Insights into the innate immunome of actiniarians using a comparative genomic approach

Chloé A. van der Burg 1,2*, Peter J. Prentis 3,4, Joachim M. Surm 1,2 and Ana Pavasovic 1,2

Abstract

Background: Innate immune genes tend to be highly conserved in metazoans, even in early divergent lineages such as Cnidaria (jellyfish, corals, hydroids and sea anemones) and Porifera (sponges). However, constant and diverse selection pressures on the immune system have driven the expansion and diversification of different immune gene families in a lineage-specific manner. To investigate how the innate immune system has evolved in a subset of sea anemone species (Order: Actiniaria), we performed a comprehensive and comparative study using 10 newly sequenced transcriptomes, as well as three publically available transcriptomes, to identify the origins, expansions and contractions of candidate and novel immune gene families.

Results: We characterised five conserved genes and gene families, as well as multiple novel innate immune genes, including the newly recognised putative pattern recognition receptor CniFL. Single copies of TLR, MyD88 and NF-κB were found in most species, and several copies of IL-1R-like, NLR and CniFL were found in almost all species. Multiple novel immune genes were identified with domain architectures including the Toll/interleukin-1 receptor (TIR) homology domain, which is well documented as functioning in protein-protein interactions and signal transduction in immune pathways. We hypothesise that these genes may interact as novel proteins in immune pathways of cnidarian species. Novelty in the actiniarian immunome is not restricted to only TIR-domain-containing proteins, as we identify a subset of NLRs which have undergone neofunctionalisation and contain 3–5 N-terminal transmembrane domains, which have so far only been identified in two anthozoan species.

Conclusions: This research has significance in understanding the evolution and origin of the core eumetazoan gene set, including how novel innate immune genes evolve. For example, the evolution of transmembrane domain containing NLRs indicates that these NLRs may be membrane-bound, while all other metazoan and plant NLRs are exclusively cytosolic receptors. This is one example of how species without an adaptive immune system may evolve innovative solutions to detect pathogens or interact with native microbiota. Overall, these results provide an insight into the evolution of the innate immune system, and show that early divergent lineages, such as actiniarians, have a diverse repertoire of conserved and novel innate immune genes.

Keywords: Actiniaria, Innate immune system, Transcriptomics, Evolution, TIR domain, NLR, Novel immune genes, Complement cascade, CniFL

* Correspondence: chloe.vanderburg@hdr.qut.edu.au
1School of Biomedical Sciences, Faculty of Health, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia
2Institute of Health and Biomedical Innovation, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia
Full list of author information is available at the end of the article

© The Author(s). 2016 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background
The immune system is an ancient and complex system that works to detect and defend against pathogens, as well as to regulate the interaction between microbes and hosts [1–4]. Defence against pathogens in vertebrates is a two-fold mechanism. It consists of the innate immune system, which provides non-specific protection to the host, and the adaptive immune system, which mounts a specific attack against foreign bodies (e.g., microbes) and displays immunological memory [1–4]. Vertebrates, such as cnidarians (corals, jellyfish, hydroids and sea anemones), however, only possess the innate immune system as their primary mode of pathogen defence [5].

Innate immune genes tend to be highly conserved in metazoans, even in early divergent lineages, such as Cnidaria and Porifera [5–9]. However, constant and diverse selection pressures on the immune system have driven the expansion and diversification of different immune gene families in a lineage-specific manner [10–13]. One particular example is the NOD-like receptor family (renamed by [14] as Nucleotide-binding and Leucine-rich Repeat containing gene family (NLR)), of which expansions have occurred independently in multiple early divergent lineages [7, 12, 15, 16].

The evolution of immune genes can be complex, as not all functionally conserved immune genes are homologous. Protein domains may evolve independently and subsequently converge on a conserved function or architecture [12, 17]. For example, the functional Toll-like receptor pathway in Hydra magnipapillata (freshwater polyp) is formed using a scaffold assembly of TIR-only proteins (Toll/interleukin-1 receptor homology domain) and LRR-only proteins (Leucine-rich repeat) [18–20]. Similarly, it appears that the immunoglobulin-TIR domain combination found in the interleukin receptor family has evolved independently in cnidarians and vertebrates, as these genes share limited sequence similarity [5, 11, 16]. For these reasons, investigating conserved domain architectures can be highly informative in identification and characterisation of the immune repertoire in cnidarians and other early divergent metazoan taxa. Using this approach, the evolution of novel immune genes can also be investigated, through interrogation of novel architectures with domains that have known immune functions. The TIR domain is a well-characterised example; it functions in protein-protein interactions and signal transduction in immune pathways. Such an approach has previously [11] been successfully applied to identify novel immune genes by interrogating TIR-domain-containing architectures.

Like other members of phylum Cnidaria, actiniarians are anatomically simple and develop from only two germ layers (diploblastic). These animals are typically sedentary with no physical protective barriers such as a shell, cuticle or exoskeleton and are therefore directly exposed to the abiotic and biotic conditions surrounding them. Consequently, these organisms have evolved immune defence mechanisms that tend to rely on mucous secretions which consist of antimicrobial peptides, as well as a diverse range of pattern recognition receptors (PRRs) which work in concert with the glycocalyx, to provide a physicochemical barrier [21, 22]. As with other eumetazoans, pathogen recognition in actiniarians is thought to occur primarily via the detection of pathogen associated molecular patterns (PAMPs), using a diverse array of PRRs. Cnidarian immune genes, in particular PRRs, also have a major role in maintaining homeostasis between the host and the beneficial native microbiota, which primarily reside on the epithelium [19, 23], although many cnidarians also undergo endosymbiosis with dinoflagellates [24].

Current genomic resources for cnidarians have been limited to a few key model species, including Nematostella vectensis (starlet sea anemone) [6], Acropora digitifera (coral) [25], H. magnipapillata. [26] and Aiptasia sp. (sea anemone) [27]. Interrogation of these genomic resources has revealed that the cnidarian genome is surprisingly complex. In fact, cnidarians have maintained both eumetazoan and early divergent metazoan gene families, some of which have been lost in invertebrate models Drosophila melanogaster and Caenorhabditis elegans [5, 9, 28–30]. In particular, the genome of the sea anemone Nematostella, has been shown to be more similar to vertebrates than other invertebrate model species, with gross genomic structure and exon/intron arrangement similar to vertebrates [6]. While these few cnidarian species have been crucial in elucidating the complexity of the cnidarian genome, in this highly speciose phylum, more resources are needed to facilitate further in-depth comparative evolutionary and phylogenetic analyses.

Actiniarians are uniquely suited for the study of the origin and evolution of the innate immune system. This is principally attributed to their phylogenetic placement as members of Cnidaria, the sister phylum to Bilateria. Their simple anatomical features and complex, vertebrate-like genomes make Actiniarians an interesting candidate for understanding how novel innate immune genes evolve within eumetazoan species that have limited physical defences. Furthermore, using Actiniarians as a model for understanding anthozoan immunity will help provide insights into other issues facing anthozoan species, such as coral bleaching and disease. To better understand the evolution of the innate immune system, here we performed a comprehensive and comparative investigation of the actiniarian innate immune gene repertoire with the aim to provide preliminary insights into the key processes that have shaped the cnidarian immunome. Specifically, we
have examined gene family evolution of candidate immune genes, to better understand the role of gene gain and loss events, the generation of new genes and the acquisition of genes from other species, as key drivers of the innate immune gene repertoire in actiniarians.

**Results**

**Sequencing, assembly and annotation**

Whole organism extractions of total RNA were performed from *Actinia tenebrosa* (red, brown, green and blue colourmorphs, designated 1, 2, 3 and 4 respectively) *Anthopleura buddemeieri*, *Aulactinia veratra*, *Calliactis polypus* (designated 1 and 2), *Telmatactis* sp. and *Nemanthus annamensis*. Sequencing was performed on an Illumina NextSeq 500, and raw reads per sample ranged between 51,620,970 and 220,632,160. These reads were assembled into 116,120–212,774 contiguous sequences (contigs) for each transcriptome. All raw reads were submitted to the NCBI Sequence Read Archive (SRA) under one BioProject (PRJNA313244); accession numbers for individual transcriptomes can be viewed in Additional file 1: Table S1. Assembly statistics as well as completeness scores can be found in Additional file 1: Tables S2 and S3, which also includes assembly metrics for raw reads downloaded from the SRA for *Aiptasia pallida*, *Anthopleura elegantissima* and *Nematostella vectensis*. A total of 13 transcriptomes were used in this study, from 9 actiniarian species.

**Gene ontology**

Gene ontology (GO) terms were assigned to between 21,750 and 43,791 transcripts using Trinotate (Additional file 1: Table S4). WEGO analysis indicated a similar distribution of GO terms among assemblies (Additional file 1: Figure S1). CateGoRizer analysis showed that the number of genes assigned immune class GOs ranged from 7239 to 12,515 across species, with the exception of *A. pallida* which had only 1500 genes (Additional file 1: Table S5 and Figure S2). Stress response and protein metabolism were the most frequently assigned immune class GO terms for all species, while the ‘immunology, immune response’ term corresponded to approximately 2.7 % of the total number of genes in this GO category, except *A. pallida* which had 0.5 %. The results from CateGoRizer showed that most assemblies have a similar distribution of GO terms associated with the immune class. *A. pallida* showed the most variation from the other transcriptomes, with a much lower fraction of genes in all categories except categories related to metabolism (i.e., catabolism and protein, lipid and carbohydrate metabolism).

**RSEM**

RNA-Seq by Expectation-Maximization (RSEM) analysis of transcriptomes was performed to verify candidate transcripts are expressed and as a guide for choosing the most highly expressed candidate genes for PCR validation i.e., transcripts with the highest FPKM (Fragments Per Kilobase of transcript per Million mapped reads) score were chosen in most instances for PCR validation. The most highly expressed immune transcripts in all transcriptomes for which RSEM was performed (*A. tenebrosa* (1), *A. buddemeieri*, *A. veratra*, *C. polypus* (2)) were all NLR transcripts, with FPKMs of 84.69 for *A. tenebrosa*, 22.76 for *A. buddemeieri*, 65.74 for *A. veratra* and 370.7 for *C. polypus*.

**Candidate gene counts and architecture**

Candidate gene analysis revealed that for most species, a single gene copy of *TLR*, *MyD88* and *NF-kB* was identified (Table 1), with very few species having multiple isoforms of these genes (Additional file 2: Table S6). In contrast, multiple copies of *NLR* and *IL-1R*-like were found in all species. Complete copies of *NLR* genes were found in most species (with the exception of *A. elegantissima* and *Telmatactis* sp., where only partial sequences were found) ranging from 2 to 4 complete and 1–8 partial copies of *NLR* genes, with total (complete and partial) gene copy numbers ranging from 3 to 10 (Table 1 and Additional file 2: Table S7). Complete copies of *IL-1R*-like genes were found in all species, ranging

**Table 1 Candidate gene counts in each transcriptome**

| Species                  | TLR | MyD88 | NF-kB | NLR | IL-1R-like |
|--------------------------|-----|-------|-------|-----|------------|
| *A. tenebrosa* (1)       | 1   | 1     | 1     | 2   | 3          |
| *A. tenebrosa* (2)       | 1   | 1     | 1     | 2   | 7          |
| *A. tenebrosa* (3)       | 1   | 1     | 1     | 2   | 4          |
| *A. tenebrosa* (4)       | 1   | 1     | 1     | 4   | 3          |
| *A. pallida*             | -4  | 1     | 0*    | 2   | 2          |
| *A. buddemeieri*         | 1   | 1     | 1     | 3   | 2          |
| *A. elegantissima*       | 1   | 1     | 1     | 1†  | 3          |
| *A. veratra*             | 1   | 1     | 1     | 2   | 4          |
| *C. polypus* (1)         | 0   | 1     | 1     | 2   | 3          |
| *C. polypus* (2)         | 1   | 1     | 1     | 4   | 5          |
| *N. annamensis*          | 1   | 1     | 2     | 2   | 4          |
| *N. vectensis*           | 1   | 1     | 0*    | 3   | 2          |
| *Telmatactis* sp.        | 0   | 0     | 1     | 1†  | 1          |

Counts are conservative, i.e., they refer to the number of genes with a complete ORF only, not isoforms or partial genes found in each transcriptome (see Additional file 2: Tables S6 and S7 for gene, isoform and partial gene counts). Genes were considered complete when a full open reading frame was found (start and stop codons) and contained the canonical Pfam domains (see Methods). Shown are the Toll-like receptor (*TLR*), Nucleotide-binding and Leucine-rich Repeat containing gene family (*NLR*), Myeloid Differentiation primary response gene 88 (*MyD88*) and Nuclear Factor kappa-light-chain-enhancer of activated B cells (*NF-kB*) and Interleukin-1 receptor (*IL-1R-like*). *No TLR in Aiptosis genome [27]. †NF-kB in Nematostella does not have ankyrin repeats [71], here NF-kB was only considered complete if it contained the RHD and ankyrin repeats (see Methods). *Aiptosis also has only a RHD-IPT (no ankyrin repeats) containing NF-kB [AIPGENE8848 sp|P19838|NFKB1]. †Partial genes found, see Additional file 2: Table S7.
from 1 to 7 complete gene copies and 3–7 total gene copies when considering partial sequences as well (Table 1 and Additional file 2: Table S7).

All TLR and IL-1R-like predicted peptides had transmembrane domains (TMDs) and all MyD88 and NF-κB predicted peptides did not. Transmembrane domain analysis of NLRs, however, revealed 8 out of 35 total (including complete and partial) NLRs had TMDs. Multiple (3–5) TMDs were identified in TMD containing NLRs and were clustered at the N-terminus. TMD containing NLRs were found in *A. pallida*, *A. tenebrosa*, *A. buddemeieri* and *C. polypus*. The most common protein domain architecture of all candidate genes and their hypothesised location in the cell are shown in Fig. 1.

**Candidate and novel gene validation**

To validate candidate and novel genes, PCR amplification from *A. tenebrosa*, *A. buddemeieri*, *A. veratra* and *C. polypus* was performed. PCR products were sequenced and realigned to the contig from which primers were designed for each transcript. Sequences were considered complete when they aligned to the transcript with at least 94 % pairwise sequence similarity to the ORF, following trimming. A total of 15 complete and 3 partial sequences were validated, with at least one

![Fig. 1](https://example.com/fig1.png)

**Fig. 1** Typical protein domain architecture for the five candidate genes and their hypothesised cellular location. Structures depicted show typical architecture; other variations exist. Depictions based on Pfam annotations and SMART visualisation. Shown are: TLR, Toll-like receptor; NLR, Nucleotide-binding domain and Leucine-rich Repeat containing gene family; MyD88, Myeloid Differentiation primary response gene 88; NF-κB, Nuclear Factor kappa-light-chain-enhancer of activated B cells; IL-1R-like, Interleukin-1 receptor. Depictions of ankyrin repeats, LRRs, TMD and Igs do not reflect the exact number in each gene.
representative sequence from each of the 5 candidate genes, and one representative sequence from each of the 3 novel genes. All validated sequences were submitted to GenBank® (NCBI) and accession numbers can be viewed in Additional file 3: Table S14.

**Presence/absence of other innate immune genes**

Additional innate immune genes were investigated in this study for their presence/absence in actiniarian species. These included key receptors and proteins in the complement pathway (i.e., CniFL, which is hypothesised to function in the lectin-complement pathway and complement proteins MASP, C3, Factor B, C6 and Factor I), as well as genes encoding protein domains known to have immune function (i.e., SRCR and C-type Lectin domains). The results are presented in Table 2.

**Identification of novel immune genes**

Taxonomically-restricted transcripts with novel domain architectures were identified in all species, but not all novel transcripts were found in all species examined (Additional file 2: Tables S8–S10). Novel genes with potential immune function were identified based on novel architectures that included the TIR (or TIR_2) domain. A representation of the full repertoire of these genes is shown in Fig. 2. Three of the most common transcripts were investigated further and designated as Novel Genes 1, 2 and 3.

Novel Gene 1 (NG1) was the most commonly observed transcript with novel domain architecture and was found in all species examined. NG1 includes the Pfam domains Leucine-rich repeat (LRR), Miro-like protein (Miro), Ras family (Ras) and Toll/Interleukin-1 receptor homology domain (TIR_2). SMART domain annotation represented the Miro and Ras domains as either Ras family, Ras of complex (Roc) or small GTPase, all of which are GTPase family domains. A low threshold C-terminal of Roc (COR) domain was also found ~300 aa downstream in all genes, which may be essential to the function of the GTPase domain and is always found downstream of Ras [31]. NG1 is represented in Fig. 2a.

BLASTx analysis of NG1 against the TrEMBL database (Additional file 2: Table S9) identified the only significant hit as a *Nematostella* predicted protein. Swiss-Prot BLASTx searches (Additional file 2: Table S9) identified this gene as an F-box/LRR (FBXL) protein from multiple vertebrate species. The FBXL protein, however, only has the LRR region in common with NG1, indicating there is no ortholog of NG1 in vertebrates. Further investigation of this gene with known TIR and TIR_2 architectures from the EMBL-EBI [32] domain database identified 362 unique architectures with the TIR_2 domain (PF13676) and 381 unique architectures with the TIR domain (PF01582), but the TIR domain was not found in combination with any GTPase (Roc, Ras or COR). The TIR_2 domain was identified in combination with one or more of the GTPase domains in multiple bacteria species, including cyanobacteria, filamentous bacteria, purple sulfur bacteria and flavobacteria, and with an additional C2 domain in the pacific oyster, *Crassostrea gigas* (C2 domain – targets proteins to cell membranes), and a MBT domain in the placozoan, *Trichoplax adhaerens* (MBT – repeat of unknown function). Table S10 (Additional file 2) provides details of the variable domain architectures found that could represent orthologous genes to NG1.

### Table 2 Gene counts and presence/absence of other innate immune genes

| Species            | CniFL | MASP | SRCR | C3 family | Factor B (Bf) family | C6 family | Factor I (f) family |
|--------------------|-------|------|------|-----------|----------------------|-----------|-------------------|
| *A. tenebrosa* (1) | 5*    | 1    | 114  | 83        | +                    | +         | -                 |
| *A. tenebrosa* (2) | 7     | 1    | 131  | 89        | +                    | +         | -                 |
| *A. tenebrosa* (3) | 2     | 1    | 121  | 76        | +                    | +         | -                 |
| *A. tenebrosa* (4) | 2     | 1    | 105  | 70        | +                    | +         | -                 |
| *A. pallida*       | 4     | 1    | 51   | 87        | +                    | +         | -                 |
| *A. buddemeieri*   | 2*    | 1    | 135  | 88        | +                    | +         | -                 |
| *A. elegantissima*| 0     | 1    | 60   | 57        | +                    | +         | -                 |
| *A. veratra*       | 5     | 1    | 129  | 111       | +                    | -         | -                 |
| *C. polypus* (1)  | 6     | 1    | 61   | 65        | +                    | +         | -                 |
| *C. polypus* (2)  | 5     | 1    | 74   | 84        | +                    | +         | -                 |
| *N. annamensis*    | 3     | 1    | 39   | 57        | +                    | -         | -                 |
| *N. vectensis*     | 1     | 1    | 39   | 67        | +                    | -         | -                 |
| *Telmatactis* sp. | 2     | 1    | 59   | 41        | +                    | -         | -                 |

Gene counts shown are non-conservative, i.e., they refer to the number of contigs identified by Trinity as different genes, and may include contigs with partial ORFs. Only complete ORFs used in subsequent analysis for CniFL genes. *PCR validated. + present; - absent. Abbreviations are as follows: CniFL, Cnidarian Ficolin-like protein; SRCR, Scavenger-Receptor Cysteine Rich; MASP, Mannan-binding lectin (or mannose-associated) Serine Protease 1
Novel Gene 2 (NG2) was identified in six species, *A. tenebrosa* (2, 3 and 4), *A. pallida*, *A. elegantissima*, *C. polypus* (2), *N. annamensis* and *N. vectensis*, and included one or two TIR_2 domains upstream of a BTK motif (Bru-ton’s tyrosine kinase-type zinc-finger motif). Only a single predicted *Nematostella* protein was identified in TrEMBL and no proteins were identified in Swiss-Prot (Additional file 2: Table S9). No sequences in EMBL-EBI were identified with a BTK motif and TIR domain and only two sequences containing both a TIR_2 and BTK motif were found in EMBL-EBI. These predicted proteins were found in *Phytophthora ramorum* (oomycete) and *Guillardia theta* (cryptomonad algae) (UniProt IDs in Additional file 2: Table S10). NG2 is represented in Fig. 2b.

Novel Gene 3 (NG3) was identified in only three species, *A. pallida*, *C. polypus* and *N. vectensis*. NG3 includes the Pfam domains protein tyrosine kinase or protein kinase (Pkinase_Tyr or Pkinase; always overlapping), a death domain and a TIR (or TIR_2) domain. Additional domains are seen in this gene when represented by SMART; these included central Roc/COR domains and a SH3 domain (SRC Homology 3 domain). Only a single *Nematostella* predicted protein was found to match this transcript in the TrEMBL database (Additional file 2: Table S9). BLASTx analyses against the Swiss-Prot database (Additional file 2: Table S9) identified two slime mould predicted proteins with similarity; however, neither contained either the DEATH or TIR domains, and were therefore disregarded. EMBL-EBI
database search returned no TIR domain architectures that were similar to NG3. A few sequences were found in the EMBL-EBI database with TIR_2 domain combinations similar to NG3. The most similar domain combination found was with a ‘neuralized’ domain in place of the ‘Pkinase’ and ‘SH3’ domains in *Strigamia maritima* (centipede) and *Crassostrea gigas* (pacific oyster). The ‘Pkinase’ and ‘SH3’ domains are also missing in a less similar domain architecture found in *Strongylocentrotus purpuratus* (purple sea urchin) (Additional file 2: Table S10). NG3 is represented in Fig. 2c.

**Evolutionary and Phylogenetic analyses**

**Non-synonymous vs synonymous substitution rates**
Non-synonymous vs synonymous substitution rates (d_{NS}/d_{S}) were calculated using PAML for all TLR, *MyD88*, *NF-κB* and TMD containing *NLRs* genes, as well as for Novel Genes 1, 2 and 3. Table 3 shows a summary of the results and all PAML results for the Yang and Nielsen [33] method can be viewed in Additional file 4. Conserved and novel genes where all found to have patterns of synonymous and non-synonymous mutations consistent with the action of purifying selection (i.e., dN/dS ratio < 1) across all genes.

**NLR family phylogenetic analysis**
Maximum likelihood analysis was used to infer phylogenetic relationships of NLRs between actinarian species in this study with a selection of additional metazoan NLRs. Figure 3 shows a radial maximum likelihood tree, generated using a LG + G protein model. Highlighted on the tree are two clades, which correspond to NLRs with certain classes of N-terminal domains. All known metazoan TMD containing NLRs cluster into one clade, which only consists of anthozoan species (*Actiniaria* and *Scleractinia* (*A. digitifera*)). Most of the other anthozoan NLRs all cluster into another clade. Interestingly the DED sequences from anemone species from this study cluster more closely with the Chordata, Annelida and Mollusca BIR, CARD and DD clade, than with other anthozoan species (i.e., *N. vectensis* and *A. digitifera*), although this is lowly supported. *N. vectensis* NLRs consistently cluster outside of other actinarian species with high bootstrap support.

**CniFL family phylogenetic analysis**
A total of 42 protein sequences from all species except *A. elegantissima* and *Telmatactis* sp, were identified with a complete ORF. Removing duplicate sequences resulted in a final alignment of 30 sequences, which was trimmed to remove poorly aligned regions to a final 132 aa alignment. The phylogeny represented in Fig. 4 was generated using the JTT + G protein model of evolution and the alignment used can be found in Additional file 5. Most CniFL genes were identified with an additional TMD which is represented in the domain architecture illustrations in Fig. 4. One clade also contains CniFLs with an additional domain WAP (whey acidic protein) found in *N. vectensis*, *A. tenebrosa* and *C. polypus*. Mapping back the *N. vectensis* CniFL transcript to the genome (v1.0) identified the *N. vectensis* CniFL gene at scaffold_24:467371-486150 (gene model: e_gw.24.49.1). Multiple Ig domains were annotated by Pfam in the genome; however, they are annotated in a region that has been annotated as intronic.

**Table 3**

| Gene                  | d_{NS} (±SE) | d_{S} (±SE) | d_{NS}/d_{S} |
|-----------------------|--------------|-------------|--------------|
| TLR                   | 0.3610 (±0.0162) | 1.5845 (±0.2657) | 0.2278       |
| *MyD88*               | 0.3140 (±0.0298) | 2.4488 (±1.1913) | 0.1283       |
| *NF-κB*               | 0.2938 (±0.0299) | 2.4713 (±2.9559) | 0.1189       |
| TMD containing *NLRs* | 0.5990 (±0.0244) | 1.8720 (±0.6182) | 0.3200       |
| Novel Gene 1          | 0.1757 (±0.0083) | 1.4509 (±0.1338) | 0.1211       |
| Novel Gene 2          | 0.2480 (±0.0145) | 2.2503 (±0.4263) | 0.1102       |
| Novel Gene 3          | 0.3757 (±0.0144) | 3.9824 (±8.6207) | 0.0943       |

All values show as averages for each gene, calculated using the Yang and Nielsen [33] method. Average d_{NS}/d_{S} ratio does not take into account standard error.

**Discussion**
To better understand the evolution of the innate immune system we have performed a comprehensive and comparative investigation of the actinarian innate immune gene repertoire, including TLR, *NLR,IL-1R*-like, *MyD88*, *NF-κB*, *CniFL*, *SRCR*, MASP; C-type lectin domains, complement control proteins and multiple novel innate immune genes. By generating multiple high-quality transcriptome assemblies we were able to undertake a detailed study of innate immune genes in actinarians. Here we note that in some cases specific gene families appear to have undergone lineage-specific expansions and potential neofunctionalisation (e.g. *NLR*), while other genes such as *TLR* have fewer copies than observed in other eumetazoan phyla. Although these observations need further functional validation, they provide an insight into the core eumetazoan immune gene set and show that the majority of actinarian innate immune genes show evidence of purifying selection.

**Candidate and novel gene counts and architectures**
The identification of candidate and novel genes in this study is mostly consistent with previously reported information on gene copy number in cnidarians. For example [11], reported similar counts for some TIR domain containing proteins identified in nine anthozoan species. Their study included three of the species interrogated in our study: *A. pallida*, *A.
elegantissima and N. vectensis and confirmed single copies of TLR and MyD88. For IL-1R-like, of which there are multiple copies, fewer complete genes were identified in our study for the three species, whilst numbers of all TIR domain containing proteins were higher. The study [11] also identified an expansion of TIR only proteins restricted to coral species. Here we present evidence for an anthozoan-specific expansion of TIR only proteins, as we identify similar counts of TIR only proteins in all anemone species (Additional file 2: Table S8).

In this study, the TIR domain, not TIR_2, was always found in proteins with evolutionarily conserved architectures (i.e., TLR, MyD88, IL-1R-like). Notably, in most instances TIR_2, a common bacterial TIR domain [34] also found in some marine invertebrates [35–37] was found in novel genes identified here. This highlights that horizontal gene transfer (HGT) may have played a role in the introduction of the TIR_2 domain to the genome of actiniarian species, which then underwent domain shuffling to form the observed novel genes. Whilst the possibility of contamination cannot conclusively be ruled out, in most instances no similar domain architectures to the three novel genes were identified in any other species. This study identifies two previously uncharacterised genes (NG1 and NG2) and extends on the previously reported novel genes for Actiniarians [11], including NG3 and other novel genes shown in Fig. 2, which have been
identified by [11]. Interestingly, NG3 identified here was found with additional domains (SH3, Roc and COR domains) not identified by [11]. These novel domain architectures found in lineage-specific actiniarian species are likely the result of domain shuffling.

Expanded repertoire and neofunctionalisation of NLRs

The diversity in domain architectures observed in the NLR family across metazoans suggests potential evidence for neofunctionalisation in this gene family. The expansion and diversification of actiniarian NLRs, such
as those with transmembrane domains, observed in our study is significant as it indicates that some of these duplicated copies have likely undergone independent neofunctionalisation within Anthozoa. This is supported by phylogenetic analysis of NLRs across Metazoa (Fig. 3) where all 15 TMD containing NLRs clustered into one clade, but other actiniarian NLRs clustered in separate clades. The presence of multiple transmembrane domains in NLRs identified in our study indicates that they may be membrane-bound. This appears to be an anthozoan specific innovation as all other metazoan NLRs are exclusively localised to the cytoplasm and do not have these domains [38]. Interestingly, only two other anthozoan species (N. vectensis and A. digitifera) have so far been reported with TMD containing NLRs [7], but our results indicate a wide distribution of TMD containing NLRs across multiple actiniarian species. This highlights that novel innovations and neofunctionalisation have occurred in innate immune gene families of Actiniaria. *Amphimedon queenslandica* (sponge) NLRs may also contain TMDs, however, no anthozoan TMD containing NLRs cluster with these sequences and do not currently support this hypothesis [7]. Phylogenetic analysis of the metazoan NLRs resulted in a similar tree to the one obtained by [7] in their study on sponge NLRs, providing further evidence that the TMD containing NLR clade are correctly clustered into a single clade, indicating this novel structure has likely evolved from a single origin within Anthozoa. Potential functional implications of the neofunctionalisation of NLRs include allowing cooperation between other pattern recognition receptors or a modified or novel response to an immune signal.

**Conservation of innate immune pathways and the interaction of novel genes**

A functioning innate immune system may depend on the conservation of signalling pathways, more so than the conservation of particular immune genes. For example, in *Hydra* the TLR pathway is conserved through a scaffold of TIR-only and LRR-only proteins despite the lack of a full length TLR gene [18–20]. In our study, however, we found canonical TLR genes in most species (Table 1), with the exception of *A. pallida* and *Telmatactis* sp. In fact, no canonical TLR gene was identified in the *Aiptasia* genome [27]. Whether *A. pallida* and *Telmatactis* sp. also use scaffold TIR-only and LRR-only proteins remains unexamined, however, the expansion of TIR-only and TIR-domain-containing proteins in all species in this study may contribute to the building of scaffold-like immune pathways. Interestingly, the presence of 4 full length canonical TLR genes in coral (i.e., *A. digitifera*) [25] suggests that the canonical TLR was present in the common ancestor of anthozoans.

This principle of pathway conservation is demonstrated by the evolution of the interleukin-1 receptor gene family. The Ig-TIR domain combination (found in interleukin receptors such as IL-1R) is hypothesised to have evolved independently in both cnidarians and vertebrates [11, 12]. No IL-1R-like identified here were annotated as bona fide IL-1R protein by Swiss-Prot, rather they were all annotated as either TIR-domain-containing proteins or Ig containing genes. Further domain analysis demonstrated that these genes do indeed possess the canonical IL-1R-like architecture, with a C-terminal TIR, transmembrane domain and multiple Ig domains (Fig. 1). Previous studies [5, 11] have reported the independent evolution of the Ig-TIR domain combination and thus concluded that cnidarian IL-1R (or rather IL-1R-like) are not orthologs of the vertebrate IL-1R family. In particular, one study [12] argued that either the LRR-TIR (as seen in TLRs) or the Ig-TIR domain combination must have evolved independently in more than one metazoan lineage. Further phylogenetic analyses may clarify whether these gene families or domain combinations have evolved independently in multiple metazoan lineages.

Expansions in genes encoding TIR-domain-containing proteins, including both novel and ‘conserved’ (i.e., IL-1R-like) genes, may compensate for the lack of more than one TLR (or no TLR, in the case of *Aiptasia*, *Hydra* and *Telmatactis* [19, 27]). The TLR and IL-1R-like pathways both interact with MyD88 via their TIR domain during signal transduction, and therefore, may also interact with novel TIR-domain-containing proteins in a similar way. It is plausible to hypothesise that Novel Gene 3 (Fig. 2c) may interact in these pathways via its C-terminal death and TIR domains, which together form the canonical MyD88 domain architecture. Other novel TIR-domain-containing proteins (Fig. 2) may also interact in known or new pathways via the interactions of the TIR domain. Functional validation is required to determine the extent to which these novel genes contribute to cellular processes underpinning the innate immune response.

**Distribution of CniFL: a putative complement pathway receptor**

The initiation of known immune pathways though novel or lineage-specific immune genes is demonstrated by the discovery of the novel putative immune receptor CniFL. First reported in the recently sequenced *Aiptasia* genome [27], CniFL is hypothesised to function through the lectin-complement pathway. Here we identify a copy of the CniFL gene (Table 2) in all species (except *A. elegantissima*, however, this transcriptome was derived from a single tissue source (acrorhagi)), as well as multiple components of the complement pathway (Table 2). We
also identify three species (N. vectensis, A. tenebrosa and C. polypus) with CnFLs that contain an additional WAP domain and also identify a transmembrane domain CnFL in most species (Fig. 4). Interestingly, the single CnFL gene copy in N. vectensis was not identified previously [27] due to an incorrectly annotated ab initio gene model prediction in the Nematostella genome (v1.0). In addition, CnFL appears to have extensive copy number variation within hexacorallians (1 in N. vectensis to 7 in A. tenebrosa), but whether this variation in copy number is adaptive remains unexplored. The CnFL gene was hypothesised to have a role in symbiont uptake [27], which was partly due to its absence in the non-symbiotic N. vectensis. However, we observe that the CnFL gene is present in both symbiotic and non-symbiotic (A. tenebrosa, A. buddemeieri, C. polypus, N. vectensis) species investigated in our study (see Additional file 1: Table S2) with highest copy number reported in a non-symbiotic species (A. tenebrosa), therefore we recommend further functional validation to support this hypothesis.

Conclusion
In conclusion, actiniarians have a diverse repertoire of innate immune genes. These include novel domain architectures and potentially novel innovations, such as membrane-bound NLRs and novel putative complement pathway PRRs (i.e., CnFL). Further novelties may have been introduced to the actiniarian immune gene set through HGT followed by domain shuffling. Together, the conservation, expansion and diversification of different gene families and genes encoding TIR-domain-containing proteins have shaped the evolution of the actiniarian innate immune system. The conservation of specific protein domain architectures, immune genes and pathways provides an insight into the core gene set required for a functioning eumetazoan innate immune system. Further functional validation of these genes may elucidate conserved and novel biological functions, including how these genes may contribute to lineage-specific processes and morphological novelties.

Methods
To perform a comparative analysis of the innate immune gene repertoire, across multiple actiniarian taxa, we interrogated a suite of genomic resources, both publically available and newly generated in this study. Specifically, genomic datasets used in this study consisted of newly sequenced transcriptomes for Actinia tenebrosa (total n = 5; red colourmorph n = 2, brown n = 1, green n = 1, blue n = 1), Anthopleura buddemeieri (n = 1), Aulactinia veratra (n = 2), Calliactis polypus (n = 2), Nemanthus annamensis (n = 1) and Telmatactis sp. (n = 1), and as well as publically available transcriptomes for Anthopleura elegantissima (n = 1; SRX754678), Aiptasia pallida (n = 1; SRX231866, run SRR696721) and Nematostella vectensis (n = 1; SRX315372). These species were selected based on their phylogenetic distribution across most superfamilies within Actiniaria, availability of animals and general lack of sequence information for these species. Multiple transcriptomes were generated for A. tenebrosa and A. veratra which represent colourmorphs (red, brown, green and blue for A. tenebrosa) and aerial exposure treatments (one control and one treatment each for A. tenebrosa (red only) and A. veratra). All raw sequence reads were submitted to NCBI Sequence Read Archive under BioProject accession PRJNA313244 (see Additional file 1: Table S1 for all accession numbers).

Animal acquisition
Individual specimens of A. tenebrosa, A. buddemeieri and A. veratra were collected from the intertidal zone at Point Cartwright, (QLD, Australia) in 2014. C. polypus individuals were collected in 2014 from pumice washed onto rock pools at Snapper Rocks (QLD, Australia). Telmatactis sp. was bought from Cairns Marine (QLD, Australia) in 2014. N. annamensis was bought from Great Barrier Reef Marine Pty Ltd in 2015. All animals were housed in holding tanks, in the marine laboratory at QUT, until experimental use. Tanks were maintained at between 33 and 37 ppt salinity and 20–28 °C.

Sequencing, assembly and annotation
Total RNA was extracted from whole organisms by first homogenizing individuals in liquid nitrogen, followed by a TRIzol/chloroform RNA extraction protocol (TRIzol®, Life Technologies). Extracted RNA was assessed for quality and integrity on a Bioanalyzer 2100 (Agilent) using an RNA nano chip. Sequencing libraries were prepared using the Illumina TruSeq® Stranded mRNA Library Preparation Kit and sequenced on an Illumina NextSeq 500. 150 bp paired-end chemistry was used for all sample libraries, except N. annamensis and A. tenebrosa (4; blue) which were prepared at a different time using 75 bp paired-end chemistry. All transcriptomes were assembled using the Trinity de novo assembler [39]. Trinity assembler was used for A. tenebrosa (1, 2 and 4), A. pallida, A. buddemeieri, A. elegantissima, A. veratra, C. polypus (1, 2), N. annamensis, N. vectensis and Telmatactis sp.. Quality trimming was performed on all datasets using Trimmomatic [40] with default settings to retain only high quality reads and to remove non-biological sequence.

Redundant and chimeric sequences were removed from all assemblies using CD-Hit v.4.6.1 [41, 42], by clustering sequences with greater than 95 % similarity within each assembly. In addition, CEGMA (Core Eukaryotic Genes Mapping Approach) v.2.5 [43] was used to assess the completeness of each assembly, by determining the percentage
of full-length sequences in each transcriptome corresponding to 248 highly conserved eukaryotic proteins.

Functional annotation of individual transcriptomes was undertaken using Trinotate [44], for both nucleotide and predicted peptide sequences. Nucleotide sequences for each contig were annotated using BLASTx searches against the Swiss-Prot [45] and TrEMBL (Uniref90) [46] databases using BLAST+ v.2.2.31 [47, 48] software (E value 1 × 10^{-5}). Batch extraction and translation of the longest open reading frames (ORFs) for each contig were performed using TransDecoder v.2.0.1 [49, 50]. From the predicted peptide sequences, signal peptides and protein families were detected in each ORF, using Signal Peptide v.4.1 [51] and Pfam v.3.1b1 [32], respectively. Longest ORFs were annotated using BLASTp searches against the Swiss-Prot [45] and TrEMBL [46] databases using BLAST + v.2.2.31 [47, 48] software (E value 1 × 10^{-5}).

Gene ontology and RSEM analysis
GO terms were assigned to contigs that received significant BLASTx hits (E value 1 × 10^{-5}). WEGO [52] and CateGoRizer [53] were used to examine the distribution of top GO terms and the distribution of ‘immune class’ GO terms in each assembly, respectively. Estimates of transcript abundance for all newly generated transcriptomes were calculated using the Trinity [50] pipeline. Raw reads were aligned to assembled contigs using Bowtie 2 [54], followed by RNAseq by Expectation-Maximization (RSEM) [55] estimation of expression values as Fragments per Kilobase of transcript per Million mapped reads (FPKM).

Identification of candidate genes
The list of candidate genes interrogated in this project was limited to five innate immune gene families which included Toll-like receptor (TLR), Nucleotide-binding and Leucine-rich Repeat containing gene (NLR), Interleukin-1 receptor-like genes (IL-1R-like), Myeloid Differentiation primary response gene 88 (MyD88) and Nuclear Factor kappa-light-chain-enhancer of activated B cells (NF-κB). Transcripts containing full length ORFs of these candidate genes were identified using Pfam, BLASTp and BLASTx searches. Specifically, the search method for each candidate gene was as follows. TLR was identified by searching for all contigs with a TIR (PF01582) (or TIR_2, PF13676) domain, along with at least one leucine-rich repeat (LRR, CL0022). MyD88 was identified by searching for a TIR (or TIR_2) domain, along with a death domain (DD). IL-1R-like was identified by searching for a TIR (or TIR_2) domain, along with at least one immunoglobulin (Ig, CL0011) domain. NF-κB was identified by searching for a Rel homology domain (RHD, PF00554), along with at least one ankyrin repeat (PF00023). The presence of other domains with a RHD may be indicative of different classes of NF-κB, however, in these actiniarian species no other domains were found in the same contig with a RHD. Hence, all identified NF-κB belong to Class 1. A Rel homology domain without ankyrin repeats, as seen in N. vectensis [56] were not considered as full length ORFs. NLR was identified by searching for a NACHT (domain found in NAIP, C2TA, HET-E and TIP1) domain, along with at least one LRR. NLR genes may also include a variable N-terminal domain, which can be broadly classified in the ‘death-fold superfamily’ which includes the domains caspase activation and recruitment domain (CARD, PF00619), pyrin domain (PYD, PF02758), death domain (DD, PF00531) and death effector domain (DED, PF01335). Therefore, contigs also containing a domain from this superfamily were considered potential candidate NLRs. In addition, the translated ORF for each candidate gene was analysed by TMHMM v.2 [57], to identify transmembrane domains.

Finally, contigs were only retained as complete candidate genes if they consisted of a full length ORF which contained a start and stop codon, and received a BLASTp hit (from either the TrEMBL or Swiss-Prot databases) against other species using the open reading frame batch extracted by TransDecoder [49].

Presence/absence of other innate immune genes
Non-conservative counts of other innate immune genes in this study were identified from gene counts provided by Trinotate. CniFL was identified by searching for all genes containing the Pfam domains Collagen (PF01391), Ig (CL0011) and Fibrinogen (PF00147). MASP was identified from BLAST hits from TrEMBL with N. vectensis MASP and presence of the correct Pfam domains (CUB, EGF-like calcium binding, CUB2, Sushi 1, Sushi 2 and Peptidase S1). SRCR were identified by searching for the number of genes with Pfam annotations for the SRCR domain (PF00530). C-type lectin domains were identified by searching for the number of genes with Pfam annotations for lectin_C domains (PF0059). Complement control proteins (C3, Factor B, C6 and Factor I) were identified by searching for BLAST hits from either the TrEMBL or Swiss-Prot databases.

Identification of novel immune genes
Potential novel genes were identified by searching for novel domain architectures that included the TIR (PF01582) and/or TIR_2 (PF13676) domains. This domain is well documented in having a role in signal transduction and protein-protein interactions in immune pathways, and therefore, provides a robust target for finding putative novel genes that interact in innate immune pathways [58]. All species were also interrogated for novel actiniarian genes previously identified [11, 27]. Initially, all TIR or TIR_2 domain containing contigs, for each species, were identified in the Trinotate annotation
report, and all contigs without a BLASTx or BLASTp annotation were further investigated. In addition, contigs that received BLASTx or BLASTp hits only against N. vectensis predicted proteins or genes were also interrogated in detail. The EMBl-EBI [32] Pfam domain architecture analysis (http://pfam.xfam.org/ Accessed 28th March 2016) was used to investigate known architectures with either the TIR or TIR_2 domains, and thereby identify which species, if any, have genes with novel or infrequently used architectures. Novel architectures were identified if no or few hits were returned from either BLAST or EMBl-EBI, which may indicate potential actiniarian lineage-specific (orphan) genes, horizontally transferred genes or taxonomically-restricted genes.

Candidate and novel gene validation
To validate the transcriptome assemblies, as well as the ORF of candidate transcripts, PCR validation was performed for the five candidate genes, as well as three novel genes and CniFL. All primers were designed using Primer-BLAST [59] using the settings of [60], to amplify the ORF of each respective gene for four species: A. tenebrosa, A. buddemeieri, A. veratra and C. polypus (except CniFL, for which genes were validated in A. tenebrosa and A. buddemeieri only). For large ORFs, multiple primer sets were designed to ‘walk’ the length of the ORF, with at least 100 bp overlap between products. A detailed list of primer sequences is presented in Additional file 3: Table S1. PCRs were performed using RNA extracted using the above described protocol, followed by cDNA synthesis using the SensiFAST™ cDNA synthesis kit (Bioline). PCR products were then amplified using MyFi™ DNA polymerase mix (Bioline) (PCR protocols and thermocycler conditions can be found in Additional file 3: Tables S12–S13). Purification was performed using the ISOLATE II PCR and Gel kit (Bioline), and sequenced using BigDye® Terminator v3.1 (Thermo-Fisher). Sequences were run on a Genetic Analyser 3500 from Applied Biosystems (ThermoFisher). Sequence chromatograms were visualised in Geneious version R8.1.4 [61] and aligned to the transcript from which the primers were designed, as per [62]. To confirm the ORF for the candidate gene, sequence similarity was compared.

Evolutionary and phylogenetic analyses
PAML
To investigate whether genes were under purifying, neutral or positive selection and thereby provide insights into the selective pressures on the actiniarian immune system, PAML analysis was performed. Non-synonymous vs synonymous substitution rates (dN/dS) of all TLR, MyD88, NF-κB and the transmembrane domain containing NLRs, as well as Novel Genes 1, 2 and 3, were analysed using the method of Yang and Nielsen [33]. First, the nucleotide sequence for the complete ORF of all genes were extracted from the TransDecoder output and imported into Geneious [61] version R8.1.4. All stop codons were manually removed and then sequences were imported into MEGA version 7.0.14 [63] for alignment with MUSCLE [64] (codon), using default settings and gaps removed. Alignments were converted into PHYLIP format in Geneious [61] and then imported into pamIX [65, 66] version 1.3.1, where dN/dS ratios were calculated using YN00 [33]. All values for dN and dS were averaged for each gene to provide an average dN/dS.

RAxML
In order to understand how actiniarian NLRs have evolved in the context of the metazoan NLR family, the phylogenetic relationship of NLRs in this study and other metazoan NLRs were resolved using maximum likelihood analysis in RAxML. Of particular interest was the question of how the novel TMD containing NLRs have evolved (i.e., from a single origin or multiple independent origins). Initially, the translated ORFs for 35 complete and near-complete NLRs (near-complete only used for species where no complete NLR found) of the candidate NLR transcripts were imported into Geneious [61] version R8.1.4 and aligned along with NACHT domains from species across Metazoa (Additional file 6) which were obtained from [7]. Alignments were performed using MUSCLE [64] with default settings (duplicates removed). Sequences were trimmed to retain only the NACHT domain, as this represents the most evolutionarily conserved domain of NLRs, across metazoan species and is useful for inferring phylogenetic relationships [7]. Alignments were imported into MEGA version 7 [63] to determine the best model of protein evolution. Maximum likelihood analysis was then performed in RAxML version 8.1.15 [67], using rapid bootstrapping (1000 replicates) with a random seed and no outgroup.

To understand how CniFL genes are disturbed across Anthozoa species, maximum likelihood analysis of CniFL proteins was performed. All but two species had at least one gene copy for which a complete ORF was identified; Telmatactis sp. had only partial sequences and no CniFL was identified in A. elegantissima. An initial alignment of the amino acid sequences of the complete ORFs was performed using MUSCLE [64]. Alignments were trimmed using Gblocks [68] to remove poorly aligned sections, allowing for gap positions within final blocks and not allowing many contiguous non-conserved positions. Protein model testing in MEGA [63] and maximum likelihood analysis in RAxML [67] was performed the same as above for NLR. The outgroups (Hs FCN1 and Mus FCN1) were set post-analysis in Geneious [61]. SMART analysis [69, 70] was used to identify the number of domains in each CniFL shown in the tree.
Acknowledgements

The authors would like to thank the Evolutionary and Physiological Genomics Lab (ePGL), in particular Hayden Smith and Matthilde Klein, for their continual help and support. The authors would also like to thank QUT Marine group for their advice caring for the animals. The authors would like to acknowledge QUT Marine Group, Queensland University of Technology, Brisbane, Australia.

Funding

This project was funded by research support provided to A.P. by the School of Biomedical Sciences, Faculty of Health, Queensland University of Technology, Brisbane, Australia.

Availability of data and materials

Raw sequence read data sets and validated sequences have been deposited at NCBI Sequence Read Archive (SRA) (BioProject PRJNA313244) and GenBank® (https://www.ncbi.nlm.nih.gov/nuccore/) respectively. Accession numbers can be found in Additional files 1 and 3.

Authors’ contributions

CVDB, AP, JS and PP conceived and designed the project. CVDB and JS assembled and annotated the transcriptomes. CVDB analysed the data. CVDB, AP, JS and PP wrote and edited the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

Not applicable.

Ethics approval and consent to participate

This project did not require animal ethics approval, however, sample collection was authorised under the Fisheries Act 1994 (General Fisheries Permit), permit number: 166312.

Author details

1School of Biomedical Sciences, Faculty of Health, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia. 2Institute of Health and Biomedical Innovation, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia. 3School of Earth, Environmental and Biological Sciences, Science and Engineering Faculty, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia. 4Institute of Future Environments, Queensland University of Technology, GPO Box 2434, Brisbane, Qld 4000, Australia.

Received: 28 May 2016 Accepted: 25 October 2016

References

1. Akera S, Uematsu S, Takeuchi O. Pathogen recognition and innate immunity. Cell. 2006;124:783–801.
2. Parkin J, Cohen B. An overview of the immune system. Lancet. 2001;357:1777–89.
3. Charles A, Janeway J, Medzhitov R. Innate immune recognition. Annu Rev Immunol. 2002;20:197–216.
4. Fijnheer M, Kasahara M. Origin and evolution of the adaptive immune system: genetic events and selective pressures. Nat Rev Genet. 2010;11:47–59.
5. Miller DJ, Hemmrich G, Ball EE, Hayward DC, Khatthiru K, Funayama N, et al. The innate immune repertoire in Crinata – ancestral complexity and stochastic gene loss. Genome Biol. 2007;8:R59.
6. Putnam NH, Srivastava M, Hellsten U, Dirks B, Chapman J, Salamov A, et al. Sea anemone genome reveals ancestral eumetazoan gene repertoire and genomic organization. Science. 2007;317:86–94.
7. Yuan B, Bayes JM, Denman SM. The characterization of sponge NLRs provides insight into the origin and evolution of this innate immune gene family in animals. Mol Biol Evol. 2014;31:106–20.
8. Gauthier MEA, Du Pasquier L, Denman BM. The genome of the sponge Amphimedon queenslandica provides new perspectives into the origin of Toll-like and interleukin 1 receptor pathways. Dev Biol. 2010;295:139–53.
9. Technau U, Radd S, Maxwell P, Gordon PM, Sainia M, Grasso LC, et al. Maintenance of ancestral complexity and non-metazoan genes in two basal cnidarians. Trends Genet. 2005;21:633–9.
10. Buckley KM, Rast JP. Diversity of animal immune receptors and the origins of recognition complexity in the deuterostomes. Dev Comp Immunol. 2015;49:151–9.
11. Poole AZ, Weis VM. TLR-domain-containing protein repertoire of nine anthozoan species reveals coral–specific expansions and uncharacterized proteins. Dev Comp Immunol. 2014;46:480–8.
12. Zhang Q, Zmasek CM, Godzik A. Domain architecture evolution of pattern-recognition receptors. Immunogenetics. 2010;62:263–72.
13. Zhang L, Li L, Guo X, Litman GW, Dishaw LJ, Zhang G. Massive expansion and functional divergence of innate immune genes in a protostome. Sci Rep. 2015;5:8693. Available from: http://www.nature.com/articles/rep408693/full/rep408693.html.
14. Ting J-P, Verrier A, Arnett TJ, Basset F, Bohus B, et al. The NLR gene family: a standard nomenclature. Immunity. 2008;28:285–7.
15. Hamada M, Shoguchi E, Shinzato C, Kawashima T, Miller DJ, Satoh N. The complex NOD-Like receptor repertoire of the coral Acropora digitifera includes novel domain combinations. Mol Biol Evol. 2013;30:167–76.
16. Lange C, Hemmrich G, Rostembler UC, Lopez-Quintero AJ, Miller DJ, Rahn T, et al. Defining the origins of the NOD-Like receptor system at the base of animal evolution. Mol Biol Evol. 2011;28:1587–702.
17. Wu B, Huang T, Gong J, Zhou P, Bai Z. Domain combination of the vertebrate-like TLR gene family: implications for their origin and evolution. J Genet. 2011;90:401–8.
18. Franzenburg S, Fraune S, Künzel S, Baines JF, Domazet-Loso T, Bosch TC. MyD88-deficient Hydra reveal an ancient function of TLR signaling in sensing bacterial colonizers. Proc Natl Acad Sci U S A. 2012;109:19374–9.
19. Bosch TC, Augustin R, Anton-Enleben F, Fraune S, Hemmrich G, Zill H, et al. Uncovering the evolutionary history of innate immunity in the simple metazoan Hydra uses epithelial cells for host defence. Dev Comp Immunol. 2009;33:559–69.
20. Augustin R, Fraune S, Bosch TC. How Hydra senses and destroys microbes. Semin Immunol. 2010;22:54–8.
21. Augustin R, Bosch TC. CNIDARIAN IMMUNITY: A Tale of Two Barriers. In: Soderhall K, editor. Invertebrate Immunity. Berlin: Springer-Verlag Berlin; 2010. p. 1–16.
22. Mantindale MQ. The evolution of metazoan axial properties. Nat Rev Genet. 2005;6:917–27.
23. Bosch TC. Cnidarian-microbe interactions and the origin of innate immunity in metazoans. Annu Rev Microbiol. 2013;67:499–518.
24. Davy SK, Allemand D, Weis VM. Cell biology of Cnidarian-dinoflagellate symbiosis. Microbiol Mol Biol Rev. 2012;76:229–61.
25. Shinzato C, Shoguchi E, Kawashima T, Hamada M, Hisata K, Tanaka M, et al. Using the Acropora digitifera genome to understand coral responses to environmental change. Nature. 2011;476:320–3.
