

The scavenger receptor repertoire in six cnidarian species and its putative role in cnidarian-dinoflagellate symbiois

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Many cnidarians engage in a mutualism with endosymbiotic photosynthetic dinoflagellates that forms the basis of the coral reef ecosystem. Interpartner interaction and regulation includes involvement of the host innate immune system. Basal metazoans, including cnidarians have diverse and complex innate immune repertoires that are just beginning to be described. Scavenger receptors (SR) are a diverse superfamily of innate immunity genes that recognize a broad array of microbial ligands and participate in phagocytosis of invading microbes. The superfamily includes subclades named SR-A through SR-I that are categorized based on the arrangement of sequence domains including the scavenger receptor cysteine rich (SRCR), the C-type lectin (CTLD) and the CD36 domains. Previous functional and gene expression studies on cnidarian-dinoflagellate symbiosis have implicated SR-like proteins in interpartner communication and regulation. In this study, we characterized the SR repertoire from a combination of genomic and transcriptomic resources from six chidarian species in the Class Anthozoa. We combined these bioinformatic analyses with functional experiments using the SR inhibitor fucoidan to explore a role for SRs in cnidarian symbiosis and immunity. Bioinformatic searches revealed a large diversity of SR-like genes that resembled SR-As, SR-Bs, SR-Es and SR-Is. SRCRs, CTLDs and CD36 domains were identified in multiple sequences in combinations that were highly homologous to vertebrate SRs as well as in proteins with novel domain combinations. Phylogenetic analyses of CD36 domains of the SR-B-like sequences from a diversity of metazoans grouped cnidarian with bilaterian sequences separate from other basal metazoans. All cnidarian sequences grouped together in a subclade separately from bilaterian sequences with moderate support. Functional experiments were carried out on the sea anemone Aiptasia pallida that engages in a symbiosis with Symbiodinium minutum (clade B1). Experimental blocking of the SR ligand binding site with the inhibitor fucoidan reduced the ability of *S. minutum* to colonize *Aiptasia* suggesting that host SRs play a role

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in host-symbiont recognition. In addition, incubation of symbiotic anemones with fucoidan elicited an immune response, indicating that host SRs function in immune modulation that results in host tolerance of the symbionts.



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Introduction

Cnidarians such as reef-building corals engage in an intimate mutualistic symbiosis with photosynthetic dinoflagellates in the genus *Symbiodinium* that together form the trophic and structural foundation of coral reef ecosystems. *Symbiodinium* spp. provide large amounts of reduced organic carbon to the host in exchange for inorganic nutrients, a high light environment and refuge from herbivory (Yellowlees et al. 2008). In the majority of cnidarian-*Symbiodinium* interactions, the symbionts are taken up by host cells *via* phagocytosis. Instead of being digested as food, the symbionts resist host destruction and persist in host cells by residing in vacuoles known as symbiosomes (Davy et al. 2012). The molecular interplay between host cnidarian and resident symbionts during both the establishment and ongoing maintenance of the symbiosis is critical for a healthy holobiont (Weis & Allemand 2009).

Animal innate immune systems are central to managing microbes by both tolerating and promoting the survival of beneficial symbionts and resisting and destroying negative invaders (Bordenstein & Theis 2015; McFall-Ngai et al. 2013; Schneider & Ayres 2008). With the increased availability of sequence resources, there is now ample evidence that innate immune pathways are ancestral and that basal metazoans including cnidarians possess many of these pathways originally described in mammals and flies (Fuess et al. 2016; Miller et al. 2007; Yuen et al. 2014). Furthermore there are numerous examples of expansions of some innate immune gene families in invertebrates that are larger than those in vertebrate genomic repertoires, including NOD-like receptors, scavenger receptors, TIR-domain-containing proteins and ficolins (Baumgarten et al. 2015; Buckley & Rast 2015; Hamada et al. 2013; Pancer 2000; Poole & Weis 2014; Shinzato et al. 2011). A class of well-described host-microbe molecular interactions mediated by innate immunity are the PRR-MAMP interactions where microbe-associated molecular patterns (MAMPs) on the surface of microbes, such as lipopolysaccharide or glycans, are recognized by pattern recognition receptors (PRRs) on the surface of host cells (Janeway & Medzhitov 2002). These steric interactions launch a series of downstream signalling cascades in the host that serve to resist and destroy negative invaders or tolerate and nurture positive microbes. Genomic and transcriptomic studies of cnidarians are revealing the presence of many classical PRRs that have been extensively characterized in higher metazoans (Fuess et al. 2016; Miller et al. 2007).

One group of PRRs in the Metazoa are the scavenger receptors (SRs), so-named for their role in the scavenging and clearing of microbial invaders, modified host molecules, and apoptotic cell debris (Areschoug & Gordon 2009; Canton et al. 2013). SRs have a high affinity for a wide range of ligands and this flexibility of ligand binding has led them to be described as 'molecular fly paper' (Krieger 1992). A key role of SRs in innate immune function is their action as PRRs on phagocytic cells where they mediate direct non-opsonic phagocytosis of pathogenic microbes (Areschoug & Gordon 2009). SRs are thought to engage in heteromultimeric signalling



99 complexes, known as signal osomes, involving multiple PRRs and other molecules that together effect signal transduction in cells, thereby alerting them to microbes or modified host molecules 100 (Canton et al. 2013). The SR superfamily is a large group of structurally diverse transmembrane 101 102 cell surface glycoproteins, divided into nine classes SR-A through SR-I (Canton et al. 2013; 103 Krieger 2001). The classes have overlapping specificities that result in an enormous breadth of 104 MAMP recognition (Krieger 1992). Members within a given class share some sequence homology, with little-to-no homology occurring between classes. The classes are grouped by 105 their multiple domains with no single domain common to all (Gordon 2002; Gough & Gordon 106 107 2000). SR domains occur on the extracellular portion of the protein; the proteins are anchored in 108 the cell membrane with transmembrane domain(s) and contain short cytoplasmic tail(s). Figure 1 109 depicts the four SR classes that are relevant to this study. SRs are a potential target for manipulation by invading parasites, pathogens and potentially mutualists. Several pathogens 110 111 have evolved mechanisms to evade SR-mediated recognition (Areschoug & Waldemarsson 112 2008; Faure & Rabourdin-Combe 2011). Indeed, certain human pathogens exploit specific SRs for their own benefit. For example, the Hepatitis C virus (HCV) (Catanese et al. 2007) and the 113 malaria parasite *Plasmodium falciparum* (Ndungu et al. 2005; Rodrigues et al. 2008) have 114 surface ligands that are recognized by SR-B1, and both use this recognition to gain entry to host 115 cells. 116

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128 129 SR-As and SR-Is contain the scavenger receptor cysteine rich (SRCR) domain, which consists of a 110 aa motif with conserved spacing of six to eight cysteines (Hohenester et al. 1999). The SRCR domain is found in a wide range of membrane and soluble proteins and often occurs in multiple repeats arrayed on the protein (Hohenester et al. 1999; Martinez et al. 2011; Sarrias et al. 2004). Some SR-As and SR-Es contain C-type lectin domains (CTLDs), a common domain in many proteins, that are often involved in lectin-glycan interactions (Cambi et al. 2005). SR-Bs contain the CD36 domain and have two cytoplasmic tails rooted in the membrane with two transmembrane regions, forming an extracellular loop (Silverstein & Febbraio 2009). SR genes encoding SRCR, CTLD and CD36 domains have been described in invertebrates (Hibino et al. 2006; Lehnert et al. 2014; Pancer et al. 1997; Schwarz et al. 2007; Wood-Charlson & Weis 2009). However, a detailed bioinformatic characterization of cnidarian SR genes homologous to vertebrate SR-As, SR-Bs, SR-Es and SR-Is is lacking, as are any functional studies exploring the function of these proteins.

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132 SRs are of interest in studies of cnidarian immunity and symbiosis. First, interactions between SR-E-like host lectin-like proteins and symbiont surface glycans play an important role in host-133 134 symbiont recognition during onset of symbiosis (reviewed in Davy et al. 2012). In addition, SR-B homologues in two species of sea anemone, Anthopleura elegantissima (Rodriguez-Lanetty et 135 136 al. 2006) and Aiptasia pallida (Lehnert et al. 2014), were found to be highly expressed in 137 symbiotic compared to aposymbiotic individuals. For A. pallida this was a dramatic difference in

138 expression where symbiotic anemones had 28-fold greater expression than aposymbiotic



139	animals. These studies suggest that SR-E and SR-B homologues are playing a role in host-
140	symbiont communication.
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142	There were two aims for this study. The first was to identify SRs in six cnidarian species, all in
143	Class Anthozoa (corals, sea anemones and others), using a variety of genomic and transcriptomic
144	resources, and compare the repertoire to vertebrate SRs of known function. This provides a
145	platform for identifying potential roles of enidarian SR proteins in immunity and symbiosis. The
146	second aim was to perform simple functional experiments to examine the role of SRs in
147	symbiont recognition and uptake by the sea anemone A. pallida, a well-studied model system for
148	the study of coral-dinoflagellate symbiosis. We hypothesized that if a symbiont is co-opting host
149	SRs to initiate tolerogenic pathways that dampen or prevent an immune response, blocking SR-
150	ligand-binding capabilities would induce an immune response.
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152 **Materials and Methods** 153 154 Anthozoan genomic and transcriptomic resources To characterize the SR protein repertoire in chidarians, six species with publically available 155 156 resources were searched. These included three anemone species: A. elegantissima (Kitchen et al. 157 2015), Aiptasia pallida (Baumgarten et al. 2015; Lehnert et al. 2012), and Nematostella vectensis (Putnam et al. 2007), and three coral species: Acropora digitifera (Shinzato et al. 2011), A. 158 159 millepora (Moya et al. 2012) and Fungia scutaria (Kitchen et al. 2015). These species were selected based on the availability of transcriptomic and genomic resources, and to include a 160 161 diversity of organisms within Class Anthozoa. These resources were derived from various 162 developmental stages and symbiotic states (Table 1). All resources were used as provided, with 163 the exception of the A. pallida transcriptome, for which raw Illumina sequence reads for 164 accession SRR696721 were downloaded from the sequence read archive entry 165 (http://www.ncbi.nlm.nih.gov/sra/SRX231866) and reassembled using Trinity (Grabherr et al. 2011) resulting in a better assembly than the original one performed. 166 167 168 SR sequence searching 169 Twenty-four non-cnidarian sequences were obtained, primarily from GenBank and other 170 publically available databases (Table S1), for use in creating multiple sequence alignments and 171 phylogenetic trees. Eleven human SR genes were chosen for production of reference protein 172 domain architecture diagrams, to compare predicted enidarian proteins with human SR proteins 173 of known function (Figure 2). 174 To search for chidarian SR proteins, databases were queried using several search strategies to 175 176 ensure all sequences were recovered. BLASTp or tBLASTn searches with mouse and human SR protein sequences (SR-A1, MARCO, SRCL, CD36, SRB1, LMP2, and LOX1) (Supplementary 177 178 file 1) and consensus sequences (pfam01130: CD36, pfam00530: SRCR) from the conserved 179 domain database (http://www.ncbi.nlm.nih.gov/cdd) as queries were performed for each resource. Keyword searches were used with the terms SR, CD36, LMP2, SRCR, and scavenger 180 where GO or KEGG annotations were available. Lastly, representative N. vectensis sequences of 181 182 each protein type (SRCR-domain-containing, CD36, SRB1, and LOX1) were also used as queries for tBLASTn searches of the other five chidarian resources. A high e-value cutoff (1x10⁻¹) 183 1) was used in the BLAST searches to recover divergent sequences. All BLAST searches were 184 performed using the default settings in Geneious pro version 7.1.8 with the exception of N. 185 vectensis, for which searches were performed through the Joint Genome Institute online portal 186 187 using the default settings (Kearse et al. 2012). A list of metazoan resources searched are listed in 188 Table S1. Blast query sequences and cnidarian sequences identified are tabulated in 189 Supplementary File 1. 190

- 191 To confirm that the sequences obtained contained SR domains, nucleotide sequences were
- translated using Geneious and then annotated using the InterProScan plugin (Quevillon et al.
- 193 2005). Only sequences in which two or more databases within InterProScan found either SRCR,
- 194 CD36, or CTLD domains with an e-value of <1x10⁻⁴ were used. Where the InterProScan plugin
- was unable to resolve protein domains, (this occurred for approx. 1 in 10 sequences) the
- sequences were analysed using the online protein domain database PfamA
- 197 (http://pfam.sanger.ac.uk) using the default program settings. Sequences for each species were
- aligned and those that were identical or almost identical (<5 aa difference in the conserved
- domains) were omitted from the analysis as they likely represented artifacts of assembly or
- 200 different isoforms of the same protein. Sequences missing a start or stop codon were removed
- 201 from the analysis.

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- 203 Only proteins that showed significant PfamA matches to a CTLD, SRCR and/or CD36 domains
- were included in the study. Diagrammatic representations of the protein domain configurations
- were produced using this information. Protein domain architectures were grouped together
- according to common domains and compared to known human SR proteins (Figure 2).
- 208 Phylogenetic analysis of SR-B homologues
- 209 A multiple sequence alignment of a subset of CD36-domain-containing sequences was
- 210 performed with the MAFFT v 7.017 plug-in (Katoh et al. 2002) through Geneious (Kearse et al.
- 211 2012), using the default settings. The program ProtTest v2.4 (Abascal et al. 2005) was used to
- apply AIC1, AIC2 and BIC2 model selection criteria to a variety of possible substitution
- 213 matrices and rate assumptions to obtain the best-fit model of protein evolution. The results from
- 214 the overall comparison of these metrics indicated that the best-fit model for the full-length
- 215 alignment was WAG+G+F. A maximum likelihood tree was produced using FastTree v2.1.5
- 216 (Price et al. 2010). Bootstrap support values were generated using the online program
- 217 SEQBOOT (Felsenstein 2005) and values above 0.6 support were displayed at the nodes. A
- 218 PhyML (Guindon et al. 2005) alternate tree produced identical topology (data not shown).

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- 220 Maintenance and preparation of anemone and dinoflagellate cultures
- 221 Symbiotic A. pallida cultures were maintained in saltwater aguaria at 26°C with a 12/12 h
- 222 light/dark photoperiod, and were fed twice weekly with live brine shrimp nauplii. Animals were
- 223 rendered aposymbiotic by incubation for 8 h at 4 °C twice weekly for six weeks, followed by
- 224 maintenance in the dark for approximately one month. Anemones were fed twice weekly with
- brine shrimp, and cleaned of expelled symbionts and food debris regularly.

- 227 Cultured dinoflagellates, Symbiodinium minutum, clade B1 (culture ID: CCMP830) were
- maintained in 50 ml flasks in sterile Guillard's f/2 enriched seawater culture medium (Sigma, St.
- 229 Louis, MO, USA). Dinoflagellate cultures were maintained at 26°C on a 12/12 h light/dark
- 230 photoperiod.



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232	In preparation for experimental manipulations, individual anemones were placed in 24-well
233	plates in 2.5 ml of 1-µm filtered seawater (FSW) and acclimated to the well-plate for 3-4 days,
234	with the FSW replaced daily. Well plates containing aposymbiotic anemones were kept in the
235	dark and symbiotic anemones were maintained in an incubator at 26°C with a 12/12 h light/dark
236	photoperiod. Animals were not fed during the experimental time period.
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238	Addition of fucoidan to block SR binding function
239	To explore a role for SRs in the onset of symbiosis, fucoidan, a known SR ligand, was added to
240	anemones to block SR binding sites. Fucoidan is a protein derived from the brown alga Fucus
241	vesiculosus; this polyanionic ligand is known to bind SRCR and CD36 domains in SR-As and
242	SR-Bs, respectively (Dinguirard & Yoshino 2006; Hsu et al. 2001; Thelen et al. 2010).
243	
244	To examine the effect of blocking SR ligand binding capabilities on symbiont colonization
245	success, aposymbiotic anemones ($n = 3$ per treatment per time point) were pre-incubated in
246	fucoidan (Sigma, St. Louis, MO, USA), at a concentration of 0 (FSW control), 100, 200 and 400
247	μg/ml for 18 h, according to Bowdish Lab protocols (online at McMaster University;
248	www.bowdish.ca/lab/protocols). Fucoidan-treated aposymbiotic anemones were subsequently re-
249	inoculated with S. minutum CCMP830. CCMP830 cells were pelleted from the culture medium,
250	re-suspended in FSW, and then added to anemones in well-plates to a final concentration of
251	2x10 ⁵ symbionts per ml. After incubation for 12 h at 26°C in the light, anemones were rinsed
252	twice with FSW and fucoidan treatments were refreshed. To test the effect of fucoidan exposure
253	on host health, a second control treatment (fucoidan-washed control) was prepared where
254	aposymbiotic anemones were pre-incubated in 200 µg/ml fucoidan for 18 h, and then washed
255	with FSW prior to being inoculated with symbionts as described above. Anemones for all
256	treatments were sampled at 48 and 96 h post-infection (three tentacles per anemone, for $n = 3$
257	anemones per treatment per time point).
258	
259	A second experiment was designed to explore a role for SR binding in host immune tolerance
260	during symbiosis. We hypothesized that if a symbiont is co-opting host SRs to initiate
261	tolerogenic pathways (such as the TGFβ pathway) that dampen or prevent an immune response,
262	blocking SR-ligand-binding capabilities could induce an immune response upon the addition of
263	lipopolysaccharide (LPS). LPS is a MAMP that has been shown to induce an anemone immune
264	response measured as increased nitric oxide (NO) production (Detournay et al. 2012; Perez &
265	Weis 2006). Symbiotic anemones were incubated at increasing concentrations of fucoidan: 0
266	(FSW control), 100, 200, 400 and 800 μ g/ml, for 4 h, prior to the addition of 1 μ g/ml of LPS
267	(Sigma, St. Louis, MO, USA) (dissolved in 0.1% v/v DMSO) for a further 12 h. The FSW
268	control was also exposed to 1 $\mu g/ml$ LPS for 12 hours. NO production by hosts was quantified as
269	described below.
270	



271 Quantifying colonization success and host NO production using confocal microscopy 272 Colonization success was assessed fluorometrically by confocal microscopy, following methods 273 described in detail by Detournay et al (2012). Briefly, following experimental manipulation, 274 solutions in wells containing anemones were replaced with 1 ml of relaxing solution (1:1 0.37 M 275 MgCl₂: FSW). Samples were observed under a Zeiss LSM 510 Meta microscope with a 40x/0.8 276 water objective lens and a working distance of 0.8–3.2 mm. Before image scanning, the focal 277 plane of the optical section was adjusted to include the gastrodermal cells within the anemone tentacle. For each experiment, all images were obtained with the same software scanning 278 279 settings, including detector gain and laser intensity. S. minutum cells present were visualized by 280 detecting chlorophyll autofluorescence with excitation and emission wavelengths of 543 and 281 600-700 nm, respectively. Fluorescence was quantified by first defining the gastrodermal tissue 282 area within the anemone tentacles as a region of interest and then measuring the mean 283 fluorescence intensity (MFI) for that region with the LSM 5 software. Intensity of chlorophyll 284 autofluorescence for each pixel was measured and a threshold value corresponding to the background was defined by measuring the MFI at 600 nm of a gastrodermal region without 285 symbionts (threshold MFI =20). Colonization success was expressed as percent of pixels with 286 287 autofluorescence intensity above the threshold. In colonization experiments, each treatment 288 represents a sample size of three anemones per treatment and time-point, with percent 289 colonization taken as the mean of six tentacles per anemone. Three untreated symbiotic 290 anemones (six tentacles per anemone) were examined to determine a baseline colonization level 291 for symbiotic anemones.

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The statistical significance of colonization success under the treatments described above was assessed using a mixed-effects model. As measures on multiple samples (i.e., tentacles) per anemone violate independence assumptions, a mixed effect was used, treating anemone as a random effect to account for correlation among samples with anemones. Main effects included time and treatment, and their interaction was estimated to account for differences between treatments at each time point. The full model can be written as:

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$$y_{i,j} = \beta X_i + \mu_j + \epsilon_{i,j}$$

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Here, $y_{i,j}$ is the logarithm of percent colonization (plus a small constant) of tentacle i within anemone j, β is a vector of effects to be estimated, X is a design matrix encoding the treatment and time point, as well as interaction term contrasts, μ_j is a normally distributed random effect for anemone j, and $\epsilon_{i,j}$ are normally distributed residuals. The model was estimated using the LME4 packages (Bates et al. 2015), for the statistical computing software R (R-Core-Team 2012) (www.R-project.org), the script and data used for statistical analyses are given in Supplementary Files 2, 3 and 4.

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To measure and visualize production of NO by confocal microscopy, animals were treated as described in detail previously (Detournay et al. 2012; Detournay & Weis 2011). Animals were transferred to a microfuge tube containing 500 ul of relaxing solution and 15 uM 4-amino-5-methylamino-2,7 difluorofluorescein diacetate (DAF-FM DA, Molecular Probes, Eugene, OR, USA) with excitation and emission wavelengths of 488 and 510–530 nm, respectively. Samples were incubated for 30 min in the dark and then rinsed twice with relaxing solution. Fluorescence of the DAF-FM DA probe was quantified as described above for chlorophyll autofluorescence quantification.

The statistical analysis of this experiment used a similar model as above, but treating the fucoidan concentration as a continuous variable and fitting a linear slope to the fluorescence intensity. We also added a random effect for tentacle within anemone to account for non-independence of readings within each tentacle. Using the notation above, the model can be written as

$$y_{k,i,j} = \alpha + \beta F_k + \mu_j + \gamma_{i(j)} + \epsilon_{k,i,j}$$

Here, $y_{k,i,j}$ is the fluorescence reading k within tentacle i of anemone j, α is the intercept and β is the slope of the regression line relating fluorescence to fucoidan concentration F_k , μ_j is a normally distributed random effect for anemone j, $\gamma_{i(j)}$ is a normally distributed random effect for tentacle i within anemone j, and $\epsilon_{k,i,j}$ are normally distributed residuals.

334 335	Results
336	Annotated predicted cnidarian SR proteins are illustrated according to their domain architecture
337	and compared with known human SR protein domain organization (Figure 2). Overall, cnidarian
338	SR-like proteins fall into four groups: SR-As, SR-Es, SR-Is and SR-Bs. The SRCR domain is
339	present in all groups except the SR-Bs.
340	
341	Cnidarian SRCR-containing proteins
342	Vertebrate SR-As are defined by a collagen domain coupled with most proteins containing either
343	an SRCR domain or a CTLD at the C terminus (Bowdish & Gordon 2009). Only two sequences
344	meeting these criteria were identified in the cnidarian resources searched. Both are in A .
345	digitifera and contain a CUB domain in addition to two collagen domains and one SRCR.
346	Human SR-Es are defined by the presence of only CTLDs (Zani et al. 2015). The human lectin-
347	like oxidized low-density lipoprotein receptor 1 (LOX1) has an N-terminal cytoplasmic tail, a
348	transmembrane domain and a single C-terminal CTLD (Canton et al. 2013). Numerous LOX1-
349	like sequences were identified in all of the cnidarian resources searched. SR-Is in humans are
350	defined by containing only SRCR domains in various numbers of repeats and are grouped into
351	three classes: CD5, CD6 and CD163. SR-I-like sequences are abundant in all cnidarian
352	resources, in the same configurations as human SR-Is. SRCR repeat numbers range from one to
353	twenty-three.
354	
355	A variety of SRCR-domain-containing proteins were also identified in cnidarian sequence
356	resources that could not be classified into any of the vertebrate classes of scavenger receptors.
357	Several cnidarians genes with SRCRs and CUB domains were identified that resemble 'human
358	deleted in malignant brain tumor' (DMBT) protein that contains eight SRCR repeats, a single
359	CUB domain and a zona pelucida domain at the C-terminal end. Predicted cnidarian proteins that
360	resemble DMBT contain one to three CUB domains combined with a range of other protein
361	domains, including MAM, fibronectin UBOX and multiple SRCRs. Five of the six cnidarian
362	resources contain sequences with a potentially novel domain configuration of multiple SRCRs,
363	several other domains, including multiple immunoglobulin domains, and a C-terminal trypsin
364	domain.
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366	Cnidarian SR-B-like proteins
367	Searches identified eighteen full-length putative cnidarian SR-B sequences, all containing a
368 369	CD36 domain. Full-length proteins were defined as those containing both transmembrane
370	regions that form the SR-B extracellular loop configuration. Humans have four distinct SR-Bs - CD36, SRB1 & 2, and LMP2 - while the six cnidarian species searched contained between two
370 371	and four full-length proteins.
372	and rour run-rengtin proteins.
373	Phylogenetic analysis of SR-B-like proteins
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374 Phylogenetic analysis was carried out on the CD36 domains from SR-B-like sequences identified 375 (Figure S1 and Figure 3). Protein sequence alignments of the predicted SR-B-like proteins from 376 cnidarians, combined with a subset of vertebrate and invertebrate sequences, revealed that there is some conservation of the CD36 domain across metazoans. Cnidarian sequences showed weak 377 homology to human SR-Bs, with 26-32%, 28-37% and 28-33% identity to human CD36, LMP2, 378 379 and SR-B1, respectively. Identities within the cnidarian group were higher, ranging from 39 to 380 95%, with the two Acropora species showing the highest homology to each other. Cnidarian sequences showed between 21 and 27% identity to the predicted SR-B-like protein sequence 381 382 from the sponge, Suberites domuncula. Predicted chidarian proteins lacked one of the three pairs 383 of cysteine residues known to form three disulphide bridges in the human CD36 protein (Figure 384 S1) (Silverstein and Febbraio, 2009). However, a pair of cysteine residues was found in all cnidarian study species at positions C107 and C117. Predicted cnidarian proteins had 8-10 N-385 386 linked glycosylation sites compared with eleven and eight sites in human SR-B1 and CD36, 387 respectively.

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Putative cnidarian SR-B proteins grouped with high support in a large clade with the bilaterians and separately from other basal metazoans. Within this large clade, cnidarians grouped together, forming a separate clade from the bilaterians. Within the cnidarian clade, there were three well-supported sub-clades, two containing both coral and anemone species and a third, containing only anemone sequences (Figure 3). Corals and sea anemones sequences formed distinct groupings within each of these clades. In contrast, bilaterian invertebrate sequences grouped with mammalian sequences in several different sub-clades of SR-Bs: LMP2, CD36, CD36-like, SR-B1, and SR-B1-like proteins.

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- Experimental blocking of SR proteins with fucoidan reduces colonization success and elicits an immune response in A. pallida
- Fucoidan-treated anemones showed significantly lower levels of colonization (0-3%) than either the FSW control or anemones pre-incubated in fucoidan and then rinsed 48 h prior to time zero Colonization success decreased significantly in a dose-dependent manner (Figure 4, Bayesian P

403 < 0.0001).

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A second fucoidan experiment investigated the possible immune-regulation role of an SR in symbiosis maintenance. Symbiotic anemones were treated with increasing concentrations of fucoidan and were subsequently immune-challenged by incubation with LPS. The FSW control-treated anemones had low levels of NO production, a proxy for an immune response, measured as MFI of the NO-specific probe DAF-FM DA in tentacles, in response to incubation in LPS. In contrast, fucoidan-treated anemones showed a significant (Bayesian P < 0.0001) dose-dependent response of increasing NO production with increasing concentrations of fucoidan (Figure 5).

413 **Discussion** 414 415 An expanded SRCR-domain-containing protein repertoire in cnidarians 416 The SRCR-domain-containing protein repertoire in chidarians, is expanded compared to that in humans, with the A. pallida genome containing the highest number at 36 genes (Figure 2). This 417 418 finding is consistent with numerous other studies describing expansions of innate immune gene 419 families in invertebrates (see Introduction). Other examples of SRCR-domain-containing protein 420 repertoire expansion have been described in invertebrates, specifically in the sea urchin, 421 Strongylocentrotus purpuratus and the cephalochordate Branchiostoma floridae, which have 218 422 and 270 SRCR-containing sequences respectively (Huang et al. 2008; Pancer 2000; Pancer et al. 423 1999; Rast & Messier-Solek 2008). These numbers are high compared to the 16 genes present in 424 humans. In addition, cnidarian SRCR-domain-containing proteins include a variety of genes with 425 novel domain combinations that have not been found in other organisms (Figure 2). 426 Identification of these novel domain combinations in chidarian immune gene repertoires is 427 consistent with other studies of basal metazoan immune genes (Hamada et al. 2013; Poole & 428 Weis 2014; Ryu et al. 2016). The searches for SR genes in the three transcriptomes (Table 1) 429 likely revealed underestimates of the total SR repertoire, given that transcriptomes represent 430 snapshots of the whole genome. 431 432 CTLD-domain-containing SRs in cnidarians 433 In contrast to the human genome, which contains a single LOX1 gene, all six cnidarian resources 434 searched contained multiple LOX1-like SR-Es (Figure 2). These searches add to previous 435 characterizations of lectin-like proteins in cnidarians, including in corals and sea anemones (Jimbo et al. 2005; Jimbo et al. 2000; Kvennefors et al. 2010; Kvennefors et al. 2008; Meyer & 436 437 Weis 2012; Vidal-Dupiol et al. 2009; Wood-Charlson & Weis 2009). Human LOX1 has a diversity of signalling functions, including in recognition of microbes via host CTLD-microbe 438 glycan binding: a PRR-MAMP interaction (Canton et al. 2013). In cnidarians, previous studies 439 440 have detailed a role for lectin-glycan interactions in the establishment of cnidarian-dinoflagellate 441 symbioses (reviewed in Davy et al. 2012). The identification of multiple LOX1-like proteins and 442 several other CTLD-containing proteins with novel domain combinations across the six species 443 examined further strengthens the hypothesis that host CTLD-symbiont glycan binding plays an 444 important role in host innate immunity and host-symbiont recognition. Cnidarian CTLD-domain-445 containing proteins described here provide potential target proteins for future experimental 446 investigation of the lectin-glycan interactions. 447 448 CD36-domain-containing SRs in cnidarians 449 Phylogenetic analysis of metazoan CD36 domains from SR-B homologues showed a well-450 supported clade of cnidarian sequences (Figure 3). A large analysis including additional 451 sequences from basal metazoans is required to more definitively reveal deep branching patterns 452 of this gene. The observed differing location of cysteine pairs within the CD36 domain in 453 cnidarian sequences compared to vertebrate ones also occurred in other invertebrates (Figure



454 S1). As with the cnidarians searched, C. elegans contained one differing pair and the three 455 sponges, Oscarella carmella, S. domuncula, and Amphimedon queenslandica, and the ctenophore *Mnemiopsis leidvi* had no sequence pairs in common with vertebrates. These 456 differences may explain why antibodies to human and mouse SR-B1 and CD36 failed to label 457 458 proteins in A. pallida in immunoblot experiments (E.F. Neubauer, unpublished data). 459 460 Functional experiments suggest that blocking SRs decreases colonization success and increases the stress response to immune challenge in A. pallida 461 462 Colonization success in aposymbiotic A. pallida challenged with S. minutum CCMP830 463 displayed a dose-dependent response to incubation in the SR inhibitor fucoidan, exhibiting decreasing colonization success with increasing concentrations of fucoidan (Figure 4). In 464 vertebrates, fucoidan blocks the positively-charged ligand binding sites on SR-As and SR-Bs. 465 and can thereby block phagocytic activity in macrophages (Dinguirard & Yoshino 2006; Hsu et 466 467 al. 2001; Li et al. 2008). The observed inhibition of colonization in cnidarians suggests that phagocytosis of symbionts is likewise inhibited and provides evidence that one or multiple SRs 468 with SRCR and/or CD36 domains function in host-symbiont recognition during onset of 469 470 symbiosis. 471 472 Previous transcriptomic studies in A. elegantissima and A. pallida have found SR-B homologues to be upregulated in symbiotic compared to aposymbiotic anemones, suggesting that they play a 473 role in the symbiosis. Our experiments showing that incubation in fucoidan causes a dose-474 475 dependent immune response in symbiotic A. pallida (Figure 5), further implicates a role for SRs 476 in immune tolerance and regulation of symbiosis. In previous work on A. pallida, we showed that symbiotic anemones produced significantly less NO in response to an immune challenge 477 478 with LPS than did aposymbiotic animals, suggesting that symbionts are modulating the host 479 immune response (Detournay et al., 2012). The increase in this response in symbiotic anemones 480 incubated in fucoidan suggests that this immune modulation involves an SR ligand-binding 481 domain. Such a response is reminiscent of immune modulation by a variety of invading microbes 482 (Janeway & Medzhitov 2002). 483 484 In summary, this study provides the first description of the diversity of SRs in cnidarians. Members include proteins with domain combinations that are highly similar to those in 485 vertebrates as well as those that possess novel combinations. Initial functional experiments using 486 the SR inhibitor fucoidan suggest that SRs play a role in the regulation of cnidarian-487 dinoflagellate symbioses. Future functional studies on candidate SRs identified in this study can 488

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491 492 further explore their role in cnidarian immunity and symbiosis.



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- 500 Abascal F, Zardoya R, and Posada D. 2005. ProtTest: selection of best-fit models of protein 501 evolution. *Bioinformatics* 21:2104-2105. DOI:
- 502 Areschoug T, and Gordon S. 2009. Scavenger receptors: role in innate immunity and microbial 503 pathogenesis. Cellular Microbiology 11:1160-1169. DOI: 10.1111/j.1462-504 5822.2009.01326.x
- 505 Areschoug T, and Waldemarsson J. 2008. Evasion of macrophage scavenger receptor A-506 mediated recognition by pathogenic Streptococci. EMBO Journal 38:3068-3079. DOI: 507 10.1002/eji.200838457
- 508 Bates D, Maechler M, Ben B, and Walker S. 2015. Fitting linear mixed-effects models using 509 lme4. Journal of Statistical Software 67:1-48. DOI: 10.18637/jss.v067.i0
- 510 Baumgarten S, Simakov O, Esherick LY, Liew YJ, Lehnert EM, Michell CT, Li Y, Hambleton 511 EA, Guse A, Oates ME, Gough J, Weis VM, Aranda M, Pringle JR, and Voolstra CR. 512 2015. The genome of Aiptasia, a sea anemone model for coral symbiosis. Proceedings of 513 the National Academy of Sciences 112:11893-11898. DOI: 10.1073/pnas.1513318112
- 514 Bordenstein SR, and Theis KR. 2015. Host Biology in Light of the Microbiome: Ten Principles 515 of Holobionts and Hologenomes. *PLoS Biol* 13:e1002226. DOI: 516 10.1371/journal.pbio.1002226
- 517 Bowdish DME, and Gordon S. 2009. Conserved domains of the class A scavenger receptors: 518 evolution and function. Immunological Reviews 227:19-31. DOI: 10.1111/j.1600-519 065X.2008.00728.x
- 520 Buckley KM, and Rast JP. 2015. Diversity of animal immune receptors and the origins of 521 recognition complexity in the deuterostomes. Developmental and Comparative 522 Immunology 49:179-189. DOI: 10.1016/j.dci.2014.10.013
- 523 Cambi A, Koopman M, and Figdor CG. 2005. How C-type lectins detect pathogens. Cellular 524 Microbiology 7:481-488. DOI: 10.1111/j.1462-5822.2005.00506.x 525
 - Canton J, Neculai D, and Grinstein S. 2013. Scavenger receptors in homeostasis and immunity. Nature Reviews Immmunology 13:621-634. DOI: 10.1038/nri3515
- 527 Catanese MT, Graziani R, von Hahn T, Moreau M, Huby T, Paonessa G, Santini C, Luzzago A, 528 Rice CM, Cortese R, Vitelli A, and Nicosia A. 2007. High-avidity monoclonal antibodies 529 against the human scavenger class B type I receptor efficiently block hepatitis C virus 530 infection in the presence of high-density lipoprotein. Journal of Virology 81:8063-8071. 531 DOI: 10.1128/JVI.00193-07
- Davy SK, Allemand D, and Weis VM. 2012. Cell biology of cnidarian-dinoflagellate symbiosis. 533 Microbiology and Molecular Biology Reviews 76:229-261. DOI: 10.1128/mmbr.05014-534
- 535 Detournay O, Schnitzler CE, Poole A, and Weis VM. 2012. Regulation of cnidarian-536 dinoflagellate mutualisms: Evidence that activation of a host TGFB innate immune 537 pathway promotes tolerance of the symbiont. Developmental and Comparative 538 Immunology 38:525-537. DOI: 10.1016/j.dci.2012.08.008
- 539 Detournay O, and Weis VM. 2011. Role of the sphingosine rheostat in the regulation of 540 cnidarian-dinoflagellate symbioses. Biological Bulletin 221:261-269. DOI:
- 541 Dinguirard N, and Yoshino TP. 2006. Potential role of a CD36-like class B scavenger receptor in 542 the binding of modified low-density lipoprotein (acLDL) to the tegumental surface of



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565

566567

- Schistosoma mansoni sporocysts. Molecular Biochemical Parasitology 146:219-230.
 DOI: 10.1016/j.molbiopara.2005.12.010
- Faure M, and Rabourdin-Combe C. 2011. Innate immunity modulation in virus entry. *Current Opinion in Virology* 1:6-12. DOI: 10.1016/j.coviro.2011.05.013
- Felsenstein J. 2005. SEQBOOT—bootstrap, jackknife or permutation resampling of molecular sequence, restriction site, gene frequency or character data. University of Washington, Seattle.
- Fuess LE, Pinzón C JH, Weil E, and Mydlarz LD. 2016. Associations between transcriptional
 changes and protein phenotypes provide insights into immune regulation in corals.
 Developmental & Comparative Immunology 62:17-28. DOI: 10.1016/j.dci.2016.04.017
- Gordon S. 2002. Pattern recognition receptors doubling up for the innate immune response. *Cell* 111:927-930. DOI: 10.1016/S0092-8674(02)01201-1
 - Gough PJ, and Gordon S. 2000. The role of scavenger receptors in the innate immune system. *Microbes and Infection* 2:305-311. DOI: 10.1016/S1286-4579(00)00297-5
- Grabherr MG, Haas BJ, Yassour M, Levin JZ, Thompson DA, Amit I, Adiconis X, Fan L,
 Raychowdhury R, Zeng Q, Chen Z, Mauceli E, Hacohen N, Gnirke A, Rhind N, di Palma
 F, Birren BW, Nusbaum C, Lindblad-Toh K, Friedman N, and Regev A. 2011. Full length transcriptome assembly from RNA-Seq data without a reference genome. *Nature Biotechnology* 29:644-652. DOI: 10.1038/nbt.1883
 - Guindon S, Lethiec F, Duroux P, and Gascuel O. 2005. PHYML Online—a web server for fast maximum likelihood-based phylogenetic inference. *Nucleic Acids Research* 33:W557-W559. DOI: 10.1093/nar/gki352
 - Hamada M, Shoguchi E, Shinzato C, Kawashima T, Miller DJ, and Satoh N. 2013. The complex NOD-like receptor repertoire of the coral *Acropora digitifera* includes novel domain combinations. *Molecular Biology and Evolution* 30:167-176. DOI: 10.1093/molbev/mss213
- Hibino T, Loza-Coll M, Messier C, Majeske AJ, Cohen AH, Terwilliger DP, Buckley KM,
 Brockton V, Nair SV, Berney K, Fugmann SD, Anderson MK, Pancer Z, Cameron RA,
 Smith LC, and Rast JP. 2006. The immune gene repertoire encoded in the purple sea
 urchin genome. *Developmental Biology* 300:349-365. DOI: 10.1016/j.ydbio.2006.08.065
- Hohenester E, Sasaki T, and Timpl R. 1999. Crystal structure of a scavenger receptor cysteinerich domain sheds light on an ancient superfamily. *Nature Structural Biology* 6:228-232. DOI: 10.1038/6669
- Hsu HY, Chiu SL, Wen MH, Chen KY, and Hua KF. 2001. Ligands of macrophage scavenger receptor induce cytokine expression via differential modulation of protein kinase signaling pathways. *Journal of Biological Chemistry* 276:28719-28730. DOI: 10.1074/jbc.M011117200
- Huang S, Yuan S, Guo L, Yu Y, Li J, and Wu T. 2008. Genomic analysis of the immune gene
 repertoire of amphioxus reveals extraordinary innate complexity and diversity. *Genome Research* 18:1112-1126. DOI: 10.1101/gr.069674.107
- Janeway CA, and Medzhitov R. 2002. Innate immune recognition. *Annual Review of Immunology* 20:197-216. DOI:
- Jimbo M, Koike K, Sakai R, Muramoto K, and Kamiya H. 2005. Cloning and characterization of a lectin from the octocoral *Sinularia lochmodes*. *Biochemical and Biophysical Research Communications* 330:157-162. DOI:



617

- Jimbo M, Yanohara T, Koike K, Koike K, Sakai R, Muramoto K, and Kamiya H. 2000. The D-galactose-binding lectin of the octocoral *Sinularia lochmodes*: characterization and possible relationship to the symbiotic dinoflagellates. *Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology* 125:227-236. DOI:
- Katoh K, Misawa K, Kuma K-i, and Miyata T. 2002. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Research* 30:3059-3066. DOI:
- Kearse M, Moir R, Wilson A, Stones-Havas S, Cheung M, Sturrock S, Buxton S, Cooper A, Markowitz S, Duran C, Thierer T, Ashton B, Meintjes P, and Drummond A. 2012.
 Geneious Basic: an integrated and extendable desktop software platform for the organization and analysis of sequence data. *Bioinformatics* 28:1647-1649. DOI:
- Kitchen SA, Crowder CM, Poole AZ, Weis VM, and Meyer E. 2015. *De novo* assembly and characterization of four anthozoan (Phylum Cnidaria) transcriptomes. *G3: Genes*|*Genomes*|*Genetics* 5:2441-2452. DOI: 10.1534/g3.115.020164
- Krieger M. 1992. Molecular flypaper and atherosclerosis: structure of the macrophage scavenger receptor. *Trends in Biochemical Sciences* 17:141-146. DOI: 10.1016/0968-0004(92)90322-Z
- Krieger M. 2001. Scavenger receptor class B type I is a multiligand HDL receptor that influences
 diverse physiologic systems. *Journal of Clinical Investigation* 108:793-797. DOI:
 10.1172/JCI14011
- Kvennefors EC, Leggat W, Kerr CC, Ainsworth TD, Hoegh-Guldberg O, and Barnes AC. 2010.
 Analysis of evolutionarily conserved innate immune components in coral links immunity and symbiosis. *Developmental and Comparative Immunology* 34:1219-1229. DOI: 10.1016/j.dci.2010.06.016
- Kvennefors ECE, Leggat W, Hoegh-Guldberg O, Degnan BM, and Barnes AC. 2008. An ancient and variable mannose-binding lectin from the coral *Acropora millepora* binds both pathogens and symbionts. *Developmental and Comparative Immunology* 32:1582-1592. DOI: 10.1016/j.dci.2008.05.010
 - Lehnert E, Burriesci M, and Pringle J. 2012. Developing the anemone *Aiptasia* as a tractable model for cnidarian-dinoflagellate symbiosis: the transcriptome of aposymbiotic *A. pallida*. *BMC Genomics* 13:271. DOI: 10.1186/1471-2164-13-271
- Lehnert EM, Mouchka ME, Burriesci MS, Gallo ND, Schwarz JA, and Pringle JR. 2014.
 Extensive differences in gene expression between symbiotic and aposymbiotic
 Cnidarians. *G3: Genes|Genomes|Genetics* 4:277-295. DOI: 10.1534/g3.113.009084
- Li B, Lu F, Wei X, and Zhao R. 2008. Fucoidan: structure and bioactivity. *Molecules* 13:1671-1695. DOI: 10.3390/molecules13081671
- Martinez VG, Moestrup SK, Holmskov U, Mollenhauer J, and Lozano F. 2011. The conserved
 scavenger receptor cysteine-rich superfamily in therapy and diagnosis. *Pharmacological Reviews* 63:967-1000. DOI: 10.1124/pr.111.004523
- McFall-Ngai M, Hadfield MG, Bosch TCG, Carey HV, Domazet-Lošo T, Douglas AE, Dubilier
 N, Eberl G, Fukami T, Gilbert SF, Hentschel U, King N, Kjelleberg S, Knoll AH, Kremer
- N, Mazmanian SK, Metcalf JL, Nealson K, Pierce NE, Rawls JF, Reid A, Ruby EG,
- Rumpho M, Sanders JG, Tautz D, and Wernegreen JJ. 2013. Animals in a bacterial
- world, a new imperative for the life sciences. Proceedings of the National Academy of
- 632 Sciences 110:3229-3236. DOI: 10.1073/pnas.1218525110



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651

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658

659

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661 662

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667

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- Meyer E, and Weis VM. 2012. Study of cnidarian-algal symbiosis in the 'omics' age. *Biological Bulletin* 223:44-65. DOI:
- 635 Miller D, Hemmrich G, Ball E, Hayward D, Khalturin K, Funayama N, Agata K, and Bosch T.
 636 2007. The innate immune repertoire in Cnidaria ancestral complexity and stochastic
 637 gene loss. *Genome Biology* 8:R59. DOI:
- Moya A, Huisman L, Ball E, Hayward D, Grasso L, Chua C, Woo H, GATTUSO JP, Forêt S,
 and Miller D. 2012. Whole transcriptome analysis of the coral *Acropora millepora* reveals complex responses to CO₂-driven acidification during the initiation of
 calcification. *Molecular Ecology* 21:2440-2454. DOI: 10.1111/j.1365 294X.2012.05554.x
- Ndungu FM, Urban BC, Marsh K, and Langhorne J. 2005. Regulation of immune response by *Plasmodium*-infected red blood cells. *Parasite Immunololgy* 27:373-384. DOI:

 10.1073/pnas.230096397
- Pancer Z. 2000. Dynamic expression of multiple scavenger receptor cysteine-rich genes in coelomocytes of the purple sea urchin. *Proceedings of the National Academy of Sciences* 97:13156-13161. DOI: 10.1073/pnas.230096397
 - Pancer Z, Munkner J, Muller I, and Muller W. 1997. A novel member of an ancient superfamily: sponge (*Geodia cydonium*, Porifera) putative protein that features scavenger receptor cysteine-rich repeats. *Gene* 193:211-218. DOI: 10.1016/S0378-1119(97)00135-2
 - Pancer Z, Rast JP, and Davidson EH. 1999. Origins of immunity: transcription factors and homologues of effector genes of the vertebrate immune system expressed in sea urchin coelomocytes. *Immunogenetics* 49:773-786. DOI: doi:10.1007/s002510050551
 - Perez S, and Weis VM. 2006. Nitric oxide and cnidarian bleaching: An eviction notice mediates the breakdown of symbiosis. *Journal of Experimental Biology* 209:2804-2810. DOI:
 - Poole AZ, and Weis VM. 2014. TIR-domain-containing protein repertoire of nine anthozoan species reveals coral–specific expansions and uncharacterized proteins. *Developmental and Comparative Immunology* 46:480-488. DOI: 10.1016/j.dci.2014.06.002
 - Price MN, Dehal PS, and Arkin AP. 2010. FastTree 2 Approximately Maximum-Likelihood Trees for Large Alignments. *PLoS One* 5:e9490. DOI: 10.1371/journal.pone.0009490
 - Putnam NH, Srivastava M, Hellsten U, Dirks B, Chapman J, Salamov A, Terry A, Shapiro H, Lindquist E, Kapitonov VV, Jurka J, Genikhovich G, Grigoriev IV, Lucas SM, Steele RE, Finnerty JR, Technau U, Martindale MQ, and Rokhsar DS. 2007. Sea anemone genome reveals ancestral eumetazoan gene repertoire and genomic organization. *Science* 317:86-94. DOI: 10.1126/science.1139158
 - Quevillon E, Silventoinen V, Pillai S, Harte N, Mulder N, Apweiler R, and Lopez R. 2005. InterProScan: protein domains identifier. *Nucleic Acids Research* 33:W116-W120. DOI: 10.1093/nar/gki442
- R-Core-Team. 2012. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL http://www.R-project.org.
- Rast JP, and Messier-Solek C. 2008. Marine invertebrate genome sequences and our evolving understanding of animal immunity. *Biological Bulletin* 214:274-283. DOI:
- Rodrigues CD, Hannus M, Prudêncio M, Martin C, Gonçalves LA, Portugal S, Epiphanio S,
 Akinc A, Hadwiger P, Jahn-Hofmann K, Röhl I, van Gemert G-J, Franetich J-F, Luty
 AJF, Sauerwein R, Mazier D, Koteliansky V, Vornlocher H-P, Echeverri CJ, and Mota
- MM. 2008. Host scavenger receptor SR-BI plays a dual role in the establishment of



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697

698

699

700

701

702

705

706

707

- 679 malaria parasite liver infection. Cell Host & Microbe 4:271-282. DOI: 680 10.1016/j.chom.2008.07.012
- 681 Rodriguez-Lanetty M, Phillips W, and Weis V. 2006. Transcriptome analysis of a cnidarian -682 dinoflagellate mutualism reveals complex modulation of host gene expression. BMC 683 Genomics 7:23. DOI: 10.1186/1471-2164-7-23
- Ryu T, Seridi L, Moitinho-Silva L, Oates M, Liew YJ, Mavromatis C, Wang X, Haywood A, 684 685 Lafi FF, Kupresanin M, Sougrat R, Alzahrani MA, Giles E, Ghosheh Y, Schunter C, 686 Baumgarten S, Berumen ML, Gao X, Aranda M, Foret S, Gough J, Voolstra CR, Hentschel U, and Ravasi T. 2016. Hologenome analysis of two marine sponges with 687 688 different microbiomes. BMC Genomics 17:1-11. DOI: 10.1186/s12864-016-2501-0
 - Sarrias M, Grønlund J, and Padilla O. 2004. The scavenger receptor cysteine-rich (SRCR) domain: an ancient and highly conserved protein module of the innate immune system. Critical Reviews in Immunology 24:1-37. DOI: 10.1615/CritRevImmunol.v24.i1.10
 - Schneider DS, and Ayres JS. 2008. Two ways to survive infection: what resistance and tolerance can teach us about treating infectious diseases. Nature Reviews Immunology 8:889-895. DOI: 10.1038/nri2432
 - Schwarz RS, Hodes-Villamar L, Fitzpatrick KA, Fain MG, Hughes AL, and Cadavid LF. 2007. A gene family of putative immune recognition molecules in the hydroid *Hydractinia*. Immunogenetics 59:233-246. DOI: 10.1007/s00251-006-0179-1
 - Shinzato C, Shoguchi E, Kawashima T, Hamada M, Hisata K, Tanaka M, Fujie M, Fujiwara M, Koyanagi R, Ikuta T, Fujiyama A, Miller DJ, and Satoh N. 2011. Using the Acropora digitifera genome to understand coral responses to environmental change. Nature 476:320-323. DOI: 10.1038/nature10249
- Silverstein RL, and Febbraio M. 2009. CD36, a scavenger receptor involved in immunity, 703 metabolism, angiogenesis, and behavior. Science Signaling 2:re3. DOI: 704 10.1126/scisignal.272re3
 - Thelen T, Hao Y, Medeiros AI, Curtis JL, Serezani CH, Kobzik L, Harris LH, and Aronoff DM. 2010. The class A scavenger receptor, macrophage receptor with collagenous structure is the major phagocytic receptor for Clostridium sordellii expressed by human decidual macrophages. Journal of Immunology 185:4328-4335. DOI: 10.4049/jimmunol.1000989
- 709 Vidal-Dupiol J, Adjeroud M, Roger E, Foure L, Duval D, Mone Y, Ferrier-Pages C, Tambutte E, 710 Tambutte S, Zoccola D, Allemand D, and Mitta G. 2009. Coral bleaching under thermal 711 stress: putative involvement of host/symbiont recognition mechanisms. BMC 712 Physiolology 9:14. DOI: 10.1186/1472-6793-9-14
- 713 Weis VM, and Allemand D. 2009. What determines coral health? Science 324:1153-1155. DOI: 714 10.1126/science.1172540
- 715 Wood-Charlson EM, and Weis VM. 2009. The diversity of C-type lectins in the genome of a 716 basal metazoan, Nematostella vectensis. Developmental and Comparative Immunology 717 33:881-889. DOI: 10.1016/j.dci.2009.01.008
- 718 Yellowlees D, Rees TAV, and Leggat W. 2008. Metabolic interactions between algal symbionts 719 and invertebrate hosts. Plant Cell & Environment 31:679-694. DOI: doi:10.1111/j.1365-720 3040.2008.01802.x
- 721 Yuen B, Bayes JM, and Degnan SM. 2014. The Characterization of Sponge NLRs Provides 722 Insight into the Origin and Evolution of This Innate Immune Gene Family in Animals.
- 723 Molecular Biology and Evolution 31:106-120. DOI: 10.1093/molbev/mst174



724	Zani IA, Stephen SL, Mughal NA, Russell D, Homer-Vanniasinkam S, Wheatcroft SB, and
725	Ponnambalam S. 2015. Scavenger receptor structure and function in health and disease.
726	Cells 4:178-201. DOI: 10.3390/cells4020178
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Table 1. Information on the cnidarian sequence resources used in this study. Non-symbiotic refers to species that do not form symbioses with dinoflagellates. Aposymbiotic refers to species that do form symbioses but the material from which the sequencing was performed did not contain symbionts.

Organism	Developmental stage	Symbiotic state	Data type	Reference
Nematostella vectensis	Larvae	Non-symbiotic	Genome	(Putnam et al. 2007)
Anthopleura elegantissima	Adult	Aposymbiotic	Transcriptome	(Kitchen et al. 2015)
Aiptasia pallida	Adult	Aposymbiotic	Transcriptome	(Lehnert et al. 2012)
Aiptasia pallida	Adult	Aposymbiotic	Genome	(Baumgarten et al. 2015)
Acropora digitifera	Sperm	Aposymbiotic	Genome	(Shinzato et al. 2011)
Acropora millepora	Adult and larvae	Symbiotic	Transcriptome	(Moya et al. 2012)
Fungia scutaria	Larvae	Aposymbiotic	Transcriptome	(Kitchen et al. 2015)



Figure Legends

Figure 1. Domain architecture of vertebrate SRs relevant to this study. All SR sequences are anchored in the membrane with one or two transmembrane domains. All have very short cytoplasmic tails and extensive extracellular ligand-binding domains. SR-As contain a collagen domain(s) and can include an SRCR or a CTLD. SR-Bs have two cytoplasmic tails on either side of a CD36 domain that forms an extracellular loop. SR-Es are defined by the presence of a CTLD. SR-Is have multiple SRCR repeats and no other identifiable extracellular domains. C, carboxy terminus; CTLD, C type lectin domain; LOX1, lectin-like oxidized low density lipoprotein receptor 1; MARCO, macrophage receptor with collagenous structure; N, amino terminus; SRCL, scavenger receptor with C-type lectin; SRCR, scavenger receptor cysteine-rich

domain.

Figure 2. Domain architecture of cnidarian SR domains in the six resources searched compared to human SRs. Identified cnidarian SR-A-like and SR-E-like sequences display diverse domain architecture and include novel domain combinations not found in vertebrates. SR-I-like sequences had a varying number of SRCR repeats. A variety of SRCR-domain-containing cnidarian sequences identified did not fit the criteria of any vertebrate SR classes and are presented as SRCR + CUB domains or SRCR + trypsin domains. SR-B-like domain combinations closely resembled vertebrate SR-Bs with two transmembrane domains, two cytoplasmic tails and a CD36 domain. CTLD, C type lectin domain; CUB, complement C1r/C1s, Uegf, BMP1; DMBT, deleted in malignant brain tumor protein; Ig, immunoglobulin; I-Set, intermediate set of immunoglobulin domain; LDL, low density lipoprotein; LOX1, lectin-like oxidized low density lipoprotein receptor 1; EGF, epidermal growth factor; MAM meprin/A5-protein/PTPmu; MARCO, macrophage receptor with collagenous structure; SCARA5, scavenger receptor class A member 5; SRCL, scavenger receptor with C-type lectin; SRCR, scavenger receptor cysteine-rich domain; U-box, ubiquitin box. Human SR data taken from Canton et al (2013). (See Supplementary File 1 for sequence information.)

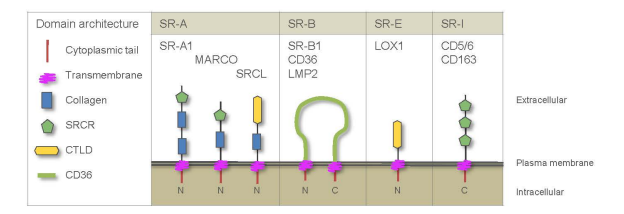
Figure 3. Maximum-likelihood tree of SR-Bs from across the Metazoa. The tree was constructed with the CD36 domain of each protein using FastTree v 2.1.5. Bootstrap support values were generated using SEQBOOT, values above 0.6 are displayed at nodes. The alignment, including organism names, is displayed in Figure S1.

Figure 4. Experimental colonization by *S. minutum* CCMP830 of aposymbiotic *A. pallida* treated with increasing levels of the SR inhibitor, fucoidan. Graph shows percent colonization success as measured by surface area of host gastrodermis occupied by symbionts (see Methods for details) as a function of time after inoculation. Two controls were included: FSW alone and an 18 h incubation in 200 μg/ml fucoidan in FSW followed by a 48 h recovery in FSW to test for fucoidan toxicity to the animals. Anemones in experimental fucoidan treatments exhibited a



777	dose-dependent response with decreased colonization success with increasing fucoidan
778	concentrations. Bars represent means \pm SD, n = 3 anemones per treatment. Asterisks indicate
779	high (p $>$ 0.999) posterior probability of treatment effects being different from controls under the
780	Bayesian ANOVA model.
781	
782	Figure 5. Effect of SR inhibition by fucoidan on immune stimulation in symbiotic <i>A. pallida</i> .
783	Immune stimulation of animals was elicited by incubation in 1 µg/ml LPS overnight prior to the
784	experiment. Immune stimulation was measured by quantifying DAF-FM DA, a probe for the
785	presence NO, itself a marker for immune stress. Graph shows MFI of DAF-FM DA in tentacles
786	in response to incubation in increasing concentrations of fucoidan. Animals exhibited a
787	significant dose-dependent response to fucoidan (Bayesian generalized linear mixed model, P <
788	0.0001), with increasing NO production with increasing SR inhibition by fucoidan. Bars
789	represent means \pm SD; n = 3 anemones. Inset: representative confocal images of tentacles
790	incubated in FSW only and 800 μg/ml fucoidan. DAF-FM DA (green) symbiont
791	autofluorescence (red).
792	
793	







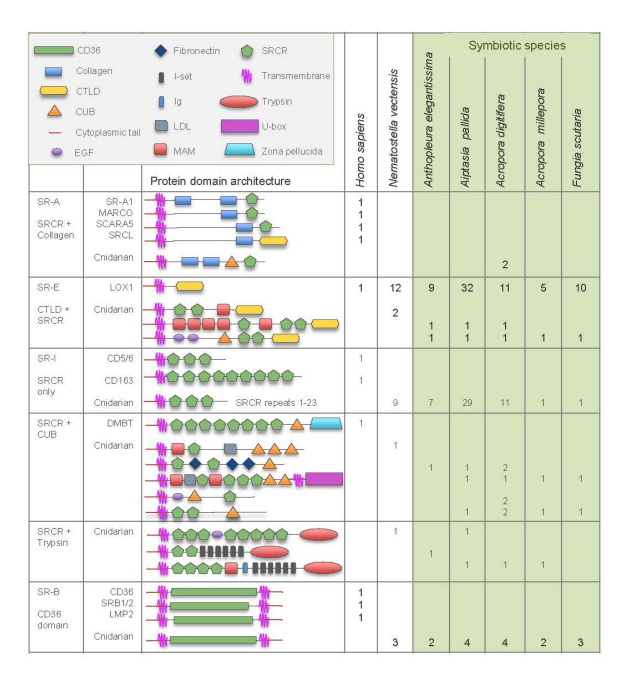


Figure 2



