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Citation

Version: Corrected Publisher’s Version
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Note: To cite this publication please use the final published version (if applicable).
MyD88 innate immune function in a zebrafish embryo infection model

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Abstract

Innate immunity signalling mechanisms during vertebrate embryogenesis are largely unknown. To study Toll-like receptor (TLR) signalling function in the zebrafish embryo model, we designed an experimental setup for antisense morpholino knockdown under conditions of bacterial infection. Clearance of *Salmonella typhimurium* Ra bacteria was significantly impaired after knockdown of myeloid differentiation factor 88 (MyD88), a common adaptor protein in TLR and interleukin-1 receptor signalling. Thereby, we demonstrate for the first time that the innate immune response of the developing embryo involves MyD88-dependent signalling, which further establishes the zebrafish embryo as a model to study vertebrate innate immunity.

Introduction

Innate immunity relies heavily on signalling by members of the Toll-like receptor (TLR) family (25). TLRs and associated adaptor molecules are highly conserved between zebrafish (*Danio rerio*) and other vertebrates (13, 18). Bacterial and viral infections were found to induce expression levels of different zebrafish TLR genes (18, 20). However, direct functional evidence to confirm the role of TLR signalling in the innate immune response of zebrafish has not yet been reported.

The exploitation of zebrafish as an animal model to study immunity and infectious diseases is attractive for three main reasons (34, 28, 29, 30). First, high-throughput forward genetic screens in zebrafish are a powerful means to uncover novel immune functions. Second, the optical transparency of the free-living zebrafish embryos makes it possible to examine the early development of the immune system and the progression of microbial infections in real-time (11, 32, 12, 6, 31). Third, the zebrafish embryo is easily accessible to experimental manipulations and efficient inactivation of gene functions can be achieved by injection of antisense morpholino oligonucleotides (19). However, a major obstacle is that many of the immunological details and research tools that are available for more established animal models have not yet been resolved and developed for zebrafish.

The innate immune system of the zebrafish embryo starts developing during the first day post fertilization (dpf). Myeloid precursors originate from the anterior lateral plate mesoderm and migrate to the yolk sac, where they differentiate before the onset of blood circulation (11). Differentiated myeloid cells invade the head mesenchyme tissue or join the blood circulation (11, 32, 12). It has been shown that they are able to phagocytose apoptotic cell corpses (11). Furthermore, myeloid cells show specific adherence to bacteria injected into the blood and phagocytose them rapidly (11, 6, 31). They are also able to sense the presence of bacteria injected into one of the closed body cavities and to respond by migration to the infection site (11). All cells of
the myeloid lineage initially express the transcription factor gene Pu.1 (Spi1), which is essential for their differentiation (21). After 1 dpf Pu.1 expression decreases and two distinct populations of myeloid cells can be distinguished by the expression of two marker genes, \(L\)-plastin, which encodes a macrophage-specific actin-bundling protein, and \(mpx\), which encodes a member of the myeloperoxidase family (11, 3, 17). At 2 dpf, the \(mpx\)-positive cells show the morphological characteristics of neutrophil granulocytes and are able to migrate to sites of trauma (17, 4). Immature lymphoblasts can first be detected by 3 dpf, but T and B lymphocytes do not mature until 4 to 6 weeks after hatching (5, 16). Therefore, the zebrafish embryo model is useful to determine the role of innate immunity in responses to different infectious agents, as it is uncoupled from adaptive immunity. With this approach, Davis et al. (6) showed that, during the first days of development, innate immunity determinants are sufficient for granuloma formation resulting from a mycobacterial infection.

**Results and discussion**

To investigate the potential of the zebrafish embryo as a model to study vertebrate innate immune signalling, we first set out to determine the expression of TLRs and associated adaptor genes during embryo development. Semi-quantitative RT-PCR analysis, using the Superscript II one-step system (Invitrogen) with previously described conditions and primers (18), showed that at least 15 zebrafish TLR genes are expressed at 1 dpf, when the first functional macrophages and neutrophils enter blood circulation (Fig.1). Most of these TLRs are also maternally present, since expression was already detected at the 4-cell stage, which is prior to the onset of zygotic gene expression. Several TLRs display distinct differential expression patterns during early stages of embryogenesis. For example, \(z\mathrm{TLR}_1\) expression peaks during blastula and gastrula stages (dome to 80% epiboly) and is high during embryogenesis compared to the adult stage. Expression of \(z\mathrm{TLR}_3\) peaks during gastrulation and segmentation (80% epiboly to 5-somite stage), is reduced between 1 to 5 dpf, but returns to higher levels in the adult stage. Diffuse \(z\mathrm{TLR}_3\) expression in the developing brain of zebrafish embryos was previously reported (20). A peak in the expression of \(z\mathrm{TLR}_{5a}, z\mathrm{TLR}_{5b}, z\mathrm{TLR}_7, z\mathrm{TLR}_{8a}, z\mathrm{TLR}_{8b}\) and \(z\mathrm{TLR}_{18}\) coincides with the appearance of embryonic macrophages at 1 dpf. Expression of the \(z\mathrm{MyD88}\) adaptor gene is highest in adults. In the embryo, maternal \(z\mathrm{MyD88}\) transcript levels are reduced during blastula and gastrula stages and return to higher levels during segmentation and later stages (Fig.1). The other MyD88-like adaptor genes \(z\mathrm{MAL}, z\mathrm{TRIF}\) and \(z\mathrm{SARM}\) are also maternally present and expressed throughout embryogenesis (Fig.1).

To study innate immunity signalling function in the zebrafish embryo, we targeted \(z\mathrm{MyD88}\), which is known to function as a common adaptor protein in the downstream signalling pathways of all mammalian TLRs, except TLR3, and as an adaptor of the interleukin-1/-18 receptors that are activated through TLR signalling (1, 7, 2). A
morpholino knockdown approach (19) was used to interfere with MyD88 function by inhibition of its mRNA translation. An antisense morpholino (GeneTools) (5’-TAGCAAAACCTCTGTATCCAGCGA-3’) was designed, which targets the leader sequence of the zMyD88 mRNA (DQ100359) at positions -30 to -7 with respect to the ATG. A 5-basepair mismatch control morpholino was used with the sequence 5’-TACCATACCTgTGTTATCGGGA-3’ (mismatches in lower case). For microinjection morpholinos were diluted to different concentrations in Danieiu’s buffer (19) and approximately 1 nl was injected into the blastomere of the 1-2 cell stage embryo. To test the specificity of the MyD88 morpholino and its 5-mismatch control sequence, each of these morpholinos was first coinjected with a zMyD88-EGFP fusion mRNA including the 5’ leader sequence. Embryos co injected with the 5-mismatch control morpholino and zMyD88-GFP mRNA showed clear fluorescence at the 70-90% epiboly stage (Fig.2B,D). In contrast, embryos coinjected with the MyD88 morpholino and zMyD88-GFP mRNA showed only autofluorescence of the yolk (Fig.2A,C), similar as in embryos injected with morpholinos only (Fig.2E,F). Therefore, the MyD88 morpholino effectively blocks translation of zMyD88-GFP mRNA. Coinjection of each of the morpholinos with GFP mRNA (Fig.2G,H) or with a modified zMyD88-GFP mRNA lacking the 5’ leader sequence (Fig.2I,J), resulted in similar fluorescence levels in embryo tissues, confirming that the MyD88 morpholino specifically targets the zMyD88 leader sequence.
Embryos injected with 1.7 to 4 ng of MyD88 or control morpholinos showed no apparent morphological differences with wild type embryos. To determine if myelopoiesis was affected in MyD88 morphants, we analysed the expression of myeloid cell markers. From 1 dpf onwards all myeloid cells of zebrafish embryos express either one of the two markers, *L-plastin* or *mpx* (11, 3, 17, 4). Expression of the macrophage marker gene, *L-plastin*, was examined at 1 dpf, after the onset of blood circulation. To this extent a digoxigenin-labeled antisense riboprobe was synthesized with T7 RNA polymerase from an EcoRI-linearized *l-plastin* cDNA clone (AF157110) and whole-mount in situ hybridization was carried out according to the protocol of Thisse et al. (27). In MyD88 morphants, *L-plastin*-positive macrophages were dispersed over the yolk sac and some had accumulated in the ventral venous plexus (Fig.3A). This pat-
tern was similar to that observed in mismatch control morphants (Fig. 3B) and to the reported $L$-plastin expression pattern of wild type zebrafish embryos (11).

A histochemical staining for myeloperoxidase (MPX) activity (17) was performed to check for the presence of granulocytes in embryos at 2 dpf. Peroxidase-positive cells in both the MyD88 and control morphants were abundantly present in the ventral venous plexus and some were scattered over the yolk surface or had invaded the head region (Fig. 3C-F). The same distribution pattern of granulocytes was observed in non-injected control embryos (data not shown) and has been previously reported, based on both peroxidase staining and $mpx$ gene expression (3, 17). Next, we took ad-

**FIGURE 3.** Development and properties of myeloid cells in MyD88 morphants. Embryos injected with 1.7 ng of MyD88 morpholino (A,C,E,G) or with 1.7 ng of 5-mismatch control morpholino (B,D,F,H) were analysed for $L$-plastin expression in macrophages (A,B) and for myeloperoxidase activity in granulocytes (C-H). (A,B), 1 dpf embryos; (C,D), 2 dpf embryos; (E,F) tails of the embryos shown in C and D; (G,H) tails of 2 dpf embryos analysed 6 hours after wounding of the tail fin. Embryos were grown in 0.003% 1-phenyl-2-thiourea (Sigma) to prevent melanization. Composite images were made of different focal planes.
vantage of an acute inflammation assay devised by Lieschke et al. (17) to determine if the granulocytes of MyD88 morphants were functional. After wounding of the caudal fin with a sharp forceps, peroxidase-positive granulocytes of both MyD88 and control morphants accumulated at the site of trauma within 6 hours, indicating their functional involvement in acute inflammation (Fig.3G,H). In conclusion, based on L-plastin and MPX marker analyses, MyD88 morphants showed no apparent myelopoietic defects.

To demonstrate a function for MyD88 in the innate immune response of the zebrafish embryo, we made use of a Salmonella typhimurium infection model that was previously established (31). In this infection model, a low dose of DsRed-labeled bacteria is injected into the embryo’s bloodstream just after the onset of circula-
tion at 1 dpf. While injection of the wild type *S. typhimurium* strain SL1027 resulted in a rapid lethal infection, its isogenic lipopolysaccharide (LPS) derivative SF1592 (Ra-type LPS mutant) proved to be non-pathogenic (31). In the present study, wild-type embryos, embryos injected with 1.7 ng of MyD88 morpholino, and embryos injected with the control morpholino were challenged in an infection experiment with *S. typhimurium* Ra mutant bacteria containing the DsRED plasmid pGMDs3 (31). Embryos were staged at 28 hpf (15) and individually infected by microinjection of approximately 100 colony forming units (cfu) into the axial vein near the blood island and the urogenital opening as previously described (31). As a control, a similar dose as used in the infection experiment was spotted onto LB agar plates for cfu counting.

Embryos were monitored daily until 6 days after infection with *S. typhimurium* Ra. No differences in survival rate were found between the infected wild-type embryos and MyD88 morphants. However, when embryos were examined for the presence of fluorescent bacteria, the MyD88 morphants showed more red spots, representing bacteria, than the wild-type embryos (data not shown). Therefore, total cfu counts were analysed from groups of five embryos that were sampled at 1 day post infection (dpi), 2 and 6 dpi. The pooled embryos were disintegrated (31) and the mixture was plated on LB agar plates. Four independent infection experiments were performed. At 1 dpi the average number of total cfu was approximately 5-fold higher in the MyD88 morphants as compared to wild type and mismatch control embryos (Fig.4). At 2 dpi, the difference between MyD88 morphants and wild type embryos was 10-fold and significant at p<0.05. Although it is not likely that morpholino knockdown is completely penetrant after 3 days of embryo development (19), a further increase of total cfu was still observed in MyD88 morphants at 6 dpi. At this stage the difference with total cfu in wild type and mismatch control embryos was significant at p<0.01. Between different experiments MyD88 morphant embryos harboured 100- to 1000-fold more bacteria than wild-type and mismatch control embryos, which had either completely cleared the infection or contained only low amounts of bacteria not higher than the inoculum size (Fig.4). Therefore we conclude that MyD88 morphants are not able to clear an infection with *S. typhimurium* Ra effectively.

Although there was a clear increase in cfu counts, it is interesting that infection with the normally non-pathogenic LPS Ra mutant of *S. typhimurium* was not lethal for MyD88 morphant embryos, indicating that multiplication of *S. typhimurium* Ra is not completely uncontrolled in MyD88 morphant embryos. Future analysis of a stable MyD88 knockout line should clarify if this was due to incomplete loss of MyD88 function in morphant embryos or due to MyD88-independent innate immunity mechanisms.

To investigate if the inability of MyD88 morphants to clear *S. typhimurium* Ra bacteria could be due to a defect in phagocytosis, embryos were examined at 1 h post infection with a Leica MZ 16 FA microscope. Ds-red labeled bacteria were observed
inside macrophages of MyD88 morphants (Fig. 5), similar as in wild type embryos or in embryos injected with the mismatch control morpholino. Although we cannot yet exclude differences in phagocytosis efficiency or phagosome maturation, our present observations suggest that MyD88 morphants are primarily affected in activation of the bacterial killing mechanisms.

In conclusion, we have shown that TLRs are broadly expressed during zebrafish embryo development and that MyD88 is required for a wild-type response of zebrafish embryos to S. typhimurium Ra infection. These results indicate that MyD88-dependent signalling functions and is important in the early innate immune responses of embryonic zebrafish, independent from coupling to adaptive signalling responses. Furthermore, we have shown that zebrafish embryos express other MyD88-like adaptor molecules, such as Mal, TRIF and SARM, suggesting that MyD88-independent signalling pathways also exist in zebrafish, similar as in other vertebrates (14, 33, 8). The critical role of zebrafish MyD88 is consistent with many infection studies in MyD88-/- adult mice, which showed increased susceptibility to a variety of pathogens (26, 9, 23, 10, 22, 24). Therefore, the present study validates the zebrafish embryo as a useful model for analysis of the vertebrate innate immune system, which creates exciting possibilities for future studies in zebrafish embryo infection models.

**FIGURE 5.** Presence of S. typhimurium Ra bacteria inside macrophages of a MyD88 morphant embryo. MyD88 morphant embryos were infected at 28 hpf by injection of DsRed-expressing S. typhimurium Ra bacteria into the axial vein and images of infected macrophages in the yolk sac circulation valley were taken after 1 h using a Leica DC500 camera and MZ 16 FA microscope. (A) Bright-field image showing a group of macrophages (m) and erythrocytes (e). (B) Fluorescence image of S. typhimurium Ra in the same location as the macrophages. (C) Overlay image of A and B, indicating the ability of macrophages of MyD88 morphants to phagocytose bacteria. Scale bar: 10 μm.
Acknowledgments

The authors thank Ben Appelmelk and Ewa Snaar-Jagalska for valuable discussions, Indira Medina Rodriguez for initial testing of morpholinos and Wim Schouten, Teun Tak and Davy de Witt for fish care. Research at the VU Medical Centre was supported by a Horizon Breakthrough grant from the Netherlands Organisation for Scientific Research (NWO) and research at Leiden University was supported by a European Commission 6th Framework Programme grant (contract LSHG-CT-2003-503496, ZF-MODELS).
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