | 2.29E-04 | - | - | - |
| comp5608_c0_seq1 | down | -3.35 | 3.50 | 2.31E-04 | c-type lectin | ABB7167 2 | 7.01E-16 |
| comp16886_c0_seq1 | down | -7.17 | 0.11 | 2.36E-04 | - | - | - |
| comp18757_c0_seq1 | down | -7.14 | 0.09 | 2.36E-04 | hla-b associated transcript 1 | XP_0032 17350 | 6.91E-21 |
| comp1190_c0_seq5 | down | down | -3.25 | 4.38 | 2.48E-04 | - | - | - |
| comp7972_c0_seq4 | down | -3.11 | 4.03 | 3.13E-04 | cubilin | XP_0026 12977 | 0.00E+00 |
| comp7814_c1_seq2 | down | -7 | 0.04 | 3.58E-04 | - | - | - |
| comp16058_c0_seq1 | down | -3.9 | 1.18 | 3.76E-04 | - | - | - |
| comp24625_c0_seq1 | down | -7.09 | - | 0.03 | 3.81E-04 | - | - | - |
| comp5396_c0_seq1 | down | -3.01 | 3.96 | 4.75E-04 | melatonin receptor 1a | ADM731 75 | 1.28E-66 |
| comp27900_c0_seq1 | down | -4.2 | 0.79 | 5.18E-04 | - | - | - |
| comp1190_c0_seq8 | down | -3.1 | 3.94 | 5.22E-04 | - | - | - |
| comp16567_c0_seq1 | down | -4.58 | 0.39 | 5.85E-04 | - | - | - |
| comp5722_c1_seq1 | down | -2.86 | 4.28 | 6.15E-04 | collagen alpha | XP_0015 12734 | 3.63E-32 |
| comp13269_c0_seq1 | down | -3.06 | 3.19 | 6.31E-04 | DAPPUD RAFT 3 09315 | EFX7073 7 | 1.87E-17 |
| comp18902_c0_seq1 | down | -3.43 | 1.62 | 6.49E-04 | rho gtpase | XP_0027 39105 | 1.37E-58 |
| comp17170_c0_seq1 | down | -3.42 | 1.63 | 6.75E-04 | - | - | - |
| comp19167_c0_seq2 | down | -3.48 | 1.42 | 7.44E-04 | - | - | - |
| comp1285_c1_seq1 | down | -4.44 | 0.25 | 9.04E-04 | - | - | - |
| comp7712_c0_seq2 | down | -3.36 | 1.33 | 1.07E-03 | - | - | - |
| comp14853_c0_seq8 | down | -3.34 | 1.12 | 1.09E-03 | - | - | - |
Differentially expressed contigs unique to GX days 1 and 5 (genes not differentially expressed in any F3L treatment) that were down-regulated are shown, ranked by false discovery rate-adjusted p-value. Magnitude of differential expression is expressed as log-10 fold change over control pool (logFC) and abundance is expressed as log-10 counts per million (logCPM). Regulation, or “Reg.”, (up- or down-regulated) is shown for each contig. ( - ) indicates that the contig is not differentially expressed at that timepoint (Reg. columns) or that the contig does not have BLASTX hit to the NCBI non-redundant protein database with associated e-value ≤ 1e-6. Where contigs are differentially expressed in both timepoints, logFC, logCPM, and p-value correspond to the timepoint in which the contig was most highly differentially expressed and that timepoint is indicated in the regulation columns in bold.
Table 4. Top 50 most highly differentially expressed genes (DEGs) among GX_early shared with F3L_early and/or F3L_late up-regulated relative to control pool

| Contig          | Reg. GX-1d | Reg. GX-5d | logFC | log CPM | p-value | Best blastx hit to nr db  | Hit accession | Hit e-value |
|-----------------|------------|------------|-------|---------|---------|---------------------------|---------------|-------------|
| comp866_c0_seq1 | -          | up         | 5.43  | 4.01    | 1.11E-12| BRAFLDRAFT_227853         | XP_002612894  | 6.68E-31    |
| comp619_c0_seq1 | up         | up         | 4.84  | 5.8     | 2.34E-11| serine protease inhibitor cvsi-2| B9A8D7        | 1.55E-11    |
| comp1523_c0_seq2| -          | up         | 4.67  | 4.37    | 5.40E-11| -                         | -             | -           |
| comp2870_c0_seq3| up         | up         | 4.82  | 3.88    | 1.19E-10| predicted protein          | XP_001632962  | 3.82E-17    |
| comp2870_c0_seq1| up         | up         | 4.34  | 5.13    | 1.97E-09| hypothetical protein       | XP_002416173  | 2.97E-59    |
| comp11276_c0_seq3| -          | up         | 4.67  | 2.68    | 2.88E-09| Polyketide synthase pk2    | XP_002734101  | 3.26E-44    |
| comp985_c0_seq1 | -          | up         | 4.34  | 4.38    | 3.18E-09| -                         | -             | -           |
| comp3562_c1_seq1| up         | -          | 4.27  | 3.64    | 3.21E-09| -                         | -             | -           |
| comp631_c0_seq2 | -          | up         | 4.09  | 5.3     | 3.99E-09| dopamine beta hydroxylase-like | XP_002117559 | 1.97E-08    |
| comp9303_c0_seq5| -          | up         | 4.17  | 2.48    | 9.67E-09| omega class glutathione s-transferase | CAD89618      | 3.39E-13    |
| comp1165_c0_seq1| -          | up         | 4.07  | 5.51    | 9.80E-09| -                         | -             | -           |
| comp928_c0_seq1 | -          | up         | 3.92  | 6.29    | 1.93E-08| serine protease inhibitor cvsi-2 | B9A8D7        | 3.21E-12    |
| comp300_c0_seq1 | -          | up         | 3.87  | 8.08    | 2.17E-08| -                         | -             | -           |
| comp1165_c0_seq3| -          | up         | 3.89  | 6.63    | 3.08E-08| -                         | -             | -           |
| comp1893_c0_seq1| up         | up         | 3.95  | 4.01    | 4.61E-08| dopamine beta hydroxylase-like | AAS92605      | 3.53E-27    |
| comp2875_c0_seq2| -          | up         | 3.9   | 3.61    | 5.23E-08| serine protease            | XP_002593726  | 2.71E-08    |
| comp8625_c0_seq1| up         | -          | 4.04  | 3.44    | 6.69E-08| fatty acid synthase-like   | ACZ55138      | 0.00E+0      |
| comp298_c0_seq1 | up         | up         | 3.74  | 7.91    | 8.70E-08| -                         | -             | -           |
| Gene         | Type | Log2FC | Log2|   |   |   |
|-------------|------|--------|-----|---|---|---|
| comp2550_c0_seq1 | up   | 3.83   | 4.73 | 9.59E-08 - | - | - |
| comp3607_c0_seq3 | -    | 3.88   | 3.27 | 1.09E-07 arylsulfatase-like | XP_002607295 | 1.00E-12 |
| comp11276_c0_seq8 | -    | 4.12   | 2.19 | 1.48E-07 - | - | - |
| comp631_c0_seq1   | -    | 3.58   | 6.81 | 1.60E-07 dopamine beta hydroxylase-like | XP_002117561 | 2.29E-31 |
| comp1880_c0_seq2  | up   | 3.85   | 3.59 | 1.62E-07 - | - | - |
| comp281_c1_seq1   | -    | 3.54   | 8.32 | 1.78E-07 - | - | - |
| comp7186_c1_seq4  | up   | -      | 3.92 | 2.39 | 2.12E-07 dna damage-regulated autophagy modulator protein 2 | NP_001230625 | 1.82E-14 |
| comp1199_c0_seq1  | up   | -      | 3.74 | 4.66 | 2.23E-07 - | - | - |
| comp335_c0_seq1   | -    | 3.6    | 5.15 | 3.26E-07 - | - | - |
| comp12125_c0_seq1 | -    | 3.49   | 3.4  | 4.15E-07 - | - | - |
| comp1479_c0_seq1  | up   | 3.67   | 3.44 | 5.43E-07 - | - | - |
| comp1199_c1_seq2  | up   | -      | 3.57 | 5.05 | 6.03E-07 - | - | - |
| comp1853_c0_seq4  | -    | 3.57   | 3.25 | 7.21E-07 - | - | - |
| comp985_c0_seq2   | -    | 3.51   | 4.9  | 7.43E-07 - | - | - |
| comp7828_c0_seq1  | up   | -      | 3.63 | 2.6  | 7.76E-07 - | - | - |
| comp1199_c1_seq3  | up   | -      | 3.52 | 5.31 | 9.49E-07 - | - | - |
| comp1037_c0_seq1  | -    | 3.27   | 6.27 | 1.28E-06 - | - | - |
| comp985_c0_seq3   | -    | 3.39   | 3.06 | 1.38E-06 - | - | - |
| comp88_c0_seq1    | up   | 3.4    | 2.85 | 1.79E-06 - | - | - |
| comp88_c0_seq2    | up   | 3.4    | 2.85 | 1.79E-06 - | - | - |
| comp88_c0_seq3    | up   | 3.4    | 2.85 | 1.79E-06 - | - | - |
| comp88_c0_seq4    | up   | 3.4    | 2.85 | 1.79E-06 - | - | - |
| comp1197_c1_seq1  | -    | 3.4    | 5.68 | 2.07E-06 - | - | - |
| comp887_c0_seq1   | up   | 3.32   | 5.73 | 2.13E-06 c1q domain containing protein 1q13 | CBX41662 | 1.54E-07 |
| comp437_c0_seq1   | up   | 3.41   | 5.69 | 2.20E-06 - | - | - |
| comp437_c0_seq2   | up   | 3.35   | 5.33 | 2.30E-06 - | - | - |
The top fifty most differentially expressed and up-regulated genes shared between GX days 1 and 5 and F3L days 1, 5, 15, and/or 30 are shown, ranked by false discovery rate-adjusted p-value. Magnitude of differential expression is expressed as log-10 fold change over control pool (logFC) and abundance is expressed as log-10 counts per million (logCPM). Regulation, or “Reg.”, (up- or down-regulated) is shown for each contig. Hyphen (-) indicates that the contig is not differentially expressed at that timepoint (Reg. columns) or that the contig does not have BLASTX hit to the NCBI non-redundant protein database with associated e-value ≤ 1e-6 (annotation columns). Where contigs are differentially expressed in both timepoints, logFC, logCPM, and p-value correspond to the timepoint in which the contig was most highly differentially expressed and that timepoint is indicated in the regulation columns in bold.

| Contig ID | Regulation | LogFC | LogCPM | P-value | Gene Name | Accession | LogCPM | P-value |
|-----------|------------|-------|--------|---------|-----------|-----------|--------|---------|
| comp1229_c0_seq1 | - up | 3.16 | 4.59 | 2.59E-06 | - | - | - |
| comp11715_c0_seq1 | - up | 3.4 | 2.6 | 2.63E-06 | - | - | - |
| comp4943_c0_seq1 | - up | 3.35 | 3.49 | 3.04E-06 | Cytochrome p450 | XP_002594971 | 1.92E-58 |
| comp2451_c0_seq15 | - up | 3.59 | 1.94 | 3.19E-06 | Galactosamine (n-acetyl)-6-sulfate sulfatase-like | XP_002605064 | 5.12E-16 |
| comp3498_c0_seq2 | up - | 3.18 | 4.32 | 3.45E-06 | Heat shock protein 60 | ABN11936 | 7.59E-83 |
| comp186_c0_seq1 | up | 3.25 | 5.44 | 3.57E-06 | - | - | - |
Table 5. Top 50 most highly differentially expressed genes (DEGs) among GX_early DEGs shared with F3L_early and/or F3L_late down-regulated relative to control pool

| Contig            | Reg. GX-1d | Reg. GX-5d | logFC | log CPM | p-value | Best blastx hit to nr db | Hit accession | Hit e-value |
|-------------------|------------|------------|-------|---------|---------|--------------------------|---------------|-------------|
| comp12483_c0_seq1 | down       | -11.19     | 4.06  | 3.29E-11|         | c1q domain containing protein 1q13 | CBX41662     | 4.49E-09    |
| comp1102_c0_seq1  | -          | -7.59      | 4.92  | 7.33E-11| -       | -                        | -             | -           |
| comp9636_c0_seq2  | -          | -10.66     | 3.55  | 2.99E-10| -       | -                        | -             | -           |
| comp1102_c0_seq3  | -          | -6.92      | 5.29  | 3.29E-10|         | collagen alpha-5 chain     | XP_002596170 | 1.40E-08    |
| comp6091_c0_seq2  | -          | -10.58     | 3.45  | 4.66E-10|         | camp responsive element binding 2 | AAU93879     | 3.04E-18    |
| comp14386_c0_seq1 | -          | -7.3       | 4.42  | 6.49E-10| -       | -                        | -             | -           |
| comp3971_c0_seq2  | -          | -6.02      | 5.04  | 7.78E-09|         | monocarboxylate transporter 14 | EGI68511     | 5.46E-27    |
| comp9636_c0_seq6  | -          | -9.83      | 2.70  | 1.11E-08| -       | -                        | -             | -           |
| comp8174_c0_seq1  | down       | -6.45      | 3.28  | 3.28E-08| -       | -                        | -             | -           |
| comp10675_c0_seq1 | -          | -6.02      | 3.86  | 4.08E-08| -       | -                        | -             | -           |
| comp30091_c0_seq2 | -          | -9.22      | 2.10  | 1.33E-07|         | Nudt9                    | EGD73755     | 1.12E-27    |
| comp35662_c0_seq1 | -          | -8.93      | 1.97  | 2.23E-07|         | Pol                      | XP_786277     | 0.00E+00    |
| comp13170_c1_seq3 | -          | -9.02      | 1.96  | 2.30E-07|         | deleted in malignant brain tumors 1 | XP_002833280 | 4.81E-30    |
| comp513_c0_seq3   | -          | -8.98      | 1.93  | 2.64E-07| -       | -                        | -             | -           |
| comp3971_c0_seq4  | -          | -4.74      | 5.33  | 5.66E-07|         | monocarboxylate transporter | XP_002573719 | 1.50E-17    |
| comp7347_c0_seq1  | -          | -6.41      | 2.21  | 5.79E-07| -       | -                        | -             | -           |
| comp23074_c0_seq1 | -          | -5.9       | 2.53  | 6.43E-07| -       | -                        | -             | -           |
| comp35580_c0_seq2 | -          | -8.66      | 1.57  | 1.15E-06| -       | -                        | -             | -           |
| comp3971_c0_seq1  | -          | -8.56      | 1.52  | 1.31E-06|         | monocarboxylate transporter | XP_00160814  | 2.19E-26    |
| comp5775_c0_seq3  | -          | -5.3       | 2.59  | 1.47E-06| -       | -                        | -             | -           |
| comp11368_c0_seq1 | -          | -6.16      | 1.88  | 2.26E-06| -       | -                        | -             | -           |
| comp9636_c0_seq8 | - down     | -5.57 | 2.14 | 3.08E-06 | - | - |
|-----------------|------------|-------|------|----------|---|---|
| comp14613_c0_seq2 | down down | -5.06 | 2.12 | 3.59E-06 | - | - |
| comp50794_c0_seq1 | - down | -8.23 | 1.15 | 5.51E-06 | novel protein human megf11 | EGW0405 | 1.99E-25 |
| comp3971_c0_seq5 | - down | -4.6 | 3.13 | 5.80E-06 | - | - |
| comp33670_c0_seq1 | - down | -8.23 | 1.10 | 6.91E-06 | - | - |
| comp30670_c0_seq1 | down down | -8.15 | 1.05 | 8.24E-06 | - | - |
| comp10350_c0_seq1 | - down | -8.17 | 1.04 | 8.74E-06 | c-type lectin 2 | XP_002603342 | 5.30E-10 |
| comp11802_c0_seq2 | - down | -8.12 | 1.03 | 8.74E-06 | - | - |
| comp869_c0_seq2 | - down | -4 | 6.49 | 9.60E-06 | x-box binding | XP_002732738 | 1.19E-07 |
| comp5787_c0_seq1 | - down | -7.99 | 0.98 | 1.05E-05 | - | - |
| comp6966_c1_seq2 | - down | -8.04 | 0.98 | 1.05E-05 | - | - |
| comp30235_c0_seq3 | down - | -8.14 | 0.97 | 1.17E-05 | - | - |
| comp47716_c0_seq1 | - down | -8.01 | 0.95 | 1.19E-05 | - | - |
| comp5396_c0_seq2 | - down | -3.91 | 4.71 | 1.25E-05 | melatonin receptor 1a | ADM73175 | 2.20E-58 |
| comp10976_c1_seq1 | - down | -4.41 | 2.80 | 1.26E-05 | - | - |
| comp55655_c0_seq1 | down - | -8.1 | 0.94 | 1.32E-05 | - | - |
| comp908_c1_seq3 | - down | -4.15 | 3.53 | 1.43E-05 | - | - |
| comp25817_c0_seq1 | - down | -7.89 | 0.83 | 1.87E-05 | protein tyrosine phosphatase | ACH42087 | 2.41E-30 |
| comp6700_c2_seq2 | - down | -3.88 | 5.01 | 2.15E-05 | - | - |
| comp27010_c0_seq1 | down - | -5.51 | 1.27 | 2.45E-05 | - | - |
| comp7978_c0_seq1 | down - | -4.37 | 2.63 | 2.63E-05 | - | - |
| comp28180_c0_seq1 | - down | -4.91 | 1.59 | 2.69E-05 | sushi-repeat-containing x-linked 2 | XP_002932840 | 3.55E-16 |
| comp908_c1_seq2 | - down | -3.85 | 4.36 | 2.82E-05 | - | - |
| comp14067_c0_seq3 | - down | -5.28 | 1.23 | 2.82E-05 | - | - |
| comp908_c1_seq1 | - down | -3.75 | 4.90 | 2.98E-05 | - | - |
| comp18320_c0_seq1 | down down | -3.93 | 4.06 | 3.30E-05 | - | - |
The top fifty most differentially expressed and down-regulated genes shared between GX days 1 and 5 and F3L days 1, 5, 15, and/or 30 are shown, ranked by false discovery rate-adjusted p-value. Magnitude of differential expression is expressed as log-10 fold change over control pool (logFC) and abundance is expressed as log-10 counts per million (logCPM). Regulation, or “Reg.”, (up- or down-regulated) is shown for each contig. Hyphen (-) indicates that the contig is not differentially expressed at that timepoint (Reg. columns) or that the contig does not have BLASTX hit to the NCBI non-redundant protein database with associated e-value $\leq 1e^{-6}$ (annotation columns). Where contigs are differentially expressed in both timepoints, logFC, logCPM, and p-value correspond to the timepoint in which the contig was most highly differentially expressed and that timepoint is indicated in the regulation columns in bold.
Table 6. Pfam families enriched among the differentially expressed genes (DEGs) from resistant (GX) and susceptible (F3L) oysters at early (1 and 5 d) and late (15 and 30 d) timepoints compared to control (CGX) oysters, with rank order of significance of enrichment and significance of enrichment (p-value).

| Pfam families | Rank order of significance of enrichment in F3L_early | p-value of enrichment, adjusted by FDR correction, F3L_early | Rank order of significance of enrichment in F3L_late | p-value of enrichment, adjusted by FDR correction, F3L_late | Rank order of significance of enrichment in GX_early | p-value of enrichment, adjusted by FDR correction, GX_early |
|---------------|-----------------------------------------------------|------------------------------------------------------------|----------------------------------------------------|------------------------------------------------------------|----------------------------------------------------|------------------------------------------------------------|
| TSP_1         | 1                                                   | 4.16E-12                                                   | 13                                                 | 0.00598                                                   | 10                                                 | 0.01299                                                   |
| Acyl_transf_3 | 2                                                   | 1.87E-06                                                   | 9                                                  | 0.00079                                                   | -                                                  | -                                                         |
| AIG1          | 3                                                   | 1.90E-05                                                   | 2                                                  | 1.44E-09                                                   | 11                                                 | 0.02949                                                   |
| MFS_1         | 4                                                   | 0.00011                                                   | -                                                  | -                                                          | 3                                                  | 0.00156                                                   |
| RVT_1         | 5                                                   | 0.00011                                                   | 6                                                  | 0.00053                                                   | -                                                  | -                                                         |
| TIMP          | 6                                                   | 0.00013                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Perilipin     | 7                                                   | 0.00018                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| SSF           | 8                                                   | 0.00019                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Sugar_tr      | 9                                                   | 0.00023                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| An_peroxidase | 10                                                  | 0.00033                                                   | 3                                                  | 1.87E-05                                                   | -                                                  | -                                                         |
| Pacifastin_1  | 11                                                  | 0.00104                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| GlcNAc_2-epim | 12                                                  | 0.00194                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Transposase_21| 13                                                  | 0.00194                                                   | 4                                                  | 5.35E-05                                                   | -                                                  | -                                                         |
| VWO           | 14                                                  | 0.00194                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| SNF           | 15                                                  | 0.00211                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Ank_2         | 16                                                  | 0.00222                                                   | 1                                                  | 8.50E-11                                                   | -                                                  | -                                                         |
| 7tm_1         | 17                                                  | 0.00243                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Nucleoplasmin | 18                                                  | 0.00271                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| T2SE          | 19                                                  | 0.00271                                                   | 7                                                  | 0.00076                                                   | -                                                  | -                                                         |
| Dam           | 20                                                  | 0.00480                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Man-6-P_recep | 21                                                  | 0.00612                                                   | 10                                                 | 0.00079                                                   | -                                                  | -                                                         |
| Pro_dh        | 22                                                  | 0.00612                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| TauD          | 23                                                  | 0.00790                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Dynamin_N     | 24                                                  | 0.00847                                                   | 11                                                 | 0.00092                                                   | -                                                  | -                                                         |
| AA_permease_2 | 25                                                  | 0.01120                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Methyltransf_FA| 26                                                 | 0.01123                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| NAD_birning_5 | 27                                                  | 0.01123                                                   | -                                                  | -                                                          | 1                                                  | 0.00069                                                   |
| Pyridoxal_dec | 28                                                  | 0.01440                                                   | 16                                                 | 0.01161                                                   | -                                                  | -                                                         |
| Sulfate_transp| 29                                                  | 0.01592                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| Cpn60_TCP1    | 30                                                  | 0.01692                                                   | -                                                  | -                                                          | 2                                                  | 0.00156                                                   |
| Peptidase_M13_N| 31                                                 | 0.02113                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| HTH_Tnp_Tc3_2 | 32                                                  | 0.02465                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| ApoL          | 33                                                  | 0.02787                                                   | 5                                                  | 0.00053                                                   | -                                                  | -                                                         |
| Gal_Lectin    | 34                                                  | 0.02841                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| zf-TAZ        | 35                                                  | 0.02841                                                   | 15                                                 | 0.00819                                                   | -                                                  | -                                                         |
| A2M           | 36                                                  | 0.03380                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| GCC2_GCC3     | 37                                                  | 0.04229                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| HSP70         | 38                                                  | 0.04229                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| KR            | 39                                                  | 0.04229                                                   | -                                                  | -                                                          | -                                                  | -                                                         |
| VOMI          | -                                                   | -                                                          | 8                                                  | 0.00076                                                   | -                                                  | -                                                         |
| CHB_HEX_C_1   | -                                                   | -                                                          | 12                                                 | 0.00299                                                   | -                                                  | -                                                         |
| RPE65         | -                                                   | -                                                          | 14                                                 | 0.00819                                                   | -                                                  | -                                                         |
| DDE_1         | -                                                   | -                                                          | 17                                                 | 0.02056                                                   | -                                                  | -                                                         |
| Mucin2_WxxW   | -                                                   | -                                                          | 18                                                 | 0.02422                                                   | -                                                  | -                                                         |
| MMR_HSR1      | -                                                   | -                                                          | 19                                                 | 0.03240                                                   | -                                                  | -                                                         |
| Peptidase_M84 | -                                                   | -                                                          | 20                                                 | 0.04388                                                   | -                                                  | -                                                         |
| zf-MYND       | -                                                   | -                                                          | 21                                                 | 0.04388                                                   | -                                                  | -                                                         |
| Pfam accession | Count | Significance p-value |
|----------------|-------|---------------------|
| Solute_trans_a | 4     | 0.00156             |
| IL17           | 5     | 0.00206             |
| HSP20          | 6     | 0.00355             |
| bZIP_2         | 7     | 0.01033             |
| Patched        | 8     | 0.01299             |
| Sulfatase      | 9     | 0.01299             |
| HSP70          | 12    | 0.02965             |
| GCC2_GCC3      | 13    | 0.04324             |

Pearson’s chi-squared test was performed to test for the enrichment of each unique HMM accession. If the expected count for any cell in the 2x2 contingency table was 5 or fewer, Fisher’s exact test was performed instead. Significance p-values listed above were adjusted for the number of independent enrichment tests performed for each group of DEGs by the False Discovery Rate correction method. Pfam families are ordered above by (1) rank order of significance among F3L_early DEGs, (2) rank order of significance among F3L_late DEGs, then (3) rank order of significance among GX_early DEGs.
Table 7. Pfam domains enriched among the differentially expressed genes (DEGs) from resistant (GX) and susceptible (F3L) oysters at early (1 and 5 d) and late (15 and 30 d) timepoints compared to control (CGX) oysters, with rank order of significance of enrichment and significance of enrichment (p-value)

| Pfam domains       | Rank order of significance of enrichment in F3L_early (days 1, 5) | p-value of enrichment, adjusted by FDR correction, F3L_early | Rank order of significance of enrichment in F3L_late (days 15, 30) | p-value of enrichment, adjusted by FDR correction, F3L_late | Rank order of significance of enrichment in GX_early (days 1, 5) | p-value of enrichment, adjusted by FDR correction, GX_early |
|--------------------|---------------------------------------------------------------|-------------------------------------------------------------|-----------------------------------------------------------------|-------------------------------------------------------------|---------------------------------------------------------------|-------------------------------------------------------------|
| C1q                | 1                                                             | 5.51E-53                                                    | 2                                                              | 1.57E-70                                                    | 4                                                             | 0.00761                                                     |
| Fibrinogen_C       | 2                                                             | 2.86E-24                                                    | 8                                                              | 1.22E-15                                                    | -                                                             | -                                                          |
| Lectin_C           | 3                                                             | 1.67E-21                                                    | 13                                                             | 3.81E-07                                                    | -                                                             | -                                                          |
| VWA                | 4                                                             | 6.91E-14                                                    | -                                                              | -                                                           | -                                                             | -                                                          |
| ShK                | 5                                                             | 8.62E-09                                                    | 17                                                             | 6.59E-06                                                    | -                                                             | -                                                          |
| Kazal_1            | 6                                                             | 8.29E-07                                                    | -                                                              | -                                                           | -                                                             | -                                                          |
| SRCR               | 7                                                             | 2.35E-06                                                    | 18                                                             | 0.00555                                                     | 15                                                            | 0.01437                                                     |
| Kunitz_BPTI        | 8                                                             | 8.56E-06                                                    | 12                                                             | 0.00770                                                     | 12                                                            | 0.03973                                                     |
| Chitin_bind_3      | 9                                                             | 6.40E-05                                                    | -                                                              | -                                                           | 6                                                             | 0.01029                                                     |
| Cu2_monooxygen     | 10                                                            | 0.00014                                                     | 4                                                              | 6.59E-06                                                    | -                                                             | -                                                          |
| E1_DerF2_DerF2     | 11                                                            | 0.00101                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| BBF                | 12                                                            | 0.00251                                                     | 1                                                              | 0.00019                                                     | -                                                             | -                                                          |
| Cu-oxidase         | 13                                                            | 0.00251                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Defensin_2         | 14                                                            | 0.00251                                                     | 6                                                              | 0.00011                                                     | -                                                             | -                                                          |
| Sulfotransfer_1    | 15                                                            | 0.00313                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| CM_14              | 16                                                            | 0.00362                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| p450               | 17                                                            | 0.00362                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| GOLD_2             | 18                                                            | 0.00438                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Kazal_2            | 19                                                            | 0.00438                                                     | 11                                                             | 0.03974                                                     | -                                                             | -                                                          |
| Sulfate_tra_GLY    | 20                                                            | 0.00438                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Cystatin           | 21                                                            | 0.00688                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| HYR                | 22                                                            | 0.00751                                                     | 10                                                             | 0.01121                                                     | 10                                                            | 0.00419                                                     |
| Thioredoxin_4      | 23                                                            | 0.00846                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| T1a                | 24                                                            | 0.00846                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Cu2_monoox_C       | 25                                                            | 0.01419                                                     | 3                                                              | 0.00301                                                     | -                                                             | -                                                          |
| Glyco_hydro_9      | 26                                                            | 0.01419                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| PAX                | 27                                                            | 0.01419                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| DOMON              | 28                                                            | 0.01976                                                     | 7                                                              | 0.00770                                                     | -                                                             | -                                                          |
| CM_4_9             | 29                                                            | 0.02046                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Lipase             | 30                                                            | 0.02509                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| adh_short_C2       | 31                                                            | 0.02823                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| F5_F8_type_C       | 32                                                            | 0.02823                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| GBP                | 33                                                            | 0.03023                                                     | 9                                                              | 0.01014                                                     | -                                                             | -                                                          |
| Cu-oxidase_2       | 34                                                            | 0.03302                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| PBP                | 35                                                            | 0.03302                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| T-box              | 36                                                            | 0.03553                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Tyrosinase         | 37                                                            | 0.03784                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| CRAL_TRIO          | 38                                                            | 0.03850                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Trypsin            | 39                                                            | 0.03850                                                     | -                                                              | -                                                           | 16                                                            | 0.00060                                                     |
| Transglut_N        | 40                                                            | 0.04209                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| FTCD               | 41                                                            | 0.04469                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Ras                | 42                                                            | 0.04469                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| DUF4218            | 43                                                            | 0.04731                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| EGF_3              | 44                                                            | 0.04731                                                     | -                                                              | -                                                           | -                                                             | -                                                          |
| Cupin_8            | -                                                             | -                                                            | 5                                                              | 0.02629                                                     | -                                                             | -                                                          |
Pearson’s chi-squared test was performed to test for the enrichment of each unique HMM accession. If the expected count for any cell in the 2x2 contingency table was 5 or fewer, Fisher’s exact test was performed instead. Significance p-values listed above were adjusted for the number of independent enrichment tests performed for each group of DEGs by the Bonferroni correction method. Pfam domains are ordered above by (1) rank order of significance among F3L_early DEGs, (2) rank order of significance among F3L_late DEGs, then (3) rank order of significance among GX_early DEGs.

| Domain             | Rank | Significance | p-value  |
|--------------------|------|--------------|----------|
| Myb_DNA-bind_4     | 14   | 0.03888      | -        |
| rve                | 15   | 0.00084      | -        |
| RVT_3              | 16   | 0.01014      | -        |
| Acyl_transf_1      | -    | -            | -        |
| Arginase           | -    | -            | -        |
| BIR                | -    | -            | -        |
| cEGF               | -    | -            | -        |
| CUB                | -    | -            | -        |
| EF_hand_5          | -    | -            | -        |
| H_lectin           | -    | -            | -        |
| ketoacyl-synt      | -    | -            | -        |
| MAM                | -    | -            | -        |
| Peptidase_S8       | -    | -            | -        |

**Significance p-values:**
- Myb_DNA-bind_4: 0.03888
- rve: 0.00084
- RVT_3: 0.01014
- Acyl_transf_1: -
- Arginase: -
- BIR: -
- cEGF: -
- CUB: -
- EF_hand_5: -
- H_lectin: -
- ketoacyl-synt: -
- MAM: -
- Peptidase_S8: -
Table 8. TribeMCL clusters enriched for differentially expressed genes (DEGs) from resistant (GX) and susceptible (F3L) oysters at early (1 and 5 d) and late (15 and 30 d) timepoints, with rank order of significance of enrichment and significance of enrichment (p-value)

| Tribe MCL cluster | # contigs in cluster | Annotation                                               | Rank order p-value F3L_early | Enrich. p-value F3L_early | Rank order p-value F3L_late | Enrich. p-value F3L_late | Rank order p-value GX_early | Enrich. p-value GX_early |
|-------------------|----------------------|----------------------------------------------------------|------------------------------|--------------------------|----------------------------|--------------------------|-----------------------------|---------------------------|
| 15                | 214                  | c-type lectin                                            | 1                           | 3.44E-28                 | 17                         | 2.49E-06                 | 17                          | 0.00713                   |
| 33                | 150                  | c1q domain containing protein                            | 2                           | 3.70E-22                 | 9                          | 2.32E-08                 | -                           | -                         |
| 19                | 196                  | unknown                                                  | 3                           | 3.14E-13                 | 1                          | 2.01E-14                 | -                           | -                         |
| 35                | 144                  | fibrinogen domain-containing*                            | 4                           | 3.87E-11                 | 2                          | 7.92E-11                 | -                           | -                         |
| 27                | 155                  | hemicentin/rhamnospondin/trombospondin*                   | 5                           | 6.43E-10                 | 26                         | 2.31E-05                 | 5                           | 0.00052                   |
| 535               | 16                   | peritrophin*                                             | 6                           | 7.87E-10                 | 41                         | 0.00064                  | -                           | -                         |
| 1084              | 9                    | vdg3                                                     | 7                           | 7.93E-10                 | 49                         | 0.00163                  | -                           | -                         |
| 493               | 17                   | dopamine beta hydroxylase                                 | 8                           | 1.39E-09                 | 4                          | 1.32E-10                 | -                           | -                         |
| 31                | 151                  | scavenger receptor cysteine-rich protein type 12/deleted in malignant brain tumors 1 | 9                           | 2.81E-09                 | 34                         | 0.00038                  | -                           | -                         |
| 179               | 40                   | c1q domain containing protein                            | 10                          | 1.36E-08                 | 3                          | 1.32E-10                 | -                           | -                         |
| 532               | 16                   | nose resistant to fluoxetine family member (nrf-6)        | 11                          | 1.78E-08                 | 40                         | 0.00064                  | -                           | -                         |
| 165               | 43                   | unknown                                                  | 12                          | 3.13E-08                 | 5                          | 4.88E-09                 | -                           | -                         |
| 32                | 151                  | unknown                                                  | 13                          | 3.63E-08                 | 6                          | 4.88E-09                 | -                           | -                         |
| 1987              | 6                    | unknown                                                  | 14                          | 4.07E-08                 | -                          | -                        | -                           | -                         |
| 325               | 24                   | c1q domain containing protein                            | 15                          | 6.03E-08                 | 24                         | 2.17E-05                 | -                           | -                         |
| 55                | 106                  | IgGFc-binding protein-like /ig-like domain-containing /fc fragment of binding protein* | 16                          | 6.22E-08                 | 52                         | 0.00193                  | -                           | -                         |
| 45                | 124                  | collagen alpha-1/3/4/5                                    | 17                          | 1.51E-07                 | -                          | -                        | -                           | -                         |
| 1416              | 7                    | unknown                                                  | 18                          | 2.15E-07                 | 25                         | 2.31E-05                 | 10                          | 0.00059                   |

76
| 50 | 112 | unknown   | 19 | 4.48E-06 | 8 | 2.32E-08 | - | - |
| 1716 | 6 | unknown | 20 | 4.76E-06 | 22 | 1.16E-05 | 6 | 0.00052 |
| 928 | 10 | hypothetical protein BRAFLDRAFT_87756* | 21 | 4.76E-06 | - | - | - | - |
| 67 | 89 | short-chain collagen c4 | 22 | 4.76E-06 | 27 | 2.31E-05 | - | - |
| 22 | 181 | von willebrand factor d and egf domain-containing | 23 | 4.76E-06 | 15 | 1.49E-06 | - | - |
| 513 | 16 | fibropellin | 24 | 6.81E-06 | 39 | 0.00064 | - | - |
| 135 | 48 | myc homolog | 25 | 6.81E-06 | 10 | 1.51E-07 | - | - |
| 1457 | 7 | unknown | 26 | 1.30E-05 | - | - | - | - |
| 3508 | 4 | unknown | 27 | 1.82E-05 | 75 | 0.00455 | - | - |
| 3509 | 4 | unknown | 28 | 1.82E-05 | 14 | 1.25E-06 | - | - |
| 24 | 167 | cytochrome p450 | 29 | 3.73E-05 | - | - | - | - |
| 142 | 47 | unknown | 30 | 3.94E-05 | 21 | 1.16E-05 | - | - |
| 36 | 139 | polyprotein | 31 | 5.97E-05 | 29 | 3.75E-05 | - | - |
| 1036 | 9 | serine protease inhibitor cvsi-2 | 32 | 5.99E-05 | - | - | 1 | 8.10E-10 |
| 2141 | 5 | unknown | 33 | 7.50E-05 | - | - | - | - |
| 335 | 23 | unknown | 34 | 8.55E-05 | 13 | 1.25E-06 | - | - |
| 888 | 10 | unknown | 35 | 0.00011 | 18 | 3.87E-06 | - | - |
| 52 | 108 | multicopper oxidase | 36 | 0.00018 | - | - | - | - |
| 1604 | 6 | unknown | 37 | 0.00019 | 93 | 0.00904 | - | - |
| 1747 | 6 | unknown | 38 | 0.00019 | - | - | - | - |
| 1817 | 6 | heat shock protein 60 | 39 | 0.00019 | - | - | 8 | 0.00052 |
| 463 | 18 | loc571499 protein | 40 | 0.00019 | 46 | 0.00096 | - | - |
| 109 | 58 | tyrosinase/cre-tyr protein | 41 | 0.00021 | - | - | - | - |
| 4174 | 3 | unknown | 42 | 0.00035 | 28 | 3.68E-05 | - | - |
| 4944 | 3 | unknown | 43 | 0.00035 | - | - | - | - |
| 5117 | 3 | unknown | 44 | 0.00035 | - | - | - | - |
| 103 | 62 | serine protease* | 45 | 0.00035 | - | - | 3 | 0.00037 |
| 658 | 13 | unknown | 46 | 0.00038 | - | - | - | - |
| 254 | 30 | chitin binding domain | 47 | 0.00042 | - | - | 16 | 0.00713 |
|   |   |   |   |   |   |
|---|---|---|---|---|---|
| 3 | 635 | nacht and ankyrin domain containing protein / ankyrin repeat protein / ankyrin unc44* | 48 | 0.00046 | 7 | 5.66E-09 |
| 97 | 65 | c1q domain containing protein | 49 | 0.00048 | 16 | 1.99E-06 |
| 39 | 137 | transient receptor potential cation subfamily member* | 50 | 0.00049 | - | - |
| 1200 | 8 | unknown | 51 | 0.00061 | 12 | 1.25E-06 |
| 1258 | 8 | AF369699_1SHG | 52 | 0.00061 | - | - |
| 341 | 23 | unknown | 53 | 0.00066 | - | - |
| 542 | 16 | hemaglutinin/amebocyte aggregation factor/dermatopontin 2 | 54 | 0.00095 | 42 | 0.00064 |
| 970 | 9 | collagen* | 55 | 0.00095 | - | - |
| 1031 | 9 | protein-glutamine gamma-glutamyltransferase | 56 | 0.00095 | - | - |
| 1079 | 9 | glycoside hydrolase | 57 | 0.00095 | - | - |
| 2593 | 4 | unknown | 58 | 0.00095 | - | - |
| 3367 | 4 | unknown | 59 | 0.00095 | - | - |
| 3386 | 4 | unknown | 60 | 0.00095 | - | - |
| 3460 | 4 | unknown | 61 | 0.00095 | - | - |
| 3921 | 4 | unknown | 62 | 0.00095 | 33 | 0.00012 |
| 30 | 152 | gtpase imap family member | 63 | 0.00095 | 19 | 4.38E-06 |
| 505 | 17 | unknown | 64 | 0.00117 | 30 | 5.17E-05 |
| 929 | 10 | epididymal secretory protein e1 precursor/niemann-pick type c2 | 65 | 0.00139 | - | - |
| 287 | 27 | pancreatic lipase-related protein | 66 | 0.00139 | - | - |
| 47 | 122 | organic cation transporter | 67 | 0.00145 | - | - |
| 840 | 11 | beta-lactamase family protein* | 68 | 0.00202 | - | - |
| 25 | 160 | neurotransmitter transporter | 69 | 0.00202 | - | - |
| 2108 | 5 | unknown | 70 | 0.00204 | - | - |
| 2140 | 5 | c1q domain containing | 71 | 0.00204 | 82 | 0.00665 | 4 | 0.00052 |
|    |    | protein                                                                 |    |    |    |    |
|----|----|-------------------------------------------------------------------------|----|----|----|----|
| 71 | 85 | von willebrand factor d and egf domain-containing*                       | 72 | 0.00313 | - | - | - |
| 1615 | 6 | myc homolog                                                             | 73 | 0.00340 | - | - | - |
| 1642 | 6 | unknown                                                                 | 74 | 0.00340 | - | - | - |
| 1656 | 6 | monocarboxylate transporter                                             | 75 | 0.00340 | - | - | - |
| 1808 | 6 | unknown                                                                 | 76 | 0.00340 | - | - | - |
| 1810 | 6 | pe-pgrs family protein                                                  | 77 | 0.00340 | - | - | - |
| 1873 | 6 | unknown                                                                 | 78 | 0.00340 | 95 | 0.00904 | - |
| 1973 | 6 | unknown                                                                 | 79 | 0.00340 | - | - | - |
| 1989 | 6 | actin binding protein                                                   | 80 | 0.00340 | 37 | 0.00054 | - |
| 654 | 13 | cell adhesion molecule                                                 | 81 | 0.00340 | - | - | - |
| 663 | 13 | sialic acid binding lectin/c1q domain containing protein*               | 82 | 0.00340 | - | - | - |
| 669 | 13 | paired box protein                                                      | 83 | 0.00340 | - | - | - |
| 700 | 13 | retinal-binding protein/sec1411 protein                                 | 84 | 0.00340 | - | - | - |
| 68 | 88 | peroxidase                                                              | 85 | 0.00350 | 43 | 0.00065 | - |
| 29 | 153 | serine threonine kinase 3/7                                            | 86 | 0.00429 | 35 | 0.00041 | - |
| 619 | 14 | hypothetical protein BRAFLDRAFT_129074*                                | 87 | 0.00442 | 36 | 0.00042 | - |
| 626 | 14 | tissue inhibitor of metalloproteinase-3                                 | 88 | 0.00442 | - | - | - |
| 137 | 48 | sodium myo-inositol cotransporter / sodium glucose cotransporter*       | 89 | 0.00472 | - | - | - |
| 1365 | 7 | unknown                                                                 | 90 | 0.00529 | - | - | - |
| 1469 | 7 | mgc154819                                                              | 91 | 0.00529 | - | - | - |
| 552 | 15 | sushi domain-containing protein 2/von willebrand factor type d domain protein | 92 | 0.00555 | - | - | - |
| 570 | 15 | hypothetical protein BRAFLDRAFT_117849                                  | 93 | 0.00555 | - | - | - |
| 294 | 26 | jagged 1-like                                                           | 94 | 0.00661 | - | - | - |
| 514 | 16 | serine protease inhibitor-1l                                            | 95 | 0.00698 | - | - | - |
| No. | Gene Name                                      | Accession | E-Value | Identity | Length | COI |
|-----|-----------------------------------------------|-----------|---------|----------|--------|-----|
| 541 | t-box transcription factor                     |           | 0.00698 |          |        |     |
| 58  | unknown                                       |           | 0.00704 | 69       | 0.00447|     |
| 1161| isopentenyl pyrophosphate:dimethylallyl pyrophosphate | 98        | 0.00743 |          |        |     |
| 1191| endo-1'4'-beta-d-glucanase                     | 99        | 0.00743 |          |        |     |
| 1226| BRAFLDRAFT_84494                              | 100       | 0.00743 |          |        |     |
| 1249| unknown                                       | 101       | 0.00743 | 47       | 0.00115|     |
| 489 | unknown                                       | 102       | 0.00829 |          |        |     |
| 490 | ring finger protein 213-like                  | 103       | 0.00829 |          |        |     |
| 278 | unknown                                       | 104       | 0.00845 |          |        |     |
| 174 | tripartite motif-containing                   | 105       | 0.00891 |          |        |     |
| 1042| endo-beta-1'4'-glucanase                      | 106       | 0.00973 |          |        |     |
| 168 | amino acid transporter                        | 107       | 0.00973 |          |        |     |
| 416 | receptor tyrosine kinase / insulin receptor*  | 108       | 0.00973 |          |        |     |
| 430 | novel protein vertebrate egf-like repeats and discoidin i-like domains 3* | 109       | 0.00973 |          |        |     |
| 4167| unknown                                       | 110       | 0.00973 | 54       | 0.00285|     |
| 4249| pacifastin                                    | 111       | 0.00973 |          |        |     |
| 4254| collagen alpha-1 chain                        | 112       | 0.00973 |          |        |     |
| 4314| unknown                                       | 113       | 0.00973 | 55       | 0.00285|     |
| 4315| unknown                                       | 114       | 0.00973 |          |        |     |
| 4316| mitogen-activated protein kinase kinase kinase 7* | 115       | 0.00973 |          |        |     |
| 4360| unknown                                       | 116       | 0.00973 |          |        |     |
| 4509| unknown                                       | 117       | 0.00973 | 56       | 0.00285|     |
| 4539| unknown                                       | 118       | 0.00973 |          |        |     |
| 4545| unknown                                       | 119       | 0.00973 |          |        |     |
| 4613| legumain                                      | 120       | 0.00973 |          |        |     |
| 4633| unknown                                       | 121       | 0.00973 | 57       | 0.00285|     |
| 4715| unknown                                       | 122       | 0.00973 |          |        |     |
| ID   | Count | Description                                      | p-value | E-value | Score |
|------|-------|--------------------------------------------------|---------|---------|-------|
| 4736 | 3     | peptidoglycan-binding domain 1 protein            | 0.00973 | -       | -     |
| 4786 | 3     | unknown                                          | 0.00973 | 58      | 0.00285 |
| 5277 | 3     | unknown                                          | 0.00973 | -       | -     |
| 5296 | 3     | predicted protein                                | 0.00973 | 60      | 0.00285 |
| 5743 | 3     | unknown                                          | 0.00973 | -       | -     |
| 5832 | 3     | unknown                                          | 0.00973 | -       | -     |
| 5912 | 3     | unknown                                          | 0.00973 | -       | -     |
| 6037 | 3     | unknown                                          | 0.00973 | -       | -     |
| 6056 | 3     | unknown                                          | 0.00973 | -       | -     |
| 6151 | 3     | unknown                                          | 0.00973 | 61      | 0.00285 |
| 6333 | 3     | unknown                                          | 0.00973 | -       | -     |
| 6428 | 3     | unknown                                          | 0.00973 | 62      | 0.00285 |
| 6438 | 3     | unknown                                          | 0.00973 | -       | -     |
| 6600 | 3     | unknown                                          | 0.00973 | 63      | 0.00285 |
| 6626 | 3     | unknown                                          | 0.00973 | -       | -     |
| 163  | 44    | rapunzel 4/5*                                    | 0.00978 | 11      | 9.26E-07 |
| 404  | 20    | apextrin-like                                    | -       | 20      | 7.96E-06 |
| 76   | 84    | neoverrucotoxin                                  | -       | 23      | 1.59E-05 |
| 1010 | 9     | unknown                                          | -       | 31      | 6.64E-05 |
| 234  | 32    | peptidoglycan-binding lysin domain*              | -       | 32      | 9.77E-05 |
| 51   | 111   | cell adhesion*                                   | -       | 38      | 0.00064  |
| 92   | 68    | unknown                                          | -       | 44      | 0.00076  |
| 61   | 92    | unknown                                          | -       | 45      | 0.00085  |
| 450  | 19    | unknown                                          | -       | 48      | 0.00115  |
| 370  | 22    | unknown                                          | -       | 50      | 0.00193  |
| 372  | 22    | unknown                                          | -       | 51      | 0.00193  |
| 838  | 11    | interferon-induced very large gtpase 1           | -       | 53      | 0.00285  |
| 4912 | 3     | unknown                                          | -       | 59      | 0.00285  |
| 299  | 26    | unknown                                          | -       | 64      | 0.00303  |
|    |   |                      |    |                      |    |                      |    |
|----|---|---------------------|----|---------------------|----|---------------------|----|
| 703| 12| unknown             |    | -                   | 65 | 0.00306             |    |
| 753| 12| unknown             |    | -                   | 66 | 0.00306             |    |
| 157| 45| predicted protein(gi|156221710|gb|ED O42562.1)*       |    | -                   | 67 | 0.00308             |    |
| 263| 29| unknown             |    | -                   | 68 | 0.00435             |    |
| 2783| 4 | unknown             |    | -                   | 70 | 0.00455             |    |
| 2857| 4 | unknown             |    | -                   | 71 | 0.00455             |    |
| 2901| 4 | legumain            |    | -                   | 72 | 0.00455             |    |
| 2902| 4 | unknown             |    | -                   | 73 | 0.00455             |    |
| 3158| 4 | unknown             |    | -                   | 74 | 0.00455             |    |
| 3606| 4 | unknown             |    | -                   | 76 | 0.00455             |    |
| 3827| 4 | creb-binding protein|    | -                   | 77 | 0.00455             |    |
| 3873| 4 | unknown             |    | -                   | 78 | 0.00455             |    |
| 124| 52| interferon-induced protein | 44 | -                   | 79 | 0.00503             |    |
| 567| 15| aromatic amino acid decarboxylase | 80 | -                   | 0.00503 |    |
| 230| 33| hypothetical protein BRAFLDRAFT_82912 |    | -                   | 81 | 0.00597             |    |
| 2238| 5 | unknown             |    | -                   | 83 | 0.00665             |    |
| 2460| 5 | unknown             |    | -                   | 84 | 0.00665             |    |
| 2463| 5 | tissue inhibitor of metalloproteinase 3 |    | -                   | 85 | 0.00665             |    |
| 2487| 5 | unknown             |    | -                   | 86 | 0.00665             |    |
| 2520| 5 | unknown             |    | -                   | 87 | 0.00665             |    |
| 2521| 5 | unknown             |    | -                   | 88 | 0.00665             |    |
| 74 | 84| SINV_05289          |    | -                   | 89 | 0.00746             |    |
| 454| 18| unknown             |    | -                   | 90 | 0.00771             |    |
| 104| 62| cell cycle checkpoint protein rad17 |    | -                   | 91 | 0.00904             |    |
| 1575| 6 | unknown             |    | -                   | 92 | 0.00904             |    |
| 1646| 6 | receptor for egg jelly |    | -                   | 94 | 0.00904             |    |
| 1962| 6 | antileukoproteinase |    | -                   | 96 | 0.00904             |    |
Pearson’s chi-squared test was performed to test for the enrichment of each group of DEGs for each TribeMCL cluster. If the expected count for any cell in the 2x2 contingency table was 5 or fewer, Fisher’s exact test was performed instead. Significance p-values listed above were adjusted for the number of independent enrichment tests performed for each group of DEGs by the False Discovery Rate correction method. Enriched TribeMCL clusters are ordered above by (1) rank order of significance of enrichment for F3L_early DEGs, (2) rank order of significance of enrichment for F3L_late DEGs, then (3) rank order of significance of enrichment for GX_early DEGs. Enriched TribeMCL clusters were annotated with a protein name if more than half of the contigs had identical or nearly identical BLASTX best hits and the remainder of contigs had no significant hits, or if ≥ 80% of the contigs had identical or nearly identical BLASTX best hits and the remainder of contigs had dissimilar best BLASTX hits (which may be the case in the sharing of domains, e.g., neurotrypsin and trypsin). In the cases in which contigs that composed a TribeMCL cluster had no best BLASTX hits or had a number of dissimilar hits that if counted conjointly did not represent a majority of the contigs, then that TribeMCL cluster was not annotated, that is, was named “unknown.” * Enriched TribeMCL clusters were reservedly annotated if less than half of the contigs had identical or nearly identical BLASTX best hits and composed a plurality while the remainder of contigs had no significant hits or a small number of dissimilar but non-repeating hits (frequency equal to one), or if a number of dissimilar hits of similar frequency that if counted conjointly composed ≥ 50% of the contigs in the cluster (e.g., “hemicentin/rhamnospondin/thrombospondin*”).

| DEG  | Contig | Protein Description | F3L_early | F3L_late | GX_early | p-value |
|------|--------|---------------------|-----------|----------|----------|---------|
| 411  | 20     | gm2 ganglioside activator | -         | -        | 97       | 0.00975 |
| 1781 | 6      | organic solute transporter subunit alpha | -         | -        | -        | 7       | 0.00052 |
| 1369 | 7      | unknown             | -         | -        | -        | 9       | 0.00059 |
| 1119 | 8      | melatonin receptor 1a-like | -         | -        | -        | 11      | 0.00071 |
| 1026 | 9      | unknown             | -         | -        | -        | 12      | 0.00083 |
| 919  | 10     | unknown             | -         | -        | -        | 13      | 0.00096 |
| 99   | 64     | fibropellin         | -         | -        | -        | 14      | 0.00210 |
| 48   | 117    | monocarboxylate transporter | -         | -        | -        | 18      | 0.00912 |
| 192  | 37     | proprotein convertase subtilisin kexin* | -         | -        | -        | 19      | 0.00922 |
Table 9. Comparison of select “diversified” groups of genes in the Eastern oyster to the Pacific oyster *C.gigas* genome and differential expression

| Select Gene Groups of Interest | SPs | SPIs | FREPs | C1qDC proteins | CTLDC proteins | DMBT1/ SRCR type 12 | IFI44 | GIMAP proteins |
|-------------------------------|-----|------|-------|----------------|---------------|-------------------|-------|---------------|
| # of contigs                  | 112 | 99   | 180   | 492            | 404           | 187               | 88    | 210           |
| # of nonredundant contigs (reduced by 95% similarity) | 73  | 81   | 128   | 391            | 276           | 122               | 59    | 173           |
| # of nonredundant contigs that map to *C. gigas* genome | 70  | 73   | 115   | 323            | 220           | 109               | 45    | 158           |
| # of unique *C. gigas* loci to which contigs map | 51  | 61   | 74    | 149           | 140           | 61                | 22    | 33            |
| # of genes in *C. gigas* genome | 22<sup>a</sup> | 40<sup>a</sup> | 190<sup>b</sup> | 321<sup>b</sup> | 266<sup>b</sup> | 43<sup>a</sup> | 27<sup>a</sup> | 19<sup>a</sup> |
| #DEGs GX early | 7   | 8    | 0     | 5             | 6             | 3                | 2     | 3            |
| #DEGs F3L early | 13  | 22   | 34    | 78            | 56            | 21               | 5     | 18           |
| #DEGs F3L late | 2   | 9    | 27    | 54            | 25            | 11               | 6     | 18           |
| #DEGs total | 14/112 | 24/99 | 41/180 | 90/492 | 65/404 | 22/187 | 9/88 | 21/210 |

a. Number of genes in the *C. gigas* genome determined by number of *C. gigas* GLEAN gene model peptide sequences with reciprocal blast hits (e-value ≤ 1e-05) to transcript nucleotide sequence in each group of interest (TBLASTN, BLASTX) and with best blast hits with the select gene group of interest.

b. Numbers of genes in the *C. gigas* genome provided by Zhang et al. 2012.
Figure 1. Cumulative percent mortality in two families of oysters challenged with the bacterial pathogen *Roseovarius crassostreae* (F3L and GX) and in unchallenged controls (CF3L and CGX).

The cumulative mortality is shown over the course of the 93-day bacterial challenge for both families (F3L and GX) and both challenge and control oysters, the latter of which is indicated by the prefix “C” for “control.” Time 0 is the moment of exposure by bath to *Roseovarius crassostreae*. A Pearson’s chi-squared test of significance performed for each pairwise comparison between groups F3L, CGX, and GX at day 28, two days before the final RNA sample collection, and at day 93, the final timepoint of the bacterial challenge. Significance values (p-values) were adjusted independently at each timepoint by the Bonferroni method to account for the multiple comparisons. The four arrows indicate days 1, 5, 15, and 30 at which timepoints RNA was isolated from CGX, F3L, and GX, for cDNA synthesis and sequencing.
Figure 2. Heatmap of all differentially expressed genes (DEGs) in two oyster families experimentally challenged with the bacterial pathogen *Roseovarius crassostreae* (GX and F3L) and unchallenged controls.

For all genes differentially expressed in any one of the treatments GX-d1, GX-d5, F3L-d1, F3L-d5, F3L-d15, and F3L-d30, the Z-score centered log2-transformed RPKM for each gene in each of eight treatment-days (including control, CGX-d15 and CGX-d30) is shown. Genes were hierarchically clustered using Euclidean distance and complete linkage of the Z-score-transformed gene expression. Sample groups were clustered using the complete linkage Euclidean distance of the Spearman correlation of the Z-score-transformed gene expression. Clustering and visualization were performed using the fastcluster and gplots packages, respectively, in the R programming environment.
Figure 3. Numbers of differentially expressed contigs (DEGs) shared and unique between GX_early (resistant oysters – days 1 and 5), F3L_early (susceptible oysters – days 1 and 5), and F3L_late (susceptible oysters – days 15 and 30)
Figure 4. Functionally enriched Gene Ontology terms among DEGs from resistant (GX) and susceptible (F3L) oysters at early (1 and 5 d) and late (15 and 30 d) time points compared to control (CGX) oysters mapped by semantic similarity using SimRel method and REVIGO

Functionally enriched Gene Ontology (GO) terms for each category of differentially expressed genes, GX_early, F3L_early, F3L_late, and for the three highest-level categories of the Gene Ontology hierarchy are displayed in SimRel semantic mapping space (Schlicker et al. 2006) by modifying the output of the REVIGO server (Supek et al. 2011) in the R programming environment and plotting using ggplot2. Semantic mapping space is
equivalent within the same broad GO term category, e.g. a x,y-coordinate in GX_early biological process space would have the same identity as the same x,y-coordinate in F3L_early biological process space. The color of nodes from cool (green) to warm (red) signifies increasing significance of enrichment as indicated in the legend. The size of nodes reflects whether the GO term is enriched among upregulated DEGs (large) or downregulated DEGs (small), while a GO term enriched among both upregulated and downregulated DEGs is represented by a medium size node and can be further identified by its unique, square shape. To enhance readability, overlapping nodes were sometimes labeled conjointly. This was done in a manual manner by selecting a term name which properly described the conjointly labeled terms. These cases were noted by the addition of the suffix “-related.”
Figure 5. Numbers of non-redundant transcripts/proteins and gene models in selected putative diversified gene families in multiple organisms from diverse taxa

| Organism                  | Serine proteases | Serine Protease Inhibitors | FREPs | C1QDC proteins | CTLDC proteins | SRCR class B | IF44 | IAN / G/MAP |
|---------------------------|------------------|----------------------------|-------|----------------|----------------|--------------|------|------------|
| Nematostella vectensis    | 69 / 57          | 28 / 23                    | 59 / 49 | 0 / 0          | 30 / 25        | 51 / 42      | 3 / 2 | 1 / 1      |
| Saccoglossus kowalevskii  | 38 / 63          | 6 / 10                     | 29 / 48 | 1 / 2          | 29 / 48        | 16 / 26      | 0 / 0 | 6 / 10     |
| Petromyzon marinus        | 64 / 62          | 14 / 14                    | 17 / 16 | 5 / 5          | 11 / 11        | 22 / 21      | 2 / 2 | 0 / 0      |
| Branchiostoma floridiae   | 96 / 78          | 23 / 19                    | 185 / 149 | 25 / 20  | 375 / 303       | 119 / 96     | 5 / 4 | 15 / 12    |
| Cliona intestinalis       | 57 / 61          | 10 / 11                    | 37 / 40 | 0 / 0          | 36 / 39        | 9 / 10       | 1 / 1 | 0 / 0      |
| Strongylocentrotus purpuratus | 72 / 85       | 21 / 86                    | 43 / 51 | 2 / 2          | 73 / 86        | 311 / 366    | 1 / 1 | 0 / 0      |
| Daphnia pulex             | 205 / 237        | 26 / 30                    | 5 / 6   | 0 / 0          | 3 / 3          | 11 / 13      | 0 / 0 | 0 / 0      |
| Drosophila melanogaster   | 231 / 239        | 38 / 39                    | 14 / 15 | 1 / 1          | 4 / 4          | 17 / 18      | 1 / 1 | 0 / 0      |
| Pinctada fucata           | 25 / 9           | 24 / 9                     | 91 / 33 | 170 / 61       | 75 / 27        | 48 / 17      | 17 / 6 | 27 / 10    |
| Crassostrea gigas         | 48 / 18          | 65 / 25                    | 78 / 30 | 264 / 109       | 177 / 68       | 57 / 22      | 53 / 20 | 27 / 10    |
| Crassostrea virginica     | 76 / 7           | 83 / 8                     | 174 / 17 | 431 / 42  | 298 / 29        | 131 / 13     | 90 / 9 | 156 / 15   |
| Lottia gigantea           | 57 / 61          | 23 / 25                    | 55 / 59 | 3 / 3          | 28 / 30        | 24 / 26      | 4 / 4 | 21 / 23    |
| Biomphalaria glabrata     | 18 / 25          | 13 / 18                    | 22 / 30 | 2 / 3          | 7 / 10         | 3 / 4        | 0 / 0 | 8 / 11     |
| Aplysia californica       | 20 / 8           | 6 / 2                      | 14 / 5  | 2 / 1          | 13 / 5         | 9 / 3        | 0 / 0 | 8 / 3      |

In each cell is shown side-by-side: 1) the numbers of non-redundant transcripts/proteins, as determined by similarity clustering and 2) numbers of gene models (see Methods). A species tree is reproduced beside the matrix to emphasize evolutionarily relationships between the featured organisms. The number of gene models (the second number in each cell) was Z-score centered by gene family and the magnitude of this Z-score was assigned a color according to the color gradient indicated in the key.
Figure 6. *C. virginica* GIMAP translated transcripts clustered by similarity with color and size reflective of differential expression.

*C. virginica* GIMAP translated transcripts represented in Cytoscape 2.8 and yFiles organic layout. An edge corresponds to a significant similarity (e-value ≤ 1e-05, hit identity ≥ 20%), and the wider the edge, the higher the hit identity percentage.
Figure 7. *C. virginica* IFI44 translated transcripts clustered by similarity with color and size reflective of differential expression.

*C. virginica* IFI44 translated transcripts represented in Cytoscape 2.8 and yFiles organic layout. An edge corresponds to a significant similarity (e-value $\leq 1e-05$, hit identity $\geq 20\%$), and the wider the edge, the higher the hit identity percentage.
A manually curated multiple alignment of GTPase of immunity associated protein (GIMAP) translated transcripts/proteins from multiple organisms in diverse taxa was used generate the above maximum likelihood phylogenetic tree. The tree is represented as a circular cladogram and bootstrap support from 1,000 replicates is indicated as a percentage next to each tree node. Leaves of the tree are colored according to the species to which the sequence belongs, as specified in the legend.
A manually curated multiple alignment of GTPase of immunity associated protein (GIMAP) translated transcripts/proteins from multiple organisms in diverse taxa was used to generate the above maximum likelihood phylogenetic tree using FastTree2 assuming a WAG model and hybrid CAT/gamma approximations (as in Fig. 8). The tree is represented as a circular cladogram. Bootstrap support from 1,000 replicates is indicated for the only node used to define the three major groupings indicated by the solid or dotted arcs that circumscribe the tree. (The group indicated by the dotted arc is dotted because it was defined negatively by exclusion.) The numbers of sequences for each species in each of the major groupings are listed in parentheses beside the species abbreviation. Some details are provided on the C. gigas genomic loci to which the C. virginica sequences mapped.
A manually curated multiple alignment of interferon-induced protein 44 (IFI44) translated transcripts/proteins from multiple organisms in diverse taxa was used to generate the above maximum likelihood phylogenetic tree using FastTree2 assuming a WAG model and hybrid CAT/gamma approximations. The tree is represented as a circular cladogram and bootstrap support from 1,000 replicates is indicated as a percentage next to each tree node. Leaves of the tree are colored according to the species to which the sequence belongs, as specified in the legend.
A manually curated multiple alignment of interferon-induced protein 44 (IFI44) translated transcripts/proteins from multiple organisms in diverse taxa was used generate the above maximum likelihood phylogenetic tree using FastTree2 assuming a WAG model and hybrid CAT/gamma approximations (as in Fig. 10). The tree is represented as a circular cladogram. Bootstrap support from 1,000 replicates is indicated for only those nodes used to define the major groupings indicated by the solid arcs that circumscribe the tree. The major groupings of the tree that contained bivalve-only sequences were indicated by red coloration in the inter-branch space and the major grouping that contained sequences from diverse organisms was indicated by light blue coloration in the inter-branch space. The numbers of sequences for each species in each of the major groupings are listed in parentheses beside the species abbreviation. Some details are provided on the C. gigas genomic loci to which the C. virginica sequences mapped.
### APPENDIX

#### A. Tables

Table S1. Sources and type of sequences downloaded for multiple organisms for multi-species similarity clustering

| Organism       | Type of seq. | Source                                                                 | Notes                        | Date downloaded | Name of seq. file             |
|----------------|--------------|------------------------------------------------------------------------|------------------------------|-----------------|--------------------------------|
| L. stagnalis   | ESTs         | http://www.ncbi.nlm.nih.gov/nucest                                    |                              | 2/28/12         |                                |
| B. glabrata    | ESTs         | http://www.ncbi.nlm.nih.gov/nucest                                    |                              | 2/28/12         |                                |
| A. californica | ESTs         | http://www.ncbi.nlm.nih.gov/nucest                                    | EST assembly*                | 2/28/12         |                                |
| C. gigas       | ESTs         | http://public-contigbrowser.sigenae.org:9090/Crassostrea_gigas/index.html | version 8                    | 2/28/12         | cgigas_all_contigs.tfa         |
| B. floribae    | Protein      | http://www.uniprot.org/uniref/, Suzek et al. 2007                     | Uniprot                       | 2/28/12         |                                |
| D. melanogaster| Protein      | http://www.uniprot.org/uniref/, Suzek et al. 2007                     | Uniprot                       | 2/28/12         |                                |
| D. pulex       | Protein      | http://www.uniprot.org/uniref/, Suzek et al. 2007                     | Uniprot                       | 2/28/12         |                                |
| N. vectensis   | Protein      | http://www.uniprot.org/uniref/, Suzek et al. 2007                     | Uniprot                       | 2/28/12         |                                |
| C. intestinalis| Protein      | http://www.uniprot.org/uniref/, Suzek et al. 2007                     | Uniprot                       | 2/28/12         |                                |
| S. purpuratus  | Protein      | http://sugp.caltech.edu/SpBase/download/                              | translations of gene models   | 2/28/12         | SPU_peptide.fasta              |
| P. marinus     | Protein      | ftp://ftp.ensembl.org/pub/release-67/fasta/petromyzon_marinus/pep/    | translations of gene models   | 2/28/12         | Petromyzon_marinus.Pmarinus_7.0.66.pep.all.fa |
| S. kowalevskii | Protein      | http://www.ncbi.nlm.nih.gov/RefSeq/                                   | translations of gene models   | 2/28/12         |                                |
| L. gigantea    | Protein      | http://genome.jgi-psf.org/Lotgi1/Lotgi1.do wnload.fpl.html             | translations of gene models   | 2/28/12         | Lotgi1_GeneModels_FilteredModels1 aa.fasta |
| P. fucata      | Protein      | http://marinegenomics.oist.jp/genomes/download?project_id=20          | translations of gene models   | 2/28/12         | pfu_aug1.0_Pall.fasta          |
* ESTs (255,605) were assembled using CLC Genomics Workbench (version 4.8) on default settings into 70,053 sequences (including contigs and singletons).
Table S2. Estimation of total number of gene models in the genomes of multiple organisms

| Organism       | Number of gene models | Source                                           |
|----------------|-----------------------|--------------------------------------------------|
| B. glabrata    | 21900                 | transferred from http://genome.jgi-psf.org/Lotgi1/Lotgi1.info.html |
| A. californica | 25000                 | (roughly estimated here)                         |
| C. gigas       | 28027                 | Zhang et al. 2012                                |
| C. virginica   | 28027                 | transferred from Zhang et al. 2012               |
| B. floridæ     | 21900                 | Dishaw et al. 2012                               |
| D. melanogaster| 14442                 | Hahn et al. 2007                                 |
| D. pulex       | 30907                 | Colbourne et al. 2012                            |
| N. vectensis   | 18000                 | Putnam et al. 2007                               |
| C. intestinalis| 16000                 | Dehal et al. 2002                                |
| S. purpuratus  | 28944                 | http://www.spbase.org/SpBase/resources/index.php |
| P. marinus     | 10402                 | http://useast.ensembl.org/Petromyzon_marinus/Info/Annotation/#genebuild |
| S. kowalevskii | 20000                 | http://www.hgsc.bcm.tmc.edu/project-species-o-Acorn%20worm.hgsc |
| L. gigantea    | 23851                 | http://genome.jgi-psf.org/Lotgi1/Lotgi1.info.html |
| P. fucata      | 23257                 | Takeuchi et al. 2012                             |
Table S3. Unique annotations in the set of GX_early DEGs not in F3L DEGs

| Contig          | Reg. GX 1d | Reg. GX 5d | logFC | log CPM | p-value | Best blastx hit to nr db | Hit accession | Hit e-value |
|-----------------|------------|------------|-------|---------|---------|--------------------------|---------------|-------------|
| comp12125_c0_seq3 | NA         | up         | 2.76  | 0.95    | 0.000178 | notch 2                  | XP_858190     | 3.06E-07    |
| comp1285_c1_seq3 | NA         | down       | -3.54 | 3.50    | 9.56E-05 | arginase ii              | XP_00213083-4 | 6.53E-12    |
| comp1285_c1_seq8 | NA         | down       | -3.90 | 3.84    | 2.38E-05 | arginase type i-like protein | AEB70965     | 5.87E-28    |
| comp18757_c0_seq1 | NA         | down       | -7.14 | 0.09    | 0.000236 | hla-b associated transcript 1 | XP_00321735-0 | 6.91E-21    |
| comp18902_c0_seq1 | NA         | down       | -3.43 | 1.62    | 0.000649 | rho otase                | XP_00273910-5 | 1.37E-58    |
| comp24124_c0_seq1 | NA         | up         | 4.49  | 2.59    | 9.31E-09 | ched related family member (ptr-19) | XP_00273410-0 | 1.39E-13    |
| comp24124_c0_seq4 | NA         | up         | 2.71  | 1.85    | 0.000232 | ched related family member (ptr-19) | XP_00273410-0 | 1.20E-07    |
| comp2906_c1_seq15 | up         | NA         | 2.62  | 1.79    | 0.000334 | acetyl-carboxylase        | AAF22966      | 9.83E-44    |
| comp4626_c0_seq4 | up         | NA         | 3.34  | 2.55    | 3.32E-06 | probable alpha-ketoglutarate-dependent hypophosphite dioxygenase-like | XP_00294490-0 | 1.40E-10    |
| comp6161_c0_seq10 | up         | NA         | 3.17  | 2.50    | 2.16E-05 | furin (paired basic amino acid cleaving enzyme) | CBY34171      | 4.38E-13    |
| comp6161_c0_seq11 | up         | NA         | 3.10  | 3.16    | 1.59E-05 | furin (paired basic amino acid cleaving enzyme) | AAA49718      | 1.42E-22    |
| comp6161_c0_seq5 | up         | NA         | 3.49  | 3.24    | 1.57E-06 | type 2 proinsulin processing endopeptidase | 2206277A      | 2.33E-42    |
| comp664_c0_seq4  | up         | NA         | 2.42  | 4.67    | 0.000621 | unc-5 homolog b           | XP_00164203-0 | 1.34E-14    |
| comp6837_c0_seq1 | up         | NA         | 3.15  | 4.45    | 1.16E-05 | interleukin 17d           | A9XE49        | 1.05E-56    |
| comp6837_c0_seq2 | up         | NA         | 3.02  | 4.92    | 2.41E-05 | interleukin 17-like       | A9XE49        | 7.77E-66    |
| comp688_c0_seq1  | up         | NA         | 2.52  | 7.53    | 0.000343 | erythrocyte membrane-associated giant protein antigen 332 | XP_00216700-6 | 6.21E-09    |
| comp7972_c0_seq1 | NA         | down       | -8.09 | 1.08    | 7.32E-06 | cubilin                  | XP_00273439-2 | 0           |
| comp7972_c0_seq4 | NA         | down       | -3.11 | 4.03    | 0.000313 | cubilin-like              | XP_00261297-7 | 0           |
| comp7992_c0_seq1 | up         | NA         | 2.52  | 4.11    | 0.000383 | aac4 protein              | XP_797207     | 2.33E-60    |

Annotations unique to DEGs in GX days 1 and 5 are shown. Significance of differential is expressed as FDR-adjusted p-value, magnitude of differential expression is expressed as log-10 fold change over control pool (logFC) and abundance is expressed as log-10 counts per million (logCPM). Regulation, or “Reg.”, (up- or down-regulated) is shown for each contig.
Table S4. Numbers of single nucleotide polymorphisms (SNPs)

|                           | Number of Contigs | Number of Contigs with SNPs | Total Number of SNPs | Number of Synonymous SNPs | Number of Non-synonymous SNPs | mean dN/dS |
|---------------------------|------------------|-----------------------------|----------------------|--------------------------|-------------------------------|-------------|
| Total Transcriptome       | 363173           | 69711                       | 185024               | NA                       | NA                            | NA          |
| Within determined coding region, no indels allowed | 272760           | 48115                       | 123987               | 86439                    | 37548                         | ND          |
| DEG groups                |                  |                             |                      |                          |                               |             |
|GX_early DEGs unique       | 201              | 56                          | 148                  | 99                       | 49                            | 0.20        |
|GX_early DEGs shared with F3L_early and/or F3L_late | 357              | 105                         | 283                  | 174                      | 109                           | 0.26        |
|Diversified Groups of Interest |                |                             |                      |                          |                               |             |
|Serine Proteases          | 73               | 42                          | 81                   | 56                       | 25                            | 0.07        |
|Serine Protease Inhibitors| 81               | 32                          | 90                   | 28                       | 62                            | 0.19        |
|C1qDC proteins            | 391              | 108                         | 251                  | 128                      | 123                           | 0.35        |
|FREP s                    | 276              | 34                          | 90                   | 58                       | 32                            | 0.22        |
|IAN/GIMAP proteins         | 173              | 27                          | 62                   | 36                       | 26                            | 0.25        |
|IFi44                      | 59               | 11                          | 29                   | 14                       | 15                            | 0.48        |
|CTLDC proteins            | 276              | 105                         | 300                  | 209                      | 91                            | 0.19        |
|DMBT1/SRCR type 12         | 122              | 56                          | 206                  | 150                      | 56                            | 0.22        |
B. Supplementary Figures

Figure S1. Sequence read processing statistics in numbers of reads and length of reads.
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