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Highly Variable Immune-Response Proteins (185/333) from the Sea Urchin, *Strongylocentrotus purpuratus*: Proteomic Analysis Identifies Diversity within and between Individuals

Nolwenn M. Dheilly,* Sham V. Nair,2* L. Courtney Smith,† and David A. Raftos*

185/333 genes and transcripts from the purple sea urchin, *Strongylocentrotus purpuratus*, predict high levels of amino acid diversity within the encoded proteins. Based on their expression patterns, 185/333 proteins appear to be involved in immune responses. In the present study, one- and two-dimensional Western blots show that 185/333 proteins exhibit high levels of molecular diversity within and between individual sea urchins. The molecular masses of 185/333-positive bands or spots range from 30 to 250 kDa with a broad array of isoelectric points. The observed molecular masses are higher than those predicted from mRNAs, suggesting that 185/333 proteins form strong associations with other molecules or with each other. Some sea urchins expressed >200 distinct 185/333 proteins, and each animal had a unique suite of the proteins that differed from all other individuals. When sea urchins were challenged in vivo with pathogen-associated molecular patterns (PAMPs; bacterial LPS and peptidoglycan), the expression of 185/333 proteins increased. More importantly, different suites of 185/333 proteins were expressed in response to different PAMPs. This suggests that the expression of 185/333 proteins can be tailored toward different PAMPs in a form of pathogen-specific immune response. *The Journal of Immunology*, 2009, 182: 2203–2212.

Recent studies of host defense have uncovered profound differences among animal phyla in the molecules used to mediate immune responses (1–3). It seems that the immune systems of different animals have evolved a variety of solutions to meet a basic requirement to combat pathogens. The resulting, highly diversified immune responses may reflect the specific physiology, lifespan, habitat, and associated microbial populations of particular animal groups, or they may have arisen by chance via evolutionary radiation. A number of metazoan phyla have now been studied, identifying a range of alternative immunological mechanisms that exhibit high levels of molecular diversity (2). These discoveries have driven a paradigm shift in our understanding of invertebrate immune responses, from systems that are simple and static, to those that are complex and have novel mechanisms for generating molecular hypervariability, a key requirement for keeping pace in the “arms race” against microbial pathogens.

Some invertebrate immune systems are proving to be surprisingly complex. Recent analyses of the purple urchin (*Strongylocentrotus purpuratus*) genome identified several large gene families, including gene models for 222 TLRs, 203 NOD-like receptors (NLR), 218 scavenger receptor cysteine-rich (SRCR) molecules, and 104 C-type lectins (4–6). The complexity and large size of these gene families suggest that the receptors they encode may recognize individual pathogen-associated molecular patterns (PAMPs)3 with a high degree of specificity. They might also act combinatorially, providing highly diverse recognize capabilities (5). In addition to diversified receptors, the *S. purpuratus* genome contains homologs of RAG1 and RAG2, the molecules responsible for the somatic recombination of immunoglobulins in vertebrates (7). This suggests that the molecular tools required to generate molecular hypervariability might also exist among invertebrates.

185/333 genes represent another high-diversity immune response system in sea urchins. This family was first identified during a transcriptome analysis of sea urchin immune response genes (8). Genes that were up-regulated in coelomocytes (immune cells) after the injection of LPS were identified by screening high-density arrayed, conventional cDNA libraries with probes generated by subtractive suppression hybridization. Surprisingly, ~60% of the expressed sequence tags (ESTs) characterized in this transcriptome analysis were members of a set of closely related transcripts with similarities to two uncharacterized sequences from *S. purpuratus*, called DD185 (GenBank accession AF228877 (9)) and EST333 (GenBank accession R62081 (10)), hence the designation 185/333.

Screening an arrayed cDNA library constructed from immunologically activated coelomocytes indicated that the frequency of 185/333 mRNAs was enhanced more than 75-fold compared with a nonactivated arrayed cDNA library (8). Northern blots also showed striking increases in 185/333 expression in coelomocytes from bacterially activated sea urchins compared with injury controls (9). Based on this significant increase in gene expression, 185/333 transcripts were investigated in more detail, revealing an

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3 Abbreviations used in this paper: PAMP, pathogen-associated molecular pattern; aCF, artificial coelomic fluid; CF, coelomic fluid; EST, expressed sequence tag; IPG, immobilized pH gradient; LC, liquid chromatography; MS, mass spectrometry; 1DE, one-dimensional electrophoresis; pI, isoelectric point; SNP, single nucleotide polymorphism; 2DE, two-dimensional electrophoresis.

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unexpected level of sequence diversity (8, 11). To date, for 185/333 cDNAs, 689 have been characterized from 14 sea urchins. These sequences are predicted to encode 286 different proteins (11, 12).

The diversity evident among 185/333 transcripts is generated in several ways. 185/333 mRNAs are comprised of 25 different blocks or “elements” of nucleotide sequence that are present or absent in numerous combinations. This results in “element patterns” that are repeatedly identified in different mRNAs (8, 12), contributing significant diversity to the family of 185/333 messages. Single nucleotide polymorphisms (SNPs) and small insertions or deletions (indels) are also frequent in all 185/333 sequences, adding to diversity. Additionally, there are surprisingly high levels of frame shifts and the insertion of early stop codons in the mRNA sequences (11). The processes that generate this diversity among 185/333 mRNAs are not yet well defined, but they may include differences among the estimated fifty 185/333 gene loci (11, 13, 14), high levels of allelic polymorphism at each locus, RNA editing, and/or low-fidelity RNA polymerases (15), followed by posttranslational modifications. All of these processes may have been driven by positive selection associated with anti-pathogen defense (8, 11, 12, 14). Nucleotide sequence variability results in high levels of nonconservative amino acid substitutions among predicted proteins of the type consistent with intense evolutionary selection pressure (8, 12, 14, 15).

185/333 mRNAs are predicted to encode proteins with a hydrophobic leader, a glycine-rich region with multiple endoprotease cleavage sites, an RGID motif, a histidine-rich region, numerous N-linked and O-linked glycosylation sites, acidic patches, and several types of tandem and interspersed repeats (11). The deduced proteins do not contain cysteines, transmembrane regions, GPI linkage sites, identifiable domains, or any predictable folding patterns. The only regions with at least some similarity to other molecules are the RGID motif and one of the histidine-rich domains, which is comparable to histatins, a group of mammalian salivary proteins with powerful antifungal activities (16, 17). Brockton et al. (18) have shown that 185/333 proteins are expressed by two subsets of coelomocytes (small phagocytes and polygonal cells), and that the number of 185/333+ cells increases in response to immunological challenge. Initial analyses of 185/333 proteins indicated that recombinant 185/333 proteins appear as multimers and that native proteins are present in a broad range of sizes in agreement with the mRNA sequences (18).

Here, we present the first comprehensive analysis of 185/333 proteins. It reveals a broad diversity of these proteins within individual sea urchins. Different sea urchins express different suites of 185/333 proteins, and expression is altered by immunological challenge. The data indicate that differential expression or posttranslational modification of 185/333 proteins might allow S. purpuratus to tailor immune responses toward specific pathogens.

Materials and Methods

Sea urchins

Adult S. purpuratus were purchased from Marinus Scientific after collection from the coast of southern California. They were maintained in the laboratory as described previously (19). S. purpuratus becomes immunquiescent after long-term housing of >8 mo without significant disturbance (19, 20). Immunouquiescence can easily be reversed by injecting PAMPs, or in response to injury (19–21).

Immunological challenge and sample collection

Animals were challenged by injecting 2 μg of LPS or 4 μg of peptidoglycan (Sigma-Aldrich) per milliliter of coelomic fluid (CF), as previously described (10, 19). Control animals were injected with an equivalent volume of artificial CF (aCF) (11). CF (100 μl) was withdrawn from each sea urchin immediately before injection, and then at various times after injection. A 23-gauge needle attached to a 1-ml syringe was inserted through the peristome into the coelomic cavity and CF was withdrawn without anticogulant. The CF was immediately expelled into a 1-ml tube and mixed with 100 μl of urea sample buffer (2.4 M Tris-HCl (pH 6.8), 0.25% SDS, 4 M urea, 20% glycerol). Samples were stored at −70°C until used. Proteins were precipitated using 2-D Clean-Up kits (GE Healthcare) according to the manufacturer’s instructions and resuspended in urea sample buffer (8 M urea, 4% CHAPS, 60 mM DTT). The total protein content of each sample was determined with 2-D Quant kits (GE Healthcare).

One-dimensional electrophoresis (1DE)

CF proteins in urea sample buffer (100 μg/well) were separated on 10% Tris-glycine precast polyacrylamide gels (Criterion gel system; Bio-Rad) at 130 V for 2 h, or on 7.5% bis-Tris polyacrylamide gels at 180 V for 1 h. After electrophoresis, proteins were visualized using Sypro Ruby (Sigma-Aldrich) following the manufacturer’s protocol, or with Coomassie blue using standard protocols. Alternatively, the proteins were transferred to nitrocellulose membranes by Western blotting as described below.

Two-dimensional electrophoresis (2DE)

Isoelectrofocusing was performed using an IPGphor IEF system (GE Healthcare). Immobilized pH gradient (IPG) gel strips (11 cm, pH 3–6; or pH 3–10; GE Healthcare) were rehydrated overnight with 200 μg of CF proteins in 200 μl of rehydration buffer (8 M urea, 2% CHAPS, 50 mM DTT, and 0.5% carrier ampholytes; Immobiline; GE Healthcare). Isolelectrofocusing was undertaken at 100 V for 3 h, 250 V for 20 min, and 8000 V for 6 h, to obtain a total of 26,000–30,000 Vh. The proteins in the IPG strips were reduced (1% DTT, 15 min) and alkylated (2.5% iodoacetamide, 15 min) before separation in the second dimension by SDS-PAGE using 10% Tris-glycine precast polyacrylamide gels (Criterion gel system; Bio-Rad) (22). After electrophoresis, protein spots on the gels were visualized by Coomassie blue staining or with Sypro Ruby. Alternatively, proteins were transferred to nitrocellulose membranes, as described see below.

Antibodies

The polyclonal antiseria against 185/333 proteins used in this study were the same as those reported by Brockton et al. (18). Antiseria were generated against synthetic peptides corresponding to elements 1, 7, and 25a, which are present in most 185/333 cDNAs (see Ref. 11 and Fig. 5). The peptides were conjugated to keyhole limpet hemocyanin and injected into two rabbits per peptide on four separate occasions (Quality Controlled Biochemicals). Only those antiseria for which the preimmunization bleeds did not cross-react with sea urchin CF proteins by Western blot (18) were used in this study. The three antiseria used were designated anti-185-66, anti-185-68, and anti-185-71.

Western blotting and immunostaining

Proteins separated by 1DE or 2DE were transferred from polyacrylamide gels to nitrocellulose membranes by electroblotting using a Criterion blotting system (Bio-Rad). Transfers were performed at 100 V for 1 h at room temperature using Towbin buffer (0.25 M Tris-HCl, 1.92 M glycine (pH 8.3)) with 20% methanol. Once transfer was complete, membranes were blocked by incubation in skim milk solution (7% skim milk powder in TBST; 10 mM Tris-HCl, 137 mM NaCl, 0.5% Tween 20 (pH 7.5)) overnight. Following several washes with TBST (three times for 5 min), membranes were incubated with anti-185/333 antiseria (1/20,000 dilution of an equal mix of anti-185-66, -68, and -71 in TBST, or each antiserum separately) for 1 h at room temperature. The blots were washed with TBST (three times for 5 min) and incubated for 1 h at room temperature with goat anti-rabbit IgG conjugated with HRP (1/30,000 in TBST; Sigma-Aldrich). After washing in TBST (three times for 5 min), 185/333+ proteins were visualized using ECL chemiluminescence (GE Healthcare) with blue light-sensitive high-performance chemiluminescence film (Hyperfilm ECL, GE Healthcare). Each blot was exposed to film for varying lengths of time (1–15 min) to optimize the exposure. In some cases, image processing (Adobe Photoshop; Adobe Systems) was used to combine autoradiographs from the various exposures into composite images. Proteome maps for 185/333 (Photoshop; Adobe Systems) was used to combine autoradiographs from the various exposures into composite images. Proteome maps for 185/333 (Photoshop; Adobe Systems) was used to combine autoradiographs from the various exposures into composite images. Proteome maps for 185/333 (Photoshop; Adobe Systems) was used to combine autoradiographs from the various exposures into composite images.
imidazole (CMFSW-EI; 10 mM KCl, 7 mM NaSO₄, 2.4 mM NaHCO₃, 460 mM NaCl, 70 mM EDTA, and 50 mM imidazole (pH 7.4)) and was mixed with an equal volume of CMFSW-EI containing 1% (v/v) Nonidet P-40 to lyse the cells, followed by centrifugation (12,000 × g, 10 s) to remove debris. The supernatant was diluted 1/20 with TBS and aliquoted in triplicate into 96-well ELISA plates (200 µl/well; Corning). Plates were incubated overnight at 4°C so that proteins could adhere to the wells. The plates were washed once with TBS and blocked for 1 h with TBS containing 4% (w/v) BSA. After blocking, 100 µl of anti-185 antisera (1/20,000 dilution of an equal mix of anti-185-66, -68, and -71 in TBST) was rehydrated with trypsin (12.5 ng/ml in 0.1 M glycine, 1 mM MgCl₂, 1 mM ZnCl₂ (pH 10.4); Sigma-Aldrich) was added per well and incubated for 30 min before absorbance was read in a microplate spectrophotometer at 415 nm. Data were corrected for the absorbance in wells prepared without sea urchin proteins. Controls included wells in which the primary (anti-185) Abs or both the primary and secondary Abs were omitted, and wells in which an irrelevant Ab (rabbit anti-tunicate collectin peptide (24, 25)) was used in place of the anti-185 antiserum.

Mass spectrometry and data analysis

Mass spectrometry (MS) was performed on CF proteins separated by either 1DE or 2DE. To extract proteins from 1DE, Coomassie blue-stained gels were cut into slices twice in water (10 min each). Individual lanes were cut into 16 slices of equal sizes so that proteins in different molecular mass ranges could be analyzed separately by MS (see Fig. 8). To extract proteins after 2DE, spots from 2DE gels that corresponded to individual 185/333+ proteins on Western blots were excised. Gel slices or excised protein spots were washed briefly with 100 mM NH₄HCO₃, before being destained (3 × 10 min) with 25 mM NH₄HCO₃/acetoni­trile (1/1) and dehydrated in 100% acetoni­trile for 5 min. After dehydration, gel pieces were air-dried, rehydrated with 10 mM DTT for 10 min and twice with 25 mM NH₄HCO₃/acetoni­trile (1/1) for 3 min before being dehy­drated with 100% acetoni­trile. The dehydrated gel slices were air-dried and rehydrated with trypsin (12.5 ng/µl in 50 mM NH₄HCO₃; Promega) for 30 min at 4°C. An additional aliquot of 50 mM NH₄HCO₃ was added before the proteins were digested at 37°C overnight. The resulting tryptic peptides were extracted from the gel pieces by washing twice with 2% formic acid in 50% acetoni­trile. Extracts were combined and concentrated to 10 µl by vacuum centrifugation.

Mass spectrometry was performed at the Australian Proteome Analysis Facility (Macquarie University). The tryptic digest extracts from 1DE gel slices were subjected to data-dependent nanocapillary reversed phase liquid chromatography followed by electrospray ionization using a Thermo LCQ Deca ion trap mass spectrometer (Thermo Scientific; liquid chromatography–MS–MS). For LC-MS/MS, a microbore HPLC system (TSP4000; Thermo Scientific) was modified to operate at capillary flow rates using a simple T-piece flow-splitter. Columns (8 cm × 100 µm inside diameter) were packed with 100 Å, 5-µm Zorbax C18 resin at 500 ř. Integrated electrospray tips for the columns were made from fused silica, pulled to a 5-µm tip using a laser puller (Sutter Instrument). An electrospray voltage of 1.8 kV was applied using a gold electrode via a liquid chromatography followed by electrospray ionization using a Thermo LCQ Deca ion trap mass spectrometer (Thermo Scientific; liquid chromatography–MS–MS). For LC-MS/MS, a microbore HPLC system (TSP4000; Thermo Scientific) was modified to operate at capillary flow rates using a simple T-piece flow-splitter. Columns (8 cm × 100 µm inside diameter) were packed with 100 Å, 5-µm Zorbax C18 resin at 500 ř. Integrated electrospray tips for the columns were made from fused silica, pulled to a 5-µm tip using a laser puller (Sutter Instrument). An electrospray voltage of 1.8 kV was applied using a gold electrode via a liquid junction upstream of the column. Samples were introduced onto the analyti­cal column using a Surveyor Autosampler (Thermo Scientific). The HPLC column eluent was eluted directly into the electrospray ionization source of the ion trap mass spectrometer. Peptides were eluted with a linear gradient of buffer A (0.1% formic acid) and buffer B (acetoni­trile containing 0.1% formic acid) at a flow rate of 500 nL/min. Automated peak recognition, dynamic exclusion, and daughter ion scanning of the top three most intense ions were performed using the Xcalibur software as previously described (23).

GPM open source software (Global Proteome Machine Organization; www.thegpm.org) was used to search peptide sequences against a combined S. purpuratus database created with sequences downloaded from the National Center for Biotechnology Information (NCBI). This FASTA format database contained 44,037 protein sequences comprising all EST sequences (8) using the Mascot search engine (Matrix Sciences; www.matrixscience.com/) set for carbomethylation (C) and oxidation (M) variable modifications, with peptide and fragment mass tolerances of ±0.05 Da, and a maximum missed cleavage of one.

Results

One-dimensional SDS-PAGE analysis of CF proteins

1DE was used to provide a preliminary assessment of the diversity of proteins in CF, particularly those that were detected by anti-185 Abs (Fig. 1). The molecular masses of CF proteins from all of the sea urchins analyzed in the present study (n = 13) ranged from 20 kDa to >193 kDa (see Fig. 6). Syp­ro Ruby-stained CF proteins from the individual sea urchin ranged from 30 kDa to >193 kDa (Fig. 1, lane 1). There were substantial differences in banding patterns when electrophoretically separated CF proteins from the same sea urchin were stained with Coomassie blue compared with Syp­ro Ruby (Fig. 1, compare lanes 1 and 2). This was probably due to different sensitivities and physio­chemical properties of the stains (26).

Immunostained 1DE Western blots of CF proteins identified numerous 185/333+ bands ranging from ~20 kDa to >193 kDa (Fig. 1, lane 3). Controls that omitted the primary (anti-185) and/or secondary (anti-IgG) Abs, and irrelevant controls using anti­tunicate collectin Abs instead of anti-185 antisera, were negative (data not shown). Most 185/333+ bands from all of the sea urchin samples analyzed (n = 13) had molecular masses ranging from 50 kDa to >193 kDa, although some were as low as 20 kDa (see Fig. 6). Many of the 185/333+ bands on Western blots were not visible on Syp­ro Ruby- and Coomassie blue-stained 1DE gels, suggesting that the 185/333 proteins were present at extremely low concentrations.
Two-dimensional Western blots of CF proteins

Given that cDNA analyses predicted a broad array of 185/333 proteins in the CF (8, 11, 12), we extended our analysis of 185/333 proteins using 2DE, which affords far greater resolution than does 1DE. Sea urchin CF proteins were initially separated by isoelectric focusing (pH 3–10) and then by 10% SDS-PAGE. Large numbers of 185/333 compounds were evident after 2DE, to the extent that they often appeared as dense smears (Fig. 2). Most of the 185/333 proteins recognized by the antisera in all of the sea urchins analyzed (n = 13) had pIs between 3 and 7 with apparent molecular masses of 40 kDa to >193 kDa.

To improve the resolution of individual 185/333 compounds on 2DE Western blots, additional isoelectric focusing separations were conducted on immobilized pH gradient (IPG) strips with a pH range of 3–6. Composite images obtained from film exposures of three different time intervals were used to visualize 185/333 compounds of varying abundance (Fig. 3). For the individual sea urchin shown in Fig. 3, image analysis of the different exposures revealed that fifty-one 185/333 compounds were evident after a 1-min film exposure. An additional 117 spots were evident after 5 min, and a further 96 spots appeared after 10 min, making a total of 264 spots that were detected on the composite image for this individual. The number of discrete 185/333 compounds varied substantially among individuals. Three of the 13 sea urchins tested did not express detectable levels of 185/333 proteins. However, the number of discrete 185/333 compounds was often >200 in other individuals.

The enhanced protein separation capabilities of 2DE also showed that each discrete 185/333 compound evident in 1DE contained numerous variants with similar molecular masses but different pIs. For example, at least fifteen 185/333 spots with different pIs were evident at ~75 kDa (Fig. 3). A further eight 185/333 spots appeared at ~60 kDa and at least six different 185/333 spots were present at ~30 kDa (Fig. 3). These pI variants were often regularly spaced from each other, differing by ~0.1–0.2 pH units. There were also numerous 185/333 compounds with identical pI but different molecular masses (Fig. 4). For instance, three proteins in the pI 7.5–7.75 range each had three different molecular mass forms at ~150 kDa, 193 kDa, and >193 kDa (Fig. 4).

The three different anti-185 sera, which were raised against different regions of the most commonly predicted 185/333 polypeptide sequences, identified subsets of 185/333 proteins (Fig. 5). Within the small region of a 2DE Western blot (pI range of 4–5 and molecular masses of 30 kDa or lower), anti-185-66 resolved 14 variant distinguishable variants, of which 7 were 30 kDa and had a pI range of 4–5 (Fig. 5A). An additional seven anti-185-66-positive spots were smaller than 30 kDa, with a pI range of 4.5–5. In comparison, anti-185-68 recognized five of the seven proteins at 30 kDa with a pI range of 4–5 that were present on the anti-185-66 blot, but did not recognize the set of seven lower molecular mass spots. Anti-185-71 recognized only two of the 30 kDa proteins that were recognized by the other two antisera. This result is consistent with the expression of truncated 185/333 cDNAs (11).

Diversity of 185/333 proteins between individuals

There were major differences in the expression profiles of 185/333 proteins among different sea urchins. Some animals did not show 185/333 expression before LPS injection (Fig. 6A, lanes 3 and 11), although expression was evident after challenge (Fig. 6B, lanes 3 and 11). In other cases, 185/333 expression, which was very low
FIGURE 6. 1DE Western blots of CF from 13 different sea urchins sampled before (A) and 96 h after (B) challenge with LPS. The equivalent lane numbers in both panels refer to the same animals from which CF samples were obtained before and after LPS challenge. The blots were immunostained with an equal mixture of the three different anti-185 sera (anti-185-66, -68, and -71; 1/20,000). The equivalent lane numbers in both panels refer to the same animals from which CF samples were obtained before and after LPS challenge. The blots were immunostained with an equal mixture of the three different anti-185 sera (anti-185-66, -68, and -71; 1/20,000).

### Table 1. Mass spectrometric (LC-MS/MS) data for peptides isolated from 1DE gels of S. purpuratus CF that match known 185/333 sequences

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<td>RDEBETDAAQQIDGLGPGQFDGPPR</td>
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*The data are from 1DE gel slices of CF from three different sea urchins. The GenBank accession numbers of the 185/333 sequences that matched peptides from 1DE gels are also shown. The amino acid sequences shown are the matching 185/333 peptides from the NCBI database. Parameter definitions are: loge, values indicate the probability that a putative peptide sequence corresponding to a mass spectrum arises stochastically. The lower the loge value, the more significant the assignment of the peptide sequence to the mass spectrum. The loge values listed in this table are all lower than the significance cutoff values assigned by the GPM algorithm, and so they are deemed to be statistically significant (43); m + h, peptide mass in Da; z, peptide charge.*

*Peptides that matched sequences for previous analysis of nucleotide sequences that might represent frameshift mutations (12, 13).*

A broad distribution of 185/333 molecular masses ranging from 20 kDa to >193 kDa was apparent among the 13 animals analyzed by 1DE Western blots (Fig. 6). Although many individuals expressed some 185/333 proteins with identical molecular masses (e.g., 112 kDa, Fig. 6, lanes 1–4; 100 and 120 kDa, Fig. 6, lanes 7 and 8), the suite of 185/333+ bands expressed by each individual sea urchin was unique. Three animals did not express detectable 185/335 proteins, even after challenge with LPS (data not shown). In total, nineteen 185/333+ bands with distinct molecular masses were identified among the 13 sea urchins analyzed by 1DE. The molecular masses of these proteins seemed to be evenly spaced at 8–10 kDa apart over the molecular mass range of 20 to >193 kDa (Fig. 6).

**MS analysis of 185/333**+ **proteins**

Proteins on Coomassie blue-stained 2DE gels that corresponded to 185/333+ spots on Western blots were analyzed by MS. This analysis failed to identify 185/333 proteins unambiguously using the default criteria on the Mascot search engine. Although MS of each in preinjection samples, increased significantly after LPS injection.
protein successfully identified ion fragments with mass/charge properties that were similar to known 185/333 sequences, these matches did not yield sufficiently high statistical probabilities to confirm the identity of any of the proteins (data not shown).

In contrast, MS analysis of CF proteins separated by 1DE unequivocally identified 185/333 proteins. A total of 41 peptides isolated from 1DE gel slices matched with known 185/333 sequences from the NCBI database (Table I). The same 185/333 peptide was often identified in more than one gel slice. For example, the peptide FDGPESGAPQMEGR appeared in gel slices 2–4 from animal 17. It is unlikely that the occurrence of the same peptide in multiple fractions was due to contamination of the different fractions with exactly the same 185/333 protein. If the same protein was present in more than one fraction, we would expect to find exactly the same combination of peptides from that protein in more than one fraction. However, this was not the case. Only single peptides were found in each fraction. Given this repetitive identification of the same peptide in different gel slices, a total of 23 unique 185/333 peptides were identified among all of the gel slices. In some cases, the GPM algorithm allocated slightly different predicted amino acid sequences to the same MS spectrum. For instance, GPM identified two peptides (GGGGGGGMMQGGPR and GGGGGGGGMMQGGPR; underlining identifies amino acids that differ) from the same MS spectrum. This occurred because the amino acid differences between these two peptides were predicted to yield peptides with indistinguishable mass/charge ratios. Matches to both of these peptides were present in the NCBI database. In some cases, the subtle differences between peptides involved the inclusion of an arginine resulting in closely related peptides of different lengths, such as RGDGEEETDAQQIQGDGLGGPG and RGDGEEETDAAQQIQGDGLGGPG. It is noteworthy that of the 41 peptides identified, 36 were located in the glycine-rich (N-terminal) and central regions of the predicted proteins. This is consistent with frameshifts and early stop codons that have been identified in about half of the cDNAs (11). In many cases, the introduction of these frameshifts or early stop codons would have resulted in the loss of epitopes for the antisera anti-185-66, -68, and -71 (1/20,000). The numbers on the left show the approximate positions of slices cut from 1DE gels for analysis by mass spectrometry (see Table I). In total, gels were cut into 16 slices. The positions of only eight of those slices containing the highest molecular mass proteins are shown here.

The titer of 185/333 proteins detected in CF by ELISA. By 48 h after LPS injection, anti-185 reactivity had increased 2.4-fold compared with the 0 h time point (p < 0.05). It returned to levels that were indistinguishable (p > 0.05) from those before LPS challenge by 96 h. In response to aCF (control) injections, the titer of 185/333 proteins in CF also increased up to 48 h postinjection, but only to a level that was 1.4-fold greater than the preinjection time point. This increase was significantly less than the response to LPS injection with LPS performed 14 days after the primary challenge. The titer of 185/333 proteins in CF increases after immune challenge. A, 1DE Western blots of CF from a single sea urchin collected at various times after the animal was injected twice with LPS. The second LPS injection was administered 360 h after the first. B, 185/333 protein expression levels determined by ELISA (ΔOD410) at various times after naive sea urchins (n = 5) were injected with LPS or aCF (controls). Bars indicate SEM. Asterisks denote time points that differed significantly (p < 0.05) between LPS-injected sea urchins and controls.

FIGURE 8. The titer of 185/333 proteins in CF increases after immune challenge. A, 1DE Western blots of CF from a single sea urchin collected at various times after the animal was injected twice with LPS. The second LPS injection was administered 360 h after the first. B, 185/333 protein expression levels determined by ELISA (ΔOD410) at various times after naïve sea urchins (n = 5) were injected with LPS or aCF (controls). Bars indicate SEM. Asterisks denote time points that differed significantly (p < 0.05) between LPS-injected sea urchins and controls.
LPS injection 2 wk later. A similar result was apparent for animals change in response to an initial injection of LPS, or after a second predominant 185/333 proteins in CF from animals 6 and 25 did not after repeated challenge (Fig. 9). The molecular masses of the there was an increase in the relative quantity of 185/333 proteins expression of 185/333 proteins when sea urchins were challenged with PG 1DE analysis did not detect changes in the types of 185/333 pro- Diversity of 185/333 protein expression after immunological challenge 1DE analysis did not detect changes in the types of 185/333 proteins expressed by individual sea urchins in response to LPS or PG injections, even though densitometry (data not shown) indicated there was an increase in the relative quantity of 185/333 proteins after repeated challenge (Fig. 9). The molecular masses of the predominant 185/333 proteins in CF from animals 6 and 25 did not change in response to an initial injection of LPS, or after a second LPS injection 2 wk later. A similar result was apparent for animals (p < 0.05). Negative controls confirmed the specificity of the ELISA. Wells in which the primary (anti-185 sera) or secondary (anti-rabbit IgG) Abs were omitted, or when the primary Ab was replaced by an irrelevant control (anti-tunicate collectin), did not yield absorbance readings that were significantly greater than background levels (p > 0.05, data not shown).

Discussion
This study has identified substantial diversity among 185/333 proteins, as reflected by their broad range of molecular masses and pls. Most importantly, there are obvious differences in the suites of 185/333 proteins expressed by different sea urchins, and these suites of proteins undergo subtle but extensive changes in response to different types of immune challenge. The diversity of 185/333 proteins and the changes in their expression detected by the current proteomic analysis correspond with previous studies of mRNAs from sea urchins responding to different PAMPs (11). All of these data suggest that sea urchins may be capable of altering the expression of 185/333 proteins to tailor specific responses against different PAMPs.

Despite the overall agreement between our current observations of 185/333 proteins and predictions based on the prior analyses of 185/333 nucleotide sequences (11, 12), there are some discrepan- cies. The predicted sizes of 185/333 proteins based on cDNA se- quences range from 4 to 55.3 kDa (11, 12), with predicted pls ranging from 5.42 to 11.54. However, the present study shows that native 185/333 proteins have a far wider range of molecular masses (20 to >193 kDa using 1DE analysis), with predominant bands often being at the high end of this range. They also have far more acidic pls than expected, mainly between 3 and 7. It is un- likely that the discrepancy between predicted and observed molecular masses is due to disulfide-bonded oligomerization because none of the predicted 185/333 amino acid sequences identified to date contains cysteines (8, 11). Even if cysteines were present in missense sequences of frame-shifted proteins, the strong reducing conditions used to prepare samples would have disrupted any oligomers held together by disulfide bonds. Another explanation for the discrepancy in molecular masses is that 185/333 proteins are glycosylated and form large complexes covalently linked to carbohydrates. There are numerous conserved sites for N-linked glycosylation within the histidine-rich region of the 185/333 proteins (elements 11–25), and there are conserved sites for O-linked glyco- sylation in the carboxyl-terminal region (8, 11). However, deg- lycosylation of N-linked oligosaccharides failed to decrease the molecular masses of 185/333 proteins to the size of predicted monomers (L. C. Smith, unpublished data). Given these results, it seems likely that the disparity between predicted and observed molecular masses reflects oligomerization based on mechanisms.
other than disulfide bond formation that are resistant to the reducing treatments used in 1DE and 2DE. This conclusion is supported by studies of recombinant 185/333 proteins, in which the expression of a single form of 185/333 protein yields a range of expressed proteins with molecular masses corresponding to monomers, homodimers, and higher order oligomers (18).

Other data also suggest that the diversity of molecular masses evident among 185/333 proteins is increased by the expression of truncated molecules. Many of the SNPs and indels found previously in 185/333 transcripts are predicted to result in frameshifts, and the encoded proteins may be either truncated and/or have missense sequences (11, 13). No such frameshifts have been identified in the 185/333 genes. However, comparisons of gene and mRNA sequences from individual animals suggest that posttranscriptional modifications may be responsible for these missense proteins (15). If these messages are translated, it would explain why the three different anti-185 antiserum used in the present study detected different subsets of 185/333 proteins. The antiserum were raised against amino acid sequences in N terminus (element 1), the middle (element 7), and the C terminus (element 25a) of the predicted proteins. The antiserum targeted to the N terminus (anti-185-66) identified the largest number of 185/333 proteins on 2DE Western blots, presumably because its epitope is most likely to be present in every protein, including the shortest truncated forms. Of the 689 translated cDNA sequences identified to date, 676 contain the epitope within element 1 that is recognized by the anti-66 antiserum (11). Antiserum directed toward more C-terminal regions recognize decreasing numbers of 185/333 proteoforms, probably because these epitopes are not present on truncated proteins or those with missense sequence at the C terminus. Current cDNA sequence data from Tervilliger et al. (11) show that 660 of 689 cDNA sequences contain the anti-68 epitope in element 7, while only 375 cDNA sequences contain the anti-71 epitope in element 25a.

The variability evident in the molecular masses of 185/333 proteins was matched by substantial diversity in their pIs. One source for this diversity is found in mRNAs where sequence variability results in changes in charged amino acids at particular positions leading to predicted proteins with very similar molecular masses but different pIs (8, 11, 12). Tervilliger et al. (see supplemental figures S1 and S2 in Refs. 11, 12) identified numerous sequence positions in 185/333 mRNAs that encode two to four different amino acids, many with different charges. In the present study, similar subtle differences were detected by MS, which identified a number of 185/333 peptides that differ by just a single amino acid. 2DE Western blots also showed a variety of pIs for 185/333 proteins that had very similar molecular masses. In many cases, the charge variants making up a single molecular mass class of 185/333 proteins had pIs spanning the full pH range from 3 to 10, although most variants had a pI within the range of 3–6. The different pI forms were often equally spaced, suggesting that adjacent protein spots represent variants that differ by just a single charged residue.

Although the variability in pI detected in the present study agrees with the diversity seen among cDNA sequences, our proteomic data also provide evidence for additional postranslational modifications. Some 185/333 proteins have identical isoelectric points but significantly different molecular masses. This suggests that individual 185/333 proteins may be conjugated with some other molecule(s) that alters their molecular mass but not their isoelectric point. Such conjugation would provide another explanation for why many 185/333 proteoforms are much larger in molecular mass than predicted. However, the cDNA data suggest that it is also feasible for 185/333 proteins with significantly different sequences and different pIs to have the same molecular mass, resulting in a vertical ladder of spots on 2DE Western blots.

The diversity of 185/333 proteins explains our inability to unequivocally identify 185/333 proteins by MS of individual proteins isolated by 2DE. MS analysis of the 185/333 proteoforms from 2DE gels did not identify any statistically significant matches to proteins in a custom database of 185/333 sequences. However, all of the 185/333 proteins isolated by 2DE yielded ion fragments that possessed mass/charge characteristics similar to those of known 185/333 protein sequences. The available evidence suggests that the existing database of 185/333 sequences may represent only a small fraction of the 185/333 variants present in sea urchin populations. Consequently, the chance of finding a statistically significant match in this restricted data set to a single 185/333 protein isolated from the CF of an individual sea urchin may be extremely low, even though the purity of the proteins isolated by 2DE would have been high. In other words, we may not have identified matches because the variability of 185/333 sequence means that many 185/333 proteins will not yet be in our database of 185/333 sequences and so cannot be identified by MS. Our difficulties in matching isolated proteins to the existing 185/333 database highlight the problems of employing MS techniques, which search for precise similarities between peptides to characterize hypervariable proteins. It is interesting that even though members of the Down syndrome cell adhesion molecule (DSCAM) family are important proteins in the immunological responses of insects, proteomic analyses of hemolymph extracts from immunologically challenged Drosophila have not yet been able to identify DSCAMs by MS.

To circumvent the problem of matching individual 185/333 proteoforms to our restricted sequence data set, we employed "shotgun" MS (LC-MS/MS), in which all of the CF proteins within a particular molecular mass range from 1DE gels were analyzed simultaneously by MS. Unlike 2DE, in which the individual proteins spots analyzed by MS presumably contain just a single 185/333 isotype, the 1DE gel slices subjected to shotgun MS were likely to contain numerous different 185/333 isotypes. This greatly increased the chance that at least one of these isotypes would match a previously identified sequence in our 185/333 database. 1DE also has the advantage that it is not limited by the complex separation characteristics of 2DE, which, for instance, is relatively inefficient in separating hydrophobic proteins. Additionally, we used an alternative search engine (GPM as opposed to Mascot) with slightly altered search stringencies (three missed tryptic cleavages) to increase our chances of identifying matching 185/333 peptides in the available databases. This process successfully identified numerous peptides that matched precisely to sequences in the NCBI database without affecting the robustness of the sequence matches. The gel slices in which matching peptides were identified corresponded to regions on 1DE and 2DE Western blots that contained high concentrations of 185/333 proteins, which confirms the specificities of the anti-185 sera used in this study. However, shotgun MS probably still identified just a small fraction of the 185/333 peptides present within the peptide mixture, because of the limited number of potentially matching peptides in the currently available dataset. The limited sensitivity of MS in identifying highly variable proteins in the absence of comprehensive sequence databases also explains why our MS analysis only identified multiple forms of 185/333 proteins in one of the sea urchins analyzed, even though both 1DE and 2DE Western blotting revealed substantial intraindividual diversity in all sea urchins.

The difficulties associated with analyzing the diversity of 185/333 proteins were compounded by the substantial variation evident in the suites of 185/333 proteins expressed by different sea urchins.
None of the 13 sea urchins analyzed by 1DE Western blotting expressed the same pattern of 185/333+ bands, even though there were a number of molecular mass forms that were shared by some individuals. On average, two to three major bands were evident in each sea urchin, along with numerous minor bands. The differences between individuals implies that there may be a variety of mechanisms acting in concert to produce novel arrays of 185/333 proteins in individual sea urchins. These factors could include the presence or absence of different 185/333 genes in different sea urchins, the presence of different alleles at a given 185/333 gene locus, differential gene expression of 185/333 family members, posttranscriptional processing or editing of the transcripts that insert frameshifts and SNPs (15), and posttranslational processing of the proteins (13).

Our data suggest that these potential mechanisms for molecular diversification may allow sea urchins to vary the suites of 185/333 proteins that they express in response to different types of immunological challenge. ELISA provided direct evidence that 185/333 expression increases after the injection of PAMPs, while 2DE analysis of 185/333 proteins from individual sea urchins identified many subtle changes in the patterns of 185/333 proteins responding to different PAMPs. There were clear differences in the Ps and molecular masses of the proteins expressed in response to LPS compared with those synthesized by the same sea urchin in response PG. These results agree with changes that are apparent in the cDNAs from animals responding to different PAMPs, including LPS, β-1,3-glucan, or dsRNA (11). Terwilliger et al. (11) showed that a diverse size range of 185/333 messages was present in the CF of immunocompetent sea urchins, but this broad expression profile changed to one single, major mRNA size after immune challenge. Sequence analysis of mRNAs from animals responding to immune challenge also indicated that the predominant 185/333 variants in CF are different before and after challenge. Such alterations in 185/333 transcription explain the changes detected during the present study in the suites of 185/333 proteins found in the CF of the same sea urchin responding to different PAMPs.

The identification of PAMP-specific responses suggests that 185/333 protein expression can be tailored to meet different forms of immunological challenge. The obvious implication is that 185/333 proteins are involved in some form of pathogen-specific immunological response. There is growing evidence for this level of immunological specificity among a variety of invertebrates (1, 2, 5, 6). Terwilliger et al. (11) showed that a diverse size range of 185/333 messages was present in the CF of immunocompetent sea urchins, but this broad expression profile changed to one single, major mRNA size after immune challenge. Sequence analysis of mRNAs from animals responding to immune challenge also indicated that the predominant 185/333 variants in CF are different before and after challenge. Such alterations in 185/333 transcription explain the changes detected during the present study in the suites of 185/333 proteins found in the CF of the same sea urchin responding to different PAMPs.

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