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Innate immunity driven by pattern recognition receptor (PRR) protects the host from invading pathogens. Aquatic animals like fish where the adaptive immunity is poorly developed majorly rely on their innate immunity modulated by PRRs like toll-like receptors (TLR) and NOD-like receptors (NLR). However, current development to improve the fish immunity via TLR/NLR signaling is affected by a poor understanding of its mechanistic and structural features. This review discusses the structure of fish TLRs/NLRs and its interaction with pathogen associated molecular patterns (PAMPs) and downstream signaling molecules. Over the past one decade, significant progress has been done in studying the structure of TLRs/NLRs in higher eukaryotes; however, structural studies on fish innate immune receptors are undermined. Several novel TLR genes are identified in fish that are absent in higher eukaryotes, but the function is still poorly understood. Unlike the fundamental progress achieved in developing antagonist/agonist to modulate human innate immunity, analogous studies in fish are nearly lacking due to structural inadequacy. This underlies the importance of exploring the structural and mechanistic details of fish TLRs/NLRs at an atomic and molecular level. This review outlined the mechanistic and structural basis of fish TLR and NLR activation.

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The mechanism of action of innate immunity involves a family of proteins characterized by a highly specialized structure often termed as pattern recognition receptors (PRRs) [13]. PRRs play an essential role in innate immunity by recognizing different conserved microbial motifs viz., carbohydrates (lipopolysaccharide, mannose, fructose, sucrose etc.) [14], nucleic acids (DNA/RNA) [15], peptides (flagellin) [16], peptidoglycans (PGN), lipoteichoic acids (LTA), lipopolysaccharides (LPS), muramyl dipeptide (MDP), γ-D-glutamyl-meso-diaminopimelic acid (iE-DAP), N-formylmethionine, lipoproteins and glucans or endogenous substances collectively known as microbial/pathogen/danger-associated molecular patterns (MAMPs/PAMPs/DAMPs) [2,17]. PRRs are outspread in extracellular, membrane and cytoplasmic compartments and are classified according to their ligand specificity, function and cellular localization. Functionally, PRRs are divided into two categories: endocytic PRRs and signaling PRRs. Signaling PRRs include large families of membrane-bound toll-like receptors (TLRs) [18–20] and cytoplasmic NOD-like receptors (NLRs) [21,22]; and other receptors viz., retinoic acid-inducible gene I-like receptors (RLRs) [23,24], C-type lectin receptors (CLRs), AIM2-like receptor (ALRs) and OAS-like receptors (OLRs) [25,26]. PRRs following the detection of PAMPs trigger the signaling cascade and activate the innate immune response via stimulation of an array of downstream signaling molecules that include chemokines, cytokines, antimicrobial peptides and interferons (Fig. 1) [27,28].

Aquatic animals especially fish; a chief resource for food and a source of micronutrients and essential acids has shown to be greatly affected by microbial diseases. Fish is one of the most commercially important agricultural product that directly influence the economy [29]. Thus, the protection of fish production by controlling the disease is a major area in aquaculture research. Fish that are exposed to bacteria, fungi and viruses in an aquatic medium largely protect themselves through their innate immunity. The innate immunity in fish is majorly contributed by the signaling PRRs i.e. TLRs and NLRs. TLRs were the first class of PRRs [30] that are extensively studied and characterized in fish and several other vertebrates [31]. Next to TLRs, NLRs are the second class of PRRs that are subjected to considerable investigation [32]. A key difference between TLRs and NLRs are the cellular localization and PAMPs selectivity. TLRs are a class of extracellular transmembrane PRRs, whereas NLRs belong to the class of intracellular cytoplasmic receptors [26]. In TLRs, the periplasmic extracellular domain (ECD) senses the PAMP following signal transduction to the cytoplasmic components via a single transmembrane (TM) domain (Fig. 1) [33]. Structural characterization of TLRs showed a conserved tripartite domain architecture consisting of ECD, a single-pass TM and a cytoplasmic Toll/IL-1 receptor (TIR) domain. The ECD of TLRs are composed of a tandem of evolutionarily conserved leucine-rich repeats (LRRs) [34] that selectively recognizes PAMPs and varies in number from species-to-species. Like TLRs, NLRs also present a tripartite domain architecture characterized by a N-terminal effector domain, a central nucleotide binding domain (NBD) and a C-terminal LRR domain [35]. The function of the effector domain is similar to the cytoplasmic TIR domain of TLRs that interacts with downstream effector molecules following activation of signaling cascade involved in innate immune defense. The effector domain also classifies NLRs to subfamilies that include NLRa, NLRb, NLRc and NLRp [22,35]. The NBD domain (NACHT: NAIP, CIITA, HET-E and TP1) is responsible for ATPase activity and mediates oligomerization [36]. The LRR domain in NLR resemble to the topology of TLR-ECR domain and recognizes PAMPs, which crosses the plasma membrane or endogenous DAMPs (Fig. 1) [37]. Unlike TLRs, NLRs show a broad category of function that include signal transduction, autophagy, apoptosis regulation,
inflammasome assembly and transcription activation. Readers interested to know more about the TLR and NLR domain architecture, signaling molecular pathway and function in human and other higher eukaryotes are referred to published review articles [21,33,38–41]. This review specifically discusses the domain classification (ECD, TIR, CARD and NACHT), structure and function of fish TLR and NLR domains that include TLR2, TLR3, TLR22, NOD1 and NOD2 and downstream molecules like MyD88-TIR, TRIF and RIP2-CARD.

2. Domain architecture of fish TLRs and NLRs

In fish several TLRs have been identified that share a close homology with other eukaryotes including humans (reviewed in [42,43]). Fish that are exposed to an array of microbes with rudimentary adaptive immunity possess more TLRs as compared to humans. While in mammals only 13 TLRs (10 TLRs in human: TLR1–10; and 12 TLRs in mouse: TLR1–10, TLR11–13) [39] have been identified, in teleost, over 20 TLRs were discovered highlighting their importance in providing the first line of defense in fish [42]. Unlike higher vertebrates, in fish TLR6 and TLR10–13 are absent. But, in fish duplicate and multiple copies of TLRs (TLR3, TLR4, TLR5, TLR7, TLR8, TLR20 and TLR22) are identified that are shown to be involved in the fish development [43]. Fish TLRs are structurally homologous to human TLRs and share a sequence identity ~30–70% in the PAMP interacting LRR domains [44–46]. As shown in Fig. 2, fish TLRs possess a signal peptide (~<30 amino acids, aa) that gets cleaved from the mature protein. These mature TLRs composed of ~800–900 amino acids (TLR2, TLR3, TLR4, TLR5 and TLR22). The ECD consists of a variable number of LRR domains that share a substantial sequence identity with the homologous human or mouse TLRs. The number of LRR repeats varies across species and TLR class. In zebrafish TLR3, 24 LRR are identified, whereas Indian carp TLR3 (rohu) presents 27 LRR [44,45]. The amino acid composition of LRR motifs (xLxxLxxxNLxxLxxxFxxLx; where L = leucine/soleucine/valine/phenylalnine; x = any amino acid residue; N = asparagine/threonine/serine/cysteine, and F = phenylalanine) varies across the TLRs making them selective to target a PAMP [47]. Rohu TLR2 and zebrafish TLR22 are composed of 23 and 26 LRR domains, respectively, and each LRR has ~20 amino acids [44,46]. It should be noted that the position and number of LRR varies depending on the algorithm used for LRR motif prediction [48–51]. In general, the secondary structure of LRR domains is evolutionarily highly conserved, but varies in primary structure (Fig. 2). The TM domain is a single-pass helix composed of ~20 amino acids that connects the cytoplasmic TIR domain with the ECD. The TIR domain is comparatively more conserved in fish and mammals than the LRR domain [52]. In general, like TLR motifs, the TIR domains are highly conserved in different species, sharing an average sequence identity >70% for the functional secondary structure units α/β that mediates a homotypic TIR–TIR interactions [46].

Unlike the TLR domain organization, the cytoplasmic NOD subgroup in fish NLRs share a structure that lacks signal peptide and TM domain. Nevertheless, both TLR and NOD receptors in fish have a common three-domain organization [53]. Importantly, the effector domain of fish NOD receptors consist of one or two caspase-activation and recruitment domains (CARD) that distinguish them from other NLR family proteins containing pyrin or baculovirus inhibitor of apoptosis protein repeat domain [22]. The five major NOD receptors NOD1–5 have been identified in different fish species that include zebrafish, Indian carps, catfish, Japanese flounder, Atlantic salmon etc. and are homologous to mammals [54–63]. The NOD proteins in fish are comparatively larger than TLRs and composed of ~940 and ~980 amino acids (aa) in NOD1 and NOD2, respectively [54,58,60,64–66]. In zebrafish, the NOD1 and NOD2 proteins contain one (94 aa) and two CARD domains (96 and 90 aa) at the N-terminus that mediate homotypic CARD–CARD interactions with downstream signaling molecules (Fig. 2). The central domain in NOD proteins is rather complex and their functions are poorly understood. Sequence analysis of the central domain highlighted three conserved domains in fish that include NBD, two helical domains (HD) flanking a winged helix domain (WHD) [67,68]. These domains are connected through a variable length flexible linker allowing multiple functions such as dNTP binding, oligomerization and signal transduction. The NBD domain is divided into subdomains Walker A, Walker B and Sensor 1. The functional motif in Walker-A ‘G-D-E-A-G-S/V-G-K-S’ and Walker-B ‘L-T-F-D-G-L-Y-D’ subdomains that binds to ATP is highly conserved in fish, human and mouse NOD1/ NOD2. Similarly, the functional motifs ‘G-S/L-C/H-S/I/L-V-P-L-V-F’ and ‘/F/L-Y-E-F-L-H’ in HD and WHD subdomains, respectively, are conserved in NOD1 and NOD2 proteins highlighting a similar mode of function and dNTP binding. The LRR domain in fish NODs is rather smaller (~250 aa) as compared to TLRs (~700 aa). The difference in the length of LRR domain between TLRs and NLRs is due to their selective PAMP binding and homo- or hetero-oligomerization. Homo and heterodimers of TLR-EC domains are known to mediate their function and oligomerization is required to recognize long-sized PAMPs like DNA and RNA of viruses [69–73]. However, the comparatively short LRR domain having less number of LRR motifs present in fish NOD receptor is not involved in homo- or hetero-oligomerization.

3. PAMP/MAMP/DAMP specificity of fish TLRs and NLRs

Water is a great source of pathogenic microorganisms and provides a suitable environment for the growth of major aquatic microbes such as bacteria, fungi, virus, algae, and protozoa. While these microorganisms are a part of the food chain, several of them are known to cause severe diseases in fish. As an example, bacteria species such as Pseudomonas [74], Vibrios [75], and Aeromonas [76]; viruses like viral hemorrhagic septicemia virus (VHSV), grass-carp reovirus (GCRV), betanodavirus are deadly and causes severe disease to both seawater and freshwater fish (reviewed in [77,78]). This not only directly affects the faster growing aquaculture industry, but also poses threat to human health due to a poor understanding of the evolution of novel aquatic viruses and their diversity. Such possible threats to human life can be realized from the recent COVID-19 pandemic that threatens millions of human life worldwide with an unknown origin.

In fish, TLRs are the most studied PRRs and their PAMP selectivity is comparatively best characterized. Common bacterial components recognized by fish TLR include PGN, LTA, LPS and flagellin (Fig. 1) [79]. TLR2 expression in fish has shown to be up-regulated in response to PGN and LTA [80]. TLR4 and TLR5 respectively have been identified to recognize LPS and flagellin in higher eukaryotes, but some fishes lacks TLR4 and do not recognize LPS and have multiple copies of TLRs indicating their PAMP selectivity [81–83]. Viral and bacterial RNA is shown to be a molecular target for TLR3, TLR9, TLR13 and TLR22 in fish [31,84–88]. PAMPs for a major class of fish TLRs such as TLR1, TLR7, TLR8, TLR13–20 and TLR23–27 remain unexplored. In addition to natural bacterial/viral components, synthetic ligands for fish TLRs have been tested that include poly(I:C), CpG DNA, CpG ODN and triacylated lipopeptide. Readers are referred to check previous review articles [42,43] to compare PAMP specificity between fish and mammal TLRs.

Like TLRs, fish NLRs recognize an array of ligands derived from microorganisms. NLRs trigger the activation of downstream molecules by sensing either PAMPs or DAMPs. Bacterial components such as LPS, PGN and its degraded products MDP and iD-PE are shown to activate NOD1 and/or NOD2 expression (Fig. 1) [59,60,89]. Major bacterial/fungal components and synthetic ligands like LTA, poly(I:C), mannan, toxins, glucan or β-1,3-glucan tested to have no effect on NOD2 (zebrafish), but are potential ligands for other class of fish NLRs (for example poly(I:C) activate NLR-C3 and NOD1) [61,64]. Bacterial flagellin is also shown to activate NLR-C in common carp [90]. The effect of intracellular activators such as cholesterol, dNTPs, metabolites, and antimicrobial peptides, on NLRs are less explored in fish. Nevertheless, to date studies of the effect of other aquatic parasites such as protozoans, fungi, algae and their molecular components modulating TLR/NLR activation
in fish remain mostly undiscovered. Furthermore, the reported stipulated number of ligands that are majorly identified in mammals and tested on fish limit our present understanding of the major classes of PAMPs that activate fish immunity via TLRs/NLRs recognition. To this end, lack of structural and mechanistic insights into the ligand recognition and specificity by fish TLRs/NLRs and innate immune activation further limits the development of novel synthetic molecules to boost fish innate immunity. While deriving such structural and mechanistic information is challenging, recently some studies shed light on the mechanism of PAMP interaction and downstream molecule interaction with TLR/NLR receptors using in silico structural biology approaches and are summarized in the following sections.

4. Comparative tertiary structure modeling of fish TLRs/NLRs

Establishment of a crosstalk between structure-to-function has been a bottleneck in the field of structural biology and drug-discovery research. Unfortunately, even after nearly a few decades since TLRs/NLRs are identified in zebrafish [91,92], experimental structure for these proteins is not available for most of the proteins except zebrafish TLR5 (ECD) [93] to understand their mechanism of action. The complex topology and molecular pathways are among the major roadblocks for the structural investigation. Nevertheless, the unavoidable limitations associated with experimental techniques add more challenges to solve these biological complex questions. For example, association of TLRs with membrane limits the application of X-ray crystallography technique to solve full-length TLR structure; the high-molecular size of TLRs/NLRs ~100 kDa (excluding homo-/hetero-oligomers) make it difficult to study using nuclear magnetic resonance (NMR). However, recent developments in cryo-EM that prefer large-molecular size protein and can be studied in a membrane environment look promising to study full-length TLR/NLR structure. In addition, advance membrane mimetic tools like nanodiscs and bicelles to study membrane proteins provide an promising alternative to study functional TLR domains by NMR [94,95]. To this end, computational techniques are an alternative and attractive approach that bridge the structure-to-function executing a coordinated sequence of structural bioinformatics tools. In this coordinated sequence, the first tool is building a tertiary model structure for fish TLRs/NLRs using a comparative/ab-initio molecular modeling approach as shown in Fig. 3.

In silico molecular modeling approach builds a 3D model structure with accuracy using the amino acid sequence of the target protein (query sequence) [96,97]. Using a homology based approach, the query protein sequence is scanned over experimentally solved structures (template) deposited in the public protein data bank (PDB) [98] database that presently contain >160,000 structures. Upon satisfying a threshold similarity/identity between the query sequence and template structure(s) using computational algorithms, 3D coordinates can be generated for the target protein using multiple steps as illustrated in Fig. 3 [reviewed in [97]]. The quality of the initial 3D model structure mostly depends on the degree of sequence identity/similarity between the query and template protein. Comparative model building using templates available in the current PDB database shows a good reliability for fish TLRs and NLRs as summarized in Table 1. Zebrafish as a model organism to study fish TLR/NLR is considered for searching for available
templates. As listed in Table 1, most of the zebrafish TLRs show a sequence identity (≥30%) except for TLR18–20. It should be noted that the identity listed in Table 1 is obtained from BLASTp search against the current PDB database for the full-length zebrafish TLR/NLR proteins. Special attention for obtaining a maximum query coverage (minimum gap in the query-template alignment) and sequence identity need to be considered for the modeling of functional domains. TLR-ECD such as rohu TLR3-ECD, zebrafish TLR3-ECD and TLR22-ECD have shown a query-template sequence identity over 25–30% which is considered to be reliable for homology modeling (Fig. 3) [97]. For instance, rohu TLR3-ECD shows an identity and similarity of 48 and 65%, respectively, with mouse TLR3-ECD; and rohu TLR3-TIR domain shares 33 and 56% sequence identity and similarity, respectively, with human TLR3-TIR [45]. Similarly, zebrafish TLR3 (identity/similarity: 48/64%)/TLR22-ECD (identity/similarity: 28 and 43%) share a reasonable homology with human TLR3-ECD for comparative structure modeling [44]. Rohu TLR2-ECD (identity/similarity: 35/52%), TLR2-TIR (identity 71%) and common carp

![Fig. 3. A schematic showing an in silico strategy for the structural and functional investigation of fish TLR and NLR proteins. Step (1–5) shows 3D structure modeling and refinement strategy for fish TLR/NLR proteins. Step (6–10) shows strategy to build TLR/NLR-PAMP or protein-protein complex following structural and functional analysis using long-range MD simulations.](image-url)

Table 1

| Zebrafish Uniprot ID | Template (PDB ID) | Description | Organism | Identity (%) |
|---------------------|-------------------|-------------|----------|--------------|
| TLR1 B3DIW3         | 1FYV              | TLR1       | Human    | 57.52        |
| TLR2 F1RIU3         | 5D3I              | TLR2       | Mouse    | 34.03        |
| TLR3 B8JL3          | 3CIG              | TLR3       | Mouse    | 47.88        |
| TLR4a B3U3W0        | 3FXI              | TLR4       | Human    | 35.21        |
| TLR5b B3O1N1        | 3V44/3V47/5GY2    | TLR5b      | zebrafish| n/a          |
| TLR6c Absent        | n/a               | n/a        | n/a      | n/a          |
| TLR7 F1QY64         | 5GMF              | TLR7       | Rhesus macaque | 56.79 |
| TLR8a F1P2P4        | 3W3G              | TLR8       | Human    | 41.66        |
| TLR9 F1QY61         | 3WPB              | TLR9       | Horse    | 35.52        |
| TLR10–13c Absent    | n/a               | n/a        | n/a      | n/a          |
| TLR16d B3O9K5       | 3A79/3WPB         | TLR6/TLR9  | Mouse/Horse | -23   |
| TLR19d A0A2R9Q6Z1   | 3JOA              | TLR5       | Human    | -23          |
| TLR20d F1QRG0       | 5ZSA              | TLR7       | Rhesus macaque | -23   |
| TLR21 F1QMN8        | 420C              | TLR13      | Mouse    | 30.52        |
| TLR22 B3O3JI6       | 3JOA              | TLR5       | Human    | 29.36        |
| NOD1 X1WQ4         | 5IRM/5IRL        | NOD2       | Rabbit   | 32.49        |
| NOD2a F8W3K2         | 5IRM/5IRL       | NOD2       | Rabbit   | 50.67        |

* Represents TLR/NLR present in zebrafish with multiple copies (templates are shown for only one copy).

* Crystal structure is available.

* Represents a few TLRs absent in zebrafish, but are present in mouse/human.

* TLRs with very low sequence identity (templates shown are based on the highest query coverage; whereas other TLR/NLR templates listed in the table are considered based on the maximum BLASTp score).
downstream molecule MyD88-TIR (identity 78%) also share a very good homology with mouse TLR and human MyD88 crystal structures (template) [46].

As compared to fish TLRs, NOD domains share a good sequence homology with the rabbit NOD2 crystal structure (Table 1) [99]. Previously, advanced techniques like ab-initio modeling [100] and protein threading [101] that employ a multi-template modeling approach was used to construct reliable 3D-model structures for rohu NOD1 and NOD2 LRR domains (Fig. 3) due to the unavailability of an experimental structure for NOD receptor [102,103]. However, at present the BLASTp search of rohu NOD1 LRR against PDB data bank resulted four hits that include the PDB IDs: 5IRM (34.66% identity), 5IRL (34.66% identity), 4R5D (29.63% identity) and 4R6G (28.64% identity). This suggests that, the better query-template identity between fish and rabbit NOD proteins [99] could provide an improved 3D model structure for rohu/zebrafish NOD proteins. Similarly, models for other NLR domains such as N-terminal CARD and C-terminal NACHT and effector molecules RIP2-CARD that have been successfully designed and tested could be further refined using rabbit NOD2 as a template [68,104]. The reliability of the previously reported model structures of fish TLR/NLR after successive and careful refinements [45,67,102,104] using several web-based validation programs are found good. These methods are also routinely used to assess the structural quality of experimentally solved structures. Interestingly, refined structures obtained using a comparative modeling approach for rohu and zebrafish TLRs/NLRs show a reliable validation score with a very minimum structural errors as shown in Fig. 4. Such errors can be further refined using atomic simulation techniques on a time-scale ranging from nanoseconds-microseconds as highlighted in Fig. 3.

5. Structural refinement using molecular dynamics simulations

Structural assessment of modeled structure requires substantial refinement and spatial rearrangement of the initial template structure to improve accuracy [105]. Clashes between side-chain atoms, bonds, dihedral angles etc. can be refined using web-based programs and disordered loops can be refined using advanced modeling approaches (Fig. 3). However, optimization of domain structure and its spatial rearrangement and folding require dynamics over a threshold time-period under a physiological environment. For example, several proteins undergo conformational dynamics and structure rearrangement under a threshold temperature, salt, pH etc. Molecular dynamics (MD) simulations provide a platform to optimize the structural dynamics and folding under a physiological condition [106,107]. Classical MD simulations using all-atom and coarse-grained models have been successfully employed in studying biomolecule structure, dynamics, conformational change, ligand binding etc. [108] This makes it a very powerful tool and has also been applied in refining experimentally obtained protein, DNA and other biomolecule structures. As mentioned in Section 4, a poor homology between the query and template (low sequence identity)
provides a poor quality model structure. In general, the target protein regions for which the template does not have a structure (low identity/similarity or due to low sequence coverage between query and template) are often built with structural errors that require substantial refinement. An example in fish TLRs are the loops connecting each LRR repeat or the linker (shown in blue, Fig. 4, bottom left) connecting the CARD domains in fish NOD2. It should be noted that the flexible loops in LRR are functionally very important and involved in the PAMP interaction. To this end, a homology model of fish TLR/NLR requires a threshold time-scale of all-atom MD simulation for structural optimization as presented in the schematic workflow (Fig. 3). In this review, the application of MD simulations are discussed in line with studying fish TLRs/NLRs, and readers who wish to know a broad application of MD simulation in studying biomolecules are referred to check previous review articles [108–110].

A major application of MD simulation in studying fish innate immune receptors is its use in tracking the interaction between PRRs and PAMPs in real-time under a physiological condition. In this approach, the target PRR molecule(s) and the PAMP(s) are initially placed at a certain distance from each other so that there exist no intermolecular interactions. Both molecules are equilibrated in an explicit solvent environment at physiological conditions and the intermolecular interactions are monitored over a time-scale ranging from nano- to micro-seconds. Such a technique is referred to as “blind docking” due to a lack of any prior knowledge of the PRR active site that binds to the PAMP. Such method is very useful for studying fish TLRs/NLRs that share <50% homology (identity) with human/mouse for which experimentally PAMP binding sites are resolved. The low sequence homology indicates differential binding pockets and PAMP specificity in fish that are very poorly understood. That said an assumption for a conserved PAMP binding site in both fish and human/mouse TLRs/NLRs need complex structure assessment (Fig. 3). To this end, MD simulations showed promising application in evaluating the fish PRR-PAMP complex stability (Fig. 3). In this method, a prior knowledge of PAMP (for example poly(I:C)) binding pocket in the targeted PRR (for example human TLR3) [111] is first retrieved. Molecular docking simulation is carried out allowing the poly(I:C) to find the optimal binding orientation around a predefined binding pocket in fish TLR3 referring to the prior knowledge obtained from human TLR3-poly(I:C) complex. The spatial arrangement of poly(I:C) in fish TLR3 yielding the lowest free energy is predicted to be the key binding site. As mentioned earlier, such exercises require further structural assessment and complex stability analysis in a physiological environment. All-atom MD simulations of the complex assist in structure refinement [112], validating the complex stability, monitoring change in conformation upon PAMP interaction and any possible dissociation/rearrangement of the PAMP around the protein active site (Fig. 3). Taken together MD simulations have proven to be very helpful in retrieving structural information in fish PRRs for which no experimental evidence is available. In the next section, a few structural studies on fish TLRs/NLRs using molecular docking and MD simulations are summarized.

6. Tertiary structure analysis of TLR/NLR domains

The validation and assessment of the quality of the modeled structures in fish are very important prior to their consideration for PAMP-PRR and/or protein–protein interaction analysis. Sahoo et al., for the first time built a 3D model structure for an Indian carp (Labeo rohita; rohu) TLR3 ECD and TIR domain and presented a comprehensive PAMP and protein–protein interaction analysis using model structure of zebrafish TRIF (TIR-domain-containing adapter-inducing interferon-β) [45]. The tertiary folding of rohu (rTLR3)-ECD model having a horse- shoe shape structure show good resemblance with its template mouse TLR3-ECD crystal structure [69] with an average Cα root mean square deviation (RMSD) of ~0.5 Å (Fig. 4, top left). The reported rTLR3-ECD model shows very good validation scores in Ramachandran plot (99.5% residues in allowed regions), a high 3D profiling score (97.2% residues with good compatibility score), acceptable coarse packing quality and planarity [45]. Similarly, rTLR3-TIR model structure consisting of four α-sheets and α-helices align well with the human TLR1-TIR crystal structure [113] (Fig. 4, top right) yielding a Co atom RMSD of ~0.86 Å. Validation reports of the rTLR3-TIR model structure satisfy the acceptable scores for structural analysis.

In another study, Sahoo et al., reported the modeled 3D structures for zebrafish TLR3 and TLR22 ECD domain (zTLR3 and zTLR22) that recognize a conserve PAMP (virus double-stranded RNA) [44]. Interestingly, they observed a significant difference between these two receptors post MD simulation. The zTLR3-ECD model structure closely resembles (horseshoe shape) with the rohu TLR3-ECD 3D model and human/mouse TLR3-ECD crystal structures. On the other hand, post MD simulation, zTLR22-ECD presents a substantial structural rearrangement measuring a backbone RMSD ~12 Å. The structural comparison provides several interesting findings. First, note that zTLR3-ECD and TLR22-ECD share a good sequence homology (25/42%; identity/similarity). Second, as TLR22 is absent in human/mouse, no specific homologies template structure (experimental structure of TLR22-ECD in any organism) is present. Therefore, Sahoo et al., generated 3D models of zTLR3-ECD and zTLR22-ECD using crystal structure of mouse/human TLR3-ECD as template(s) that initially show a less-flattened horseshoe shape structure (Fig. 4, top center). As discussed in the previous section, the powerful application of MD in optimizing the conformation and dynamics of modeled structure is demonstrated in this study using the model zTLR22-ECD structure. The initial compact zTLR22-ECD model undergo a significant conformational change (twist at both terminus; ~15° and 30° twisting at N- and C-terminus, respectively) during the 50 ns MD simulation presenting a flattened horseshoe structure measuring a distance of ~104 Å between N- and C-terminus as compared to ~45 Å before MD simulation (Fig. 4, top center) [44]. This conformational change and flattening of zTLR22-ECD is hypothesized to be crucial to recognize long-sized dsRNA molecules. Structural models of rohu TLR2-ECD, TLR2-TIR and common carp MyD88-TIR constructed using mouse/human TLR2-ECD and MyD88-TIR as templates have shown a very good secondary structure conservation between the target and template. The rohu TLR2-ECD also presents a horseshoe shape as observed for rohu TLR3-ECD composed of 23 LRR domains. Like mouse/human TLR2- or TLR3-ECD, fish TLR2/TLR3-ECD (rohu and zebrafish) LRR domain consist of an array of parallel β-strands facing the concave surface and are connected via loops. This arrangement from a stable hydrophobic core stabilized by hydrogen bonds (H-bond) between the neighboring β-strands and form a horseshoe shape that resembles TLR-ECD structure as observed in higher eukaryotes [33]. The outer or convex surface of the fish TLRs on the other hand is majorly composed of disordered loops with a few regions having short α-helices.

A key difference in fish TLR-ECD structure as compared to NOD-LRR is the secondary structure topology of convex surface. As illustrated in Fig. 4 (compare between zTLR3 and zNOD2-LRR), all fish TLR-ECD LRR domains are mostly unstructured in the convex surface, whereas structured (α-helix) in fish NOD-LRR (Fig. 4, bottom right). It should be noted that this structural difference between fish TLR- and NLR-LRR domain is not unique in fish, rather well conserved in higher eukaryotes including human [114]. The modeled structure of TLR27 consists of 19 LRR domains claimed to be exist only in three fishes by Wang et al. [115] The reported TLR27-ECD structure resembled the rohu TLR22-ECD horseshoe shape structure with the C-terminal region being slightly twisted yielding a flattened structure [115]. A heterodimeric model of TLR1–TLR2 complex is built using comparative modeling in common carp that shares a good fitting with the template human TLR1-TLR2 complex crystal structure [116]. Horseshoe structure is also observed for other fish TLRs that include Tibet fish TLR4 [117], zebrafish TLR5a and TLR5b [82] and miiyu croaker TLR28 [118]. The modeled zebrafish TLR5a-TLR5b heterodimeric structure was reported to have a
very similar structure when compared with the homodimer TLR5-flagellin complex structure resolved using X-ray crystallography [82,93].

The structural investigation of fish NLR have been carried out for all individual domains i.e. CARD, NACHT and LRR in Indian carps or zebrafish [68,102,104]. Because of a low sequence homology, Maharana et al., employed a multi-template modeling approach to build zebrafish NOD1- and NOD2-CARD and RICK(RIP2)-CARD structure for protein-protein interaction study (Fig. 5) [104]. The initial model structures of zebrafish NOD1/2-CARD(a,b) and RIP2-CARD showed a good secondary structure conservation with each CARD composed of six α-helices (α1–α6) with 100% residues fall in the allowed regions of Ramachandran plot. Except α1 that shows a kink structure, but conserved in NOD and RIP2-CARD, all other five α-subdomains in the modeled structure show a regular shape. Importantly, electrostatic surface potential mapping highlighted a significant difference between zebrafish NOD-CARD and RIP2-CARD. The positive (basic) surface potential (blue) in NOD1/2-CARD is less as compared to RIP2-CARD or the acidic surface potential (red) is higher in NOD1/2-CARD as compared to RIP2 (Fig. 5) [104]. This observation underlie NOD-CARD require an exposed negative surface potential to bind RIP2-CARD in order to transduce the downstream signaling cascade.

The simulated zebrafish NOD2-CARD structure rearranged the bilobed globular structure representing subdomains CARDa and CARDb connected via a linker showing in blue (Fig. 4, bottom left) [104]. The simulated model structures of zebrafish CARD domain show a high resemblance with the high-resolution NMR structure of NOD1 CARD [119]. Maharana et al., in another study predicted the model structures of the zebrafish NOD1 and NOD2 NACHT domain using the crystal structure of mouse NLRC4 (PDB ID: 4KXF) as a template to explore the NOD mediated signal transduction [68]. It should be noted that these models can be further refined using the rabbit NOD2 crystal structure (PDB ID: 2Y0K).

\[\text{Fig. 5. Tertiary structure of zebrafish NOD-CARD domains modeled using template-based modeling approach as indicated. The CARD modeled structures shown on the top composed of six }\alpha\text{-helices (}\alpha_1 \text{– }\alpha_6\text{) and charged residues are highlighted. Sequence alignment of zebrafish NOD1/2-CARD with human and mouse is shown in the center with conserved residues highlighted in different colors. Electrostatic surface potential of zebrafish NOD-CARD and downstream interacting RIP2 CARD domains are shown in the bottom at the indicated colors. The figures are reproduced from Ref. 104 with permission from The Royal Society of Chemistry.}\]
S1RM/S1RL) that gives a better query-template sequence-structure alignment. The modeled zebrafish NOD1–NACHT and NOD2–NACHT as shown in Fig. 4 (bottom center) presented ~97% residues within the allowed regions of Ramachandran plot. Yet again, structural comparison and interpretation between NOD1– and NOD2–NACHT in zebrafish do not show a substantial difference in secondary structure (5 conserved parallel β-sheets and 13 α-helices) and folding as per the key functional subdomains NBD, HD1 and WHD are concerned. A notable difference is the presence of two additional β-sheets that are present in NOD2–NACHT (C-terminus), but absent in NOD1–NACHT. As mentioned earlier, the reported NOD-LRR domains are comparatively short and differ from TLR-LRR having helices on the convex surface. In fish the structure for the NOD-LRR domain is reported only for rohu and zebrafish NOD1 and NOD2 although NOD proteins are identified in several other fish species.

7. Structural interactions between PAMP and PRR

The binding interactions between PAMP and PRR has been comprehensively investigated for both TLRs and NLRs using experimental and computational biophysical techniques. However, in fish these structural information are very poorly explored. In the absence of experimental PRR structures in fish (except TLR5–ECD) [93], modeled TLR/NLR proteins have been used for structural analysis. As mentioned in Section 5, molecular docking simulations and blind-docking using MD simulations have been adopted to reveal PAMP interaction with fish TLR/NLRs. In this section, studies in fish describing the structural interaction between PAMP and TLR or NOD are discussed.

Interaction of a variety of natural and synthetic PAMP that includes poly(I:C), flagellin, iE-DAP, MDP, LPS, zymosan, PGN and LTA with fish TLR/NOD receptors is reported. Molecular docking simulations using programs like AutoDock [120], FlexX [121], Glide [122], GOLD [123] and ArgusLab [124] is used to predict the binding sites for small-size PAMPs containing less number of atoms (poly(I:C), iE-DAP, MDP LPS, zymosan, PGN and LTA). For molecules with a large number of atoms (e.g. dsRNA and LPS), programs like HADDOCK [125], HexServer [126] and PatchDock [127] are used to build the PAMP-PRR complex. It is important to note that, the use of a combination of docking programs for a target molecule binding with the receptor could minimize the probability of non-specific interaction sites. For example, Sahoo et al., compared the binding energy and interaction sites for poly(I:C) using two different programs (AutoDock and ArgusLab) by varying the grid size (box surrounding specific interaction sites as reported previously for homologous proteins, and box embedding whole protein that allows the PAMP to identify the best binding pose). The results obtained in this study showed both the programs predict a conserved binding pocket for poly(I:C) in rohu and zebrafish TLR3 and TLR22 proteins [44].

Two different binding sites are predicted for poly(I:C) in zebrafish TLR3 (LRK2–3 and LRK18–19) and TLR22 (LRKNT-3 and LRK22–24), and a comparatively high poly(I:C) binding affinity is shown for TLR22 (Fig. 6, top left). An additional binding site is reported for poly(I:C) in rohu TLR3 (LRK4–6, LRK13–14 and LRK20–22) (Fig. 6, top center) [45]. Remarkably, this study further highlights the effect of mutation on the binding affinity of poly(I:C) with rohu TLR3. Selective mutations at LRK4–6 and LRK20–22 completely abolish poly(I:C) interactions, whereas LRK13–14 do not completely abolish poly(I:C) binding, but reduce the binding affinity [45]. Chakrapani et al., probed the binding interaction of poly(I:C) with a homology model structure of wild-type and mutant rohu TLR22 and observed complex instability in TLR22 mutant [128]. These findings highlights, docking simulation could be a powerful method to screen potential molecules for TLR activation. But, unfortunately, in fish no structure-based PAMP screening has been carried out yet. Voogdt et al., reported a heterodimeric complex structure for zebrafish TLR5 interacting with bacterial flagellin. They compared the structural differences in the TLR5-flagellin hetero-complex composed of two variable copies of TLR5 (TLR5a and TLR5b) discovered in zebrafish with the experimental homodimer TLR5 structure. This study identified that flagellin binding induces conformational change in the loop connecting LRK9–10, while the rest of the TLR5–ECD structure is highly similar to that of unbound flagellin structure [82]. Studies on rohu TLR2 interaction with different variants of PGN, LTA and zymo- san also reported a conserved predicted binding site by AutoDock, FlexX and GOLD [46].

The interaction of bacterial flagellin with zebrafish TLR5–ECD and its complex has been solved using X-ray crystallography. The expression and purification of TLR has been a roadblock for structure determination. Yoon et al. successfully expressed and purified zebrafish TLR5 by truncating the C-terminal residues that generates N-terminal fragments having 6, 12 and 14 LRR motifs [93]. The apo structure of zebrafish orthologue TLR5 with 12 N-terminal LRR motifs (PDB ID: 3V44) solved with a resolution of 2.83 Å show a horseshoe shape structure (Fig. 7). For PAMP interaction analysis, three variable fragments (D1–D3) of Salmonella flagellin are generated that primarily form a 1:1 complex that further homodimerizes to generate a symmetric 2:2 complex with TLR5 N-terminal domain where the flagellin domain D1 is positioned on the lateral side of TLR5 ECD (Fig. 7). The high-resolution complex structure obtained at 2.47 Å reveals a unique binding activity of flagellin that mediates a tail-to-tail dimerization of TLR5 containing 14 LRR motifs (PDB ID: 3V47). The conserved domain D1, but not the hypervariable domain D2 in flagellin is identified to directly interact with zebrafish TLR5 (LRR-NT to LRR10). Interaction of Bacillus subtilis flagellin with zebrafish TLR5 is also studied using crystallography and a 2.1 Å complex structure (PDB ID:5CV2) is reported [129]. Unlike Salmonella flagellin binding, the interaction of Bacillus subtilis flagellin composed of only two domains (D0 and D1) with zebrafish TLR5 (LRKNT-LRR14) show to form a 1:1 complex. Importantly, this study reported the binding of flagellin to TLR5 did not significantly alter the curvature of the LRR domain and its horseshoe shape (Fig. 7). Structural studies identify zebrafish TLR5 binding with the variable lymphocyte receptor (VLR) blocks flagellin interactions (Fig. 5, PDB IDs: 6BKA and 6BKC) [130]. VLRs are specialized adaptive immune receptor that binds to antigens via the TLR5 LRR domains. Two variable epitopes on TLR5 N-terminal ECD that are involved in flagellin binding are identified to interact with the VLR (Fig. 7).

The binding sites and affinity of poly(I:C), LPS and iE-DAP with rohu NOD1 receptor LRR domain was compared using four different docking programs (AutoDock, FlexX, GOLD and Glide) [102]. This study proposed two binding sites for poly(I:C) that are identified to be consistent in all the programs based on the binding score (Fig. 6, top right). An interesting observation from this study is the binding site predicted for poly(I:C) in rohu NOD1 differs from that of human NOD1 yielding a minimum docking score in all programs. On the other hand, a similar approach for iE-DAP interaction with rohu NOD1 using these programs predicted a binding site that is conserved in both human and rohu (Fig. 6, top right). The in silico structural findings could provide crucial mechanistic insights on the unique features of fish NLRs in recognizing PAMPs. In the absence of any prior knowledge to guide the docking simulations for novel PAMPs, ligand binding sites can be predicted using the model fish TLR/NLR structure. Following the predictions, grid boxes can be generated targeting the predicted pockets and binding affinity of the novel PAMP can be estimated. A summary of the interaction of fish TLR/NLR proteins with specific PAMPs and downstream signaling molecules are summarized in Table 2.

Similarly, for large-size molecules (e.g. LPS), multiple docking servers could be helpful in revealing a consensus binding site as demonstrated for rohu NOD1–LPS complex using Hex and PatchDock server [102]. In addition to the docking simulation, as discussed in Section 5, it is crucial to decipher the stability of the complex in a physiological environment (solvent, pH, salt and temperature). Following docking simulation, a next recommended step is thus to evaluate the complex stability by MD simulations on a time-scale of nanoseconds to microseconds/milliseconds (Fig. 3). Often specific/non-specific binding of a
targeted PAMP can be investigated looking at its atomistic interaction map before and after MD simulations. Maharana et al., compared the stability of the binding pocket for MDP in rohu NOD2 before (complex obtained from docking simulations) and after MD simulation (docked complex after undergoing multi-nanoseconds MD simulation) [103]. The comparative analysis showed a change in the binding site and number of H-bonds between MDP and rohu NOD2 before (Arg97 and Asn100 forms three H-bonds with MDP) and after MD simulations (Asn72 form two H-bonds with MDP), but do not alter the overall ligand binding pocket (Fig. 6, bottom right) [103]. In another approach, Maharana et al., probed the binding sites of ATP with zebrafish NOD1 or NOD2 NACHT domain using molecular docking and MD simulations. They compared the stability of ATP-NOD1/NOD2 complex in zebrafish obtained from AutoDock and a complex where ATP is manually placed ~3 Å from the active site following MD simulations [68].

Docking simulations integrated with all-atom MD simulations are proposed to be effective to probe interaction of virus dsRNA and TLRs. Unlike to small-ligand docking (blind docking), the dsRNA docking is driven by guiding the docking program with the interaction sites that are either derived from a homology analysis (human/mouse TLR3 crystal structure) [69,131], or obtained from poly(I:C) binding analysis which is a synthetic analog of dsRNA. HADDOCK docking program has shown to decipher the complex structure for TLR3/TLR22-ECD complex with dsRNA (Fig. 6, bottom left) derived from different viruses that include AGCRV, VHSV and Infectious Hematopoietic Necrosis Virus (IHNV). These large complex docked structures are simulated on a time-scale of nanoseconds to retrieve structural and mechanistic details at an atomic level. An interesting observation from MD simulation of dsRNA-TLR22 complex in zebrafish is an enhancement of the complex stability as compared to poly(I:C)-TLR22 complex [44]. The overall dynamics of the TLR22-ECD protein is shown to be greatly influenced by the dsRNA binding (Fig. 6, bottom left) which is investigated using an array of essential dynamics methods using the MD trajectory. The stability of the complex (both small-size and large-size PAMP) can be evaluated by computing the free binding energy between the receptor (TLR/NOD) and ligand (PAMP). Using this method, Sahoo et al. and Maharana et al., demonstrated the PAMP interaction specificity and stability in both rohu and zebrafish [44,67]. In this approach, from the MD trajectory of PAMP-PRR complex, a number of structures are retrieved at a specific time-interval. The driving force of PAMP interaction with fish...
TLR/NOD can be next computed using programs like MM/PBSA and MM/GBSA that provides a set of energetic parameters to better understand the molecular forces governing the PRR-PAMP interaction [132].

8. Probing downstream signaling via protein-protein interactions

The interaction between TLR/NOD receptors with downstream signaling molecules are important to understand the molecular process governing the innate immunity activation. Understanding this molecular process in real-time at an atomic-level is challenging due to the involvement of complex molecules that include PAMP, PRR, membrane and downstream effector molecules. However, this process can be dissected to reveal atomic information. To this end, unlike the PAMP-TLR/NOD interaction analysis, in fish the protein-protein interaction studies are miniscule. Moreover, to date no structural studies have been carried out for any fish TLRs in a membrane system.

The first reported study in fish reveals the protein-protein complex between rohu TLR3-TIR and zebra fish TRIF domain [45]. In the absence of any prior binding site information, the complex structure of TIR-TRIF is generated using the predicted protein-protein interactions sites by online web-servers (consPPISP [133] and InterProSurf [134]) [45]. Using a similar approach, rohu TLR2-TIR and its interaction with the

| TLR/NLR | PAMP/Protein | Organism | Experimental method |
|---------|--------------|----------|---------------------|
| TLR2    | PGN, LTA and Zymosan | Labeo rohita | Comparative modeling |
| TLR2    | MyD88-TIR    | Labeo rohita | Comparative modeling |
| TLR3    | Poly(I:C)    | Labeo rohita | Comparative modeling |
| TLR3    | dsRNA (AGCRV,VHSV and IHNV) | Labeo rohita | Comparative modeling |
| TLR3    | Poly(I:C)    | Danio rerio | Comparative modeling |
| TIR     | Flagellin (Bacillus, Salmonella) | Danio rerio | Comparative modeling |
| TLR5    | Variable lymphocyte receptor | Danio rerio | X-ray crystallography |
| TLR22   | Poly(I:C)    | Danio rerio | X-ray crystallography |
| TLR22   | dsRNA (AGCRV,VHSV and IHNV) | Danio rerio | Comparative modeling |
| TLR27   | n/a          | Latimeria chalumnae | Protein threading |
| NOD1    | Poly(I:C), iE-DAP and LPS, ATP | Labeo rohita | Protein threading |
| NOD2    | MDP          | Labeo rohita | Protein threading |
| NOD2    | MDP          | Danio rerio | Protein threading |
| NOD1    | RIP2-CARD    | Danio rerio | Protein threading |
| NOD2    | RIP2-CARD    | Danio rerio | Protein threading |
downstream molecule myD88-TIR (common carp) domain is reported (Fig. 8, top left) [46]. The TIR-TIR interface in rohu is predicted using web-programs like cons-PPISP, ProSurf and PatchDock, and a complex structural model of TIR-TIR is build using HADDOCK program. This study predicted key interface residues contributed by rohu TLR2-TIR and common carp MyD88-TIR domains that majorly include the loops and $\alpha$-helices (Fig. 8, top left) [46]. The reported model structure of golden pompano MyD88 resembles common carp structure, however no protein-protein interactions has yet been reported [135]. Fish TLRs interaction with downstream protein molecules reported to date are listed in Table 2.

The interaction between zebrafish NOD2-CARD domain and CARD domain of the adaptor molecule RIP2 (serine–threonine kinase-2) is investigated using molecular dynamics simulation. The CARD-CARD interaction between zebrafish NOD1/NOD2 and RIP2 is suggested to be mediated by charge-charge interactions as both CARD domains share an opposite electrostatic surface potential (Figs. 5 and 8) [67,104]. Maharana et al., probed the stability of NOD2-RIP2 CARD-CARD complex by generating two complexes by docking the positive surface of NOD2-CARD with two different negative surfaces (Complex-I: Asp494, Glu505, Glu508, and Asp506; Complex-II: Glu505, Glu508, Asp506, and Asp525) of RIP2-CARD [67]. The CARD-CARD protein complex is designed using only the CARDa domain of the zebrafish NOD2 (Fig. 2). From the MD simulation analysis, Complex-I is predicted to have a relatively more stable structure as compared to Complex-II and majorly stabilized by electrostatic interactions and salt-bridges. Together, the study proposed the acidic interface of zebrafish NOD1/NOD2-CARD binds to the basic interface of RIP2-CARD (Fig. 8). The study also anticipated a possible interaction of zebrafish CARDB with RIP-CARD, but are not yet investigated. Considering the importance of signal transduction mediated through CARD-CARD or TIR-TIR interactions, the reported in silico findings could provide insights into (i) specific CARD/TIR domain mutation and its association with fish disease; and (ii) strategies to modulate CARD/TIR domain interactions using small-molecule interventions.

9. Limitations of in silico structural analysis

In silico structural analysis of fish TLR/NLR proteins are often affected by a low sequence-structure homology with higher eukaryotes (Fig. 3), which eventually reflected on the comparative 3D model structure. As discussed in Section 4, model structure of fish TLR/NLR need structural re-modeling and refinement due to the periodic upgrading of the PDB databases. As an example, rohu/zebrafish NOD1/2 structures...
reported by Maharana et al., need further attention and re-modeling using available rabbit NOD2 crystal structure for future structural exercises. Structural errors in the initial model structure could affect the docking simulations and structural interpretations. Although long-range atomic simulations show an alternative promising pathway to docking simulations and structural interpretations. Although long-range atomistic simulations show an alternative promising pathway to docking simulations and structural interpretations. Although long-range atomistic simulations show an alternative promising pathway to docking simulations and structural interpretations. Although long-range atomistic simulations show an alternative promising pathway to docking simulations and structural interpretations. Nevertheless, complex systems like studying the protein-protein interactions and conformational dynamics of TLRs in a membrane bound and unbound state can be achieved on a time-scale of microseconds using the coarse-grained systems.

The inadequacy of experimental evidence for protein-protein complexes with native-like binding surface are challenging, and require experimental support and are still under development. Association of protein-protein complex with a non-native binding surface are found to be stable for hundreds of nanoseconds in MD simulation and required enhanced sampling techniques like “tempered binding” to obtain a native-like complex [136]. Thus, structural interpretation and stability analysis of the protein complex in fish TLRs/NLRs using MD simulation needs to be extended beyond the simplistic measurement of protein association and dissociation. In silico mutagenesis study using docking simulations and binding free energy computation provides important structural information in fish (rohu and zebrafish) [44,45]; however, requires experimental binding constants for structural validation. Lastly, structural interpretation in TLRs/NLRs are often carried out using a truncated functional sub-domain such as ECD, TIR, CARD and NACHT. The influence of connecting sub-domains or membrane (in case of TLRs) on the conformational dynamics (flexibility/rigidity) of the target domain of study are therefore not reflected in the structural analysis. This limits the overall in silico structural analysis both in apo and holo states. For example, the binding kinetics and interface between TIR-TIR in the absence of membrane and ECD can be influenced when studies are performed in a membrane interface using full-length TLR/NLR proteins.

10. Conclusion

A major progress in studying fish TLRs/NLRs has been witnessed over the past ten years. Many studies are centered on the identification and characterization of fish innate immune receptors. That said, very few studies to date have focused on exploring the structural and functional properties of fish innate immune receptors. On the other hand, in humans such studies have been carried out in depth with the development of several innate immune modulators that are under preclinical trials. Bearing in mind the rapid growth of world population, demand of fish in the food supply chain, and recent or past pandemic viral flux caused by viruses with animals as the reservoir hosts, it is crucial to explore fish innate immune receptors more in detail. Furthermore, our understanding of the fish TLR/NLR structure at an atomic level would enable in developing antagonist/agonist ligands to modulate fish immunity and disease, thus directly influencing the global economy. Fish, in particular zebrafish, is emerging as a valuable animal model for preclinical in vivo study and drug screening. This further highlights the importance of studying fish innate immune receptors to discover novel therapeutic strategies that can later be tested in higher organisms including humans.

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