Investigation of the Transcriptomic Response of Atlantic Salmon (Salmo Salar) Exposed to Paramoeba Perurans Using Next Generation Sequencing

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Abstract

Amoebic Gill Disease (AGD), caused by the protozoan extracellular parasite *Paramoeba perurans*, is a disease affecting Atlantic salmon (*Salmo salar*) aquaculture. Many studies to date have investigated the pathogenesis of ADG focusing on the host immune response in the gill after the appearance of clinical symptoms. This study investigated the gill transcriptomic profile of pre-clinical AGD using RNA-sequencing (RNA-seq) technology. RNA-seq libraries generated at 4, 7, 14 and 16 days post inoculation (dpi) identified 29,357 Differentially Expressed Genes (DEGs). RNA-seq data was validated using real-time, quantitative PCR (qPCR) analysis of 10 selected immune genes. DEGs mapped to 224 Gene Ontology (GO) terms, 27 reference pathways in the Kyoto Encyclopedia of Genes and Genomes (KEGG) and 15 Reactome Gene Sets. Immune suppression was evident at 7 dpi, prior to there being any evidence of ADG on the gill, involving signalling pathways for interleukins, Nod-like receptors, B-cell and T-cell receptors, and the differentiation of Th1/Th2 and Th17 cells. The results of this study suggest a mechanism for how *N. perurans* circumvents the host immune response to establish a successful infection, and could potentially lead to the development of novel strategies for AGD mitigation or prevention in aquaculture.

Introduction

Amoebic Gill Disease (AGD) is a proliferative gill disease of marine cultured Atlantic salmon, with the aetiological agent being the free-living protozoan *Paramoeba perurans*. Among the parasitic conditions affecting gill health, AGD is the most significant in terms of prevalence and economic impact. First described in Tasmania in 1985, AGD is now present in all Atlantic salmon producing countries. Infection of the gill with *Paramoeba sp.* is reported to induce a tri-phasic host response, including a localised reaction to adhered parasites, a non-specific immuno-regulatory cell infiltration and advanced hyperplasia with epithelial stratification filaments. Macroscopically, AGD lesions are visible as white raised mucoid patches on the gill, as a result of increased mucus production by mucous cells, which forms the basis of the gross gill score scheme developed by Taylor et al. (2009) to monitor the progress of AGD. In advance of gill pathology, the presence of *P. perurans* can be detected using a diagnostic real-time PCR (qPCR). The treatment for diagnosed AGD is either freshwater (< 3 PSU for 3 hours) or hydrogen peroxide (*H*₂*O*₂) (800–1300 ppm for < 20 minutes) baths.

To date, investigations focusing on differential gene expression AGD in salmon have utilised techniques such as microarray analysis, Polymerase Chain Reaction (PCR), ribonucleic acid sequencing (RNA-seq) and single nucleotide polymorphism (SNP) analysis.

Previous AGD studies identified up-regulation of the pro-inflammatory cytokine, interleukin-1beta (IL-1b) as being the hallmark of advanced stage AGD in Atlantic salmon. A recent study using 2D quantitative RT-PCR found AGD-affected gills displayed an increased mRNA expression of cellular
markers of immune cells, including professional antigen presenting cells (MHC-II, CD4), B cells (IgM, IgT, MHC-II) and T cells (TCR, CD4, CD8) 14.

The down-regulation of tumour suppressor p53 (p53) has been suggested as one of the mechanisms underlying cell proliferation in AGD7. The differential expression of genes involved in oxidative stress has also been reported17.

Th1, Th17 and T-reg pathways were found to be down-regulated in AGD infections initiated with a higher dose of amoeba (5000 amoeba/L) with the Th2 pathway up-regulated by both high and low infection doses (500 and 5000 amoeba /L)10. Th2 cytokines, (il4/13a and il4/13b2), and the mucin Muc5AC have also been found to be up-regulated in the gill in AGD12. The aim of the current study was to investigate the early-phase transcriptomic host response of naïve salmon inoculated with P. perurans using Next Generation Sequencing (NGS), specifically RNA-seq, prior to the onset of gill pathology. As RNA-seq can be used as a discovery-based technique, it does not require prior knowledge of the genomic information related to genes of interest, with the advantage of facilitating the identification of both known and unknown transcripts.

**Results**

Clinical symptoms of AGD were determined by macroscopic examination of the intact gills and were scored ranging from 0 (absence of clinical symptoms) to 5 (extensive lesions covering most of the gills surface) 4. No lesions were evident on the gills in AGD-infected groups at 4 and 7 dpi, with 3/6 fish at gill score 1 at 14 dpi, and 6/6 fish with gill score 1 at 16 dpi. Diagnostic qPCR5 analysis of the gill arch samples confirmed the presence of P. perurans in 5/6 at 4 dpi, 6/6 fish at 7 dpi and 14 dpi and in 5/6 fish at 16 dpi. There was no P. perurans detected in any of the control fish sampled at any time point. Gills were examined microscopically following staining with haematoxylin/eosin at all time points (Fig. 2). Evidence of hyperplasia and fusion of the lamellar epithelium (Fig. 2b) was observed at 16 dpi.

**Mapping RNA-seq reads to reference genome**

The analysis of the reads mapped to the Salmo salar reference genome for the 6 fish from each of the 5 sampling time points (0, 4, 7, 14 and 16 dpi) is presented in supplementary Table S1 online. The sequencing depth for each fish ranged from 21.7M to 35.4M raw reads. The % GC ranged from 47 to 49%. The % clean paired reads ranged from 88.4 to 94.7%. The number of assigned fragments or feature counts range was 14.7M to 25.1M.

**Validation of RNA-seq data using qPCR**

Ten immune genes from the gill RNA-seq data were selected for qPCR validation across all sampling time points including cathelicidin 1 (cath-1), c-type lectin receptor A (clr-A), c-c motif chemokine 4 (cxc4), immunoresponsive 1 homolog/aconitate decarboxylase 1 (irg1/acod1), interleukin-8 (il-8), interleukin-17F (il-17), leukocyte cell-derived chemotaxin-2, (lect2), inducible nitric oxide synthase 2, (nos2), serum
amyloid P (sap) and pentraxin-related protein (ptx3). Primer sequences are provided in supplementary Table S2 online. A Spearman correlation analysis of the RNA-seq and the qPCR data using the Log₂ fold change (FC) average of 6 individual fish per gene was conducted for all 4 time points: 4 dpi: s = 14.5, rho = 0.91, p < 0.0002, 7 dpi: s = 16.5, rho = 0.89, p < 0.0003, 14 dpi: s = 36.2, rho 0.78, p < 0.008, 16 dpi: s = 14.1 rho = 0.91, p < 0.0002. The results of the 16 dpi RNA-seq and qPCR validation data are presented in Fig. 3a with the corresponding correlation data in Fig. 3b.

**Differentially Expressed Genes (DEGs)**

Differentially expressed genes were identified at each of the four experimental time points (4,7,14 and 16 dpi) (Fig. 1), and further divided into up-regulated and down-regulated genes for a total of 8 DEG lists.

A total of 29,357 genes were found to be differentially regulated in the gill as a result of AGD infection, with approximately equal numbers of up- and down-regulated genes (Table 1). The %DEGs at 4, 14 and 16 dpi ranged from 16.5–21.9% with a higher % of genes down-regulated genes than up-regulated (61.7%, 52.4% and 54.7%, respectively). At 7 dpi there were 42.3% DEGs, with more up-regulated than down-regulated (59.4% vs 40.4%).

| Time (dpi) | Up-regulated | Down-regulated | Up- and down-regulated | % of total |
|-----------|--------------|----------------|------------------------|------------|
| 4         | 1,853 (38.3%)| 2,985 (61.7%)  | 4,838                  | 16.5       |
| 7         | 7,408 (59.6%)| 5,024 (40.4%)  | 12,432                 | 42.3       |
| 14        | 2,698 (47.6%)| 2,971 (52.4%)  | 5,669                  | 19.3       |
| 16        | 2,905 (45.3%)| 3,513 (54.7%)  | 6,418                  | 21.9       |
| Total     | 14,864 (50.6%)| 14,493 (49.4%)  | 29,357                 | 100        |

**Gene Ontology (GO) and Pathway Enrichment**

At 4 dpi, the GO:0002768: immune response-regulating cell surface receptor signalling pathway (Fig. 4a) was identified as being enriched up-regulated DEGs with GO: 0042254: ribosome biosynthesis, the most significant term enriched with down-regulated DEGS at 4 dpi (Fig. 4b). No enriched pathways were identified with up-regulated DEGs, however 6 pathways were identified with down-regulated DEGs including dre03008: Ribosome biogenesis in eukaryotes being the most significant (Table 2).

At 7 dpi, fourteen enriched pathways were identified with dre04146: peroxisome being the most significant (Table 2). Seven pathways were identified with the dre04110: cell cycle pathway being the most significantly enriched (Table 2). Pathways relevant to AGD infection included dre04115: p53 signalling pathway and R-DRE-168256: immune system of which subsets included R-DRE-168249: innate immune system R-DRE-1280215: cytokine signalling in immune system and R-DRE-449147: signalling by interleukins (Table 2).
At 14 dpi, the reactome gene set R-DRE-983695: antigen activates b-cell receptor (BCR) leading to generation of second messengers pathway was the only enriched pathway (Table 2). At 14 dpi eight pathways were enriched with dre03030: DNA replication as the most significant (Table 2).

At 16 dpi, one KEGG pathway was found to be enriched, dre00380: Tryptophan metabolism with 4 reactome gene sets also identified including R-DRE-983695: Antigen activates B Cell Receptor (BCR) leading to generation of second messengers (Table 2). At 16 dpi for the down-regulated genes, fifteen pathways were identified (8 KEGG, and 7 reactome gene sets), with the most significant being dre03008: Ribosome biogenesis in eukaryotes (Table 2). Of immune relevance was R-DRE-983170: antigen presentation of class I MHC.
| Time / gene expression direction | Category     | Term     | Description                                           | -10 Log(P) | InTerm / InList |
|---------------------------------|--------------|----------|-------------------------------------------------------|------------|----------------|
| 4dpi/down                       | KEGG Pathway | dre03008 | Ribosome biogenesis in eukaryotes                     | 10.2       | 22/37          |
|                                 |              | dre03040 | Spliceosome                                           | 7.1        | 23/53          |
|                                 |              | dre03050 | Proteasome                                            | 6.9        | 13/20          |
|                                 |              | dre03030 | DNA replication                                       | 6.6        | 15/27          |
|                                 |              | dre00100 | Steroid biosynthesis                                  | 4.7        | 7/9            |
|                                 |              | dre04141 | Protein processing in endoplasmic reticulum           | 4.6        | 20/57          |
| 7 dpi/up                        | KEGG Pathway | dre04146 | Peroxisome                                           | 9.9        | 31/45          |
|                                 |              | dre01212 | Fatty acid metabolism                                | 8.4        | 20/25          |
|                                 |              | dre03010 | Ribosome                                             | 7.5        | 28/45          |
|                                 |              | dre00071 | Fatty acid degradation                               | 7.4        | 17/21          |
|                                 |              | dre00380 | Tryptophan metabolism                                | 6.9        | 14/16          |
|                                 |              | dre00260 | Glycine, serine and threonine metabolism             | 6.8        | 15/18          |
|                                 |              | dre04142 | Lysosome                                             | 6.3        | 29/52          |
|                                 |              | dre00280 | Valine, leucine and isoleucine degradation          | 6.0        | 18/26          |
|                                 |              | dre00340 | Histidine metabolism                                 | 5.9        | 11/12          |
|                                 |              | dre03320 | PPAR signaling pathway                               | 4.9        | 17/27          |
|                                 |              | dre01040 | Biosynthesis of unsaturated fatty acids              | 4.8        | 9/10           |
|                                 |              | dre00410 | beta-Alanine metabolism                               | 4.7        | 12/16          |
|                                 |              | dre00310 | Lysine degradation                                   | 2.9        | 13/24          |
|                                 |              | dre00640 | Propanoate metabolism                                | 2.1        | 9/17           |
| 7 dpi/down                      | KEGG Pathway | dre04110 | Cell cycle                                           | 7.1        | 20/138         |
|                                 |              | dre03030 | DNA replication                                       | 5.3        | 9/38           |
|                                 |              | dre00520 | Amino sugar and nucleotide sugar metabolism         | 3.8        | 9/57           |
| Time / gene expression direction | Category | Term               | Description                                   | -10 Log(P) | InTerm / InList |
|--------------------------------|----------|--------------------|-----------------------------------------------|------------|-----------------|
|                               |          | dre04115           | p53 signaling pathway                         | 2.9        | 9/75            |
|                               |          | dre00230           | Purine metabolism                             | 2.4        | 15/196          |
|                               |          | dre00240           | Pyrimidine metabolism                         | 2.3        | 10/107          |
|                               | Reactome | R-DRE-168256       | Immune System                                 | 4.6        | 51/790          |
|                               | Gene Sets| R-DRE-168249       | Innate Immune System                          | 3.5        | 31/449          |
|                               |          | R-DRE-449147       | Signaling by Interleukins                     | 2.2        | 14/185          |
|                               |          | R-DRE-1280215      | Cytokine Signaling in Immune system           | -2.2       | 16/227          |
| 14 dpi/up                     | Reactome | R-DRE-983695       | Antigen activates B Cell Receptor (BCR) leading to generation of second messengers | 2.7        | 3/6             |
|                               | Gene Sets| R-DRE-983695       | DNA replication                               | 9.6        | 17/27           |
|                               |          | dre03008           | Ribosome biogenesis in eukaryotes             | 8.5        | 19/37           |
|                               |          | dre03040           | Spliceosome                                   | 4.2        | 17/53           |
|                               |          | dre00240           | Pyrimidine metabolism                         | 3.9        | 15/46           |
|                               |          | dre03050           | Proteasome                                    | 3.0        | 8/20            |
|                               |          | dre01212           | Fatty acid metabolism                         | 2.9        | 9/25            |
|                               |          | dre00230           | Purine metabolism                             | 2.2        | 18/84           |
|                               |          | dre00280           | Valine, leucine and isoleucine degradation     | 2.1        | 8/26            |
| 16 dpi/up                     | KEGG     | dre00380           | Tryptophan metabolism                         | 2.1        | 4/16            |
|                               | Pathway  | R-DRE-983695       | Antigen activates B Cell Receptor (BCR) leading to generation of second messengers | 2.6        | 3/6             |
|                               |          | R-DRE-1433557      | Signaling by SCF-KIT                          | 2.6        | 3/6             |
|                               |          | R-DRE-373752       | Netrin-1 signaling                            | 2.3        | 3/7             |
|                               |          | R-DRE-9007101      | Rab regulation of trafficking                 | 2.1        | 5/24            |
| Time / gene expression direction | Category            | Term               | Description                                      | -10 Log(P) | InTerm / InList |
|---------------------------------|---------------------|--------------------|--------------------------------------------------|------------|-----------------|
| 16 dpi/down                     | KEGG Pathway        | dre03008           | Ribosome biogenesis in eukaryotes                | 9.5        | 22/37           |
|                                 |                     | dre03030           | DNA replication                                  | 5.2        | 14/27           |
|                                 |                     | dre00240           | Pyrimidine metabolism                            | 4.4        | 18/46           |
|                                 |                     | dre00100           | Steroid biosynthesis                             | 4.4        | 7/9             |
|                                 |                     | dre00970           | Aminoacyl-tRNA biosynthesis                      | 3.9        | 12/26           |
|                                 |                     | dre03050           | Proteasome                                       | 3.7        | 10/20           |
|                                 |                     | dre03013           | RNA transport                                    | 3.6        | 20/61           |
|                                 |                     | dre03040           | Spliceosome                                      | 3.5        | 18/53           |
|                                 | Reactome Gene Sets  | R-DRE-8953854      | Metabolism of RNA                                | 5.5        | 34/106          |
|                                 |                     | R-DRE-1640170      | Cell Cycle                                       | 3.7        | 32/116          |
|                                 |                     | R-DRE-983170       | Antigen Presentation of Class I MHC             | 2.5        | 3/3             |
|                                 |                     | R-DRE-77286        | Mitochondrial fatty acid beta-oxidation          | 2.5        | 3/3             |
|                                 |                     | R-DRE-212300       | PRC2 methylates histones and DNA                | 2.5        | 3/3             |
|                                 |                     | R-DRE-2262752      | Cellular responses to stress                     | 2.4        | 15/50           |
|                                 |                     | R-DRE-8953897      | Cellular responses to external stimuli           | 2          | 19/74           |

At 4, 14 and 16 dpi process and pathway enrichment identified 14 up-regulated genes with involvement in immune regulation and activation (Table 3). Immune biological processes and a reactome gene set (Table 2) were identified from the DEG lists generated at each time point and are listed in Table 3 and included GO:0002682: Regulation of immune system process (btk, c1cq, CD79b, CD99, cgas, hmgb1a, hmgb3b, Il-34, kita, lyn, tfpi1, themis), GO:0002253: Activation of immune system process (btk, c1cq, CD79b, cgas, hmgb1a, themis), GO:0097190: B-cell differentiation (kita, cd79b and lkzf1.7), GO:0002768: immune response-regulating cell surface receptor signalling pathway (kita, btk, cd79b, themis2). The reactome gene set is R-DRE-983695: Antigen activates B Cell Receptor (BCR) leading to generation of second messengers (dapp, btk, and CD79b).
Table 3
Up-regulated genes with involvement in immune regulation and activation

| Gene ID  | Gene Name                                      | Biological Process and Pathway                                           | dpi     |
|----------|------------------------------------------------|-------------------------------------------------------------------------|---------|
| 568653   | btk                                            | Bruton agammaglobulinemia tyrosine kinase                                | 14, 16  |
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
|          |                                                | R-DRE-983695                                                            |         |
| 449803   | c1qc                                           | complement component 1, q subcomponent, C chain                         | 14, 16  |
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
| 100329481| cd79b                                          | CD79b molecule, immunoglobulin-associated beta                           | 4, 14, 16|
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
|          |                                                | GO:0097190                                                              |         |
|          |                                                | GO:0002768                                                              |         |
|          |                                                | R-DRE-983695                                                            |         |
| 559896   | cd99                                           | CD99 molecule                                                            | 14      |
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
| 557043   | cgas                                           | cyclic GMP-AMP synthase                                                  | 14, 16  |
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
| 550386   | dapp1                                          | dual adaptor of phosphotyrosine and 3-phosphoinositides                 | 14, 16  |
|          |                                                | R-DRE-983695                                                            |         |
| 321622   | hmgb1a                                         | high mobility group box 1a                                              | 14, 16  |
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
| 550466   | hmgb3b                                         | high mobility group box 3b                                              | 14      |
|          |                                                | GO:0002682                                                              |         |
| 560193   | il34                                           | interleukin 34                                                          | 14      |
|          |                                                | GO:0002682                                                              |         |
| 30256    | kita                                           | KIT proto-oncogene, receptor tyrosine kinase a                          | 4, 14, 16|
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0097190                                                              |         |
|          |                                                | GO:0002768                                                              |         |
| 30177    | ikzf1                                          | IKAROS family zinc finger 1                                             | 14, 16  |
|          |                                                | GO:0097190                                                              |         |
| 447804   | lyn                                            | LYN proto-oncogene, Src family tyrosine kinase                          | 14      |
|          |                                                | GO:0002682                                                              |         |
| 560339   | tfpi2                                          | tissue factor pathway inhibitor 2                                       | 14      |
|          |                                                | GO:0002682                                                              |         |
| 100535600| themis2                                        | thymocyte selection associated family member 2                          | 4, 14, 16|
|          |                                                | GO:0002682                                                              |         |
|          |                                                | GO:0002253                                                              |         |
|          |                                                | GO:0002768                                                              |         |

At 7 dpi the reactome gene set R-DRE-168256: immune system (Table 2) was associated with 51 down-regulated genes (Table 4). Within this reactome gene set, 16 genes were associated with R-DRE-1280215: cytokine signalling in the immune system with all but 2 of these genes associated with R-DRE-449147:
signalling by interleukins (Table 2). Screening of the DEG lists at 7 dpi identified multiple transcripts for 9 members of the interleukin family as being down-regulated (IL-1β, IL-8, IL-11, IL-12β, IL-15, IL-17D, IL-17F, IL-18, IL-34) with only one member, IL-27β significantly up-regulated (12.4 log₂FC) (supplementary Table S3 online). Functional enrichment analysis (string-db.org) using the 51 genes identified in R-DRE-168256: immune system genes at 7 dpi (Table 4) identified involvement in the Nod-like receptor signalling pathway (hsa04621, FDR 8.58e-08: cxcl8, hsp90ab1, ikkb, irf9, mapk3, nlrx1, sugt1, tbk1, traf3), B-cell receptor signalling (hsa-04662, FDR 0.0051: ikkb, mapk3, sky), T-cell receptor signalling (hsa-04660, FDR 0.0106: dlg1, ikkb, mapk3), Th1/Th2 cell differentiation (hsa-04658, FDR 0.00084: ikkb, jak3, mapk3, stat4) and Th17 cell differentiation (hsa-04659, FDR 0.0011: hsp90ab1, ikkb, jak3, mapk3).

The pattern and extent of DEGs at 7 dpi was different from the other time points with more genes showing differential expression (42%) and with more genes being up-regulated (59.6%) (Table 1).

Analysis of the DEG lists at 7 dpi found that of the top 100 up-regulated genes there were 87 characterised and 69 unique transcript (supplementary Table S4 online). The most up-regulated gene was mannan binding lectin serine peptidase 2 (masp2, 30.0 log2FC). Thirteen other complement transcripts were also up-regulated including (C1q-like protein 2 (x4), C1q-like protein 3, C2-like, C3-like (x2), C9, factor B, factor H, properdin-like (x2). Acute Phase Response (APR) genes in the top 100 included pentraxin, alpha-1-antitrypsin (x2), alpha-2-macroglobulin (x2), fibrinogen (α, β, γ), leukocyte cell-derived chemotaxin-2 (lect2), serum amyloid P-component (SAP).

The top 100 down-regulated genes at 7 dpi yielded 78 unique transcripts and 19 uncharacterised genes (supplementary Table S5 online). The genes that were represented by multiple transcripts included C-C motif chemokine 4-like (2x), collagenase 3-like (2x), interferon-induced very large GTPase 1-like (2x), NADPH oxidase organizer 1-like (2x). The gene with the most down-regulated expression was interferon-induced guanylate-binding protein 1-like (log₂FC, -7.5). Multiple members of the same gene family included the c-c chemokine family (cc4, cc20), the interleukins (il-1b, il-17f, il-17 receptor E), and mucins (muc2, muc7).
| Gene ID   | Gene Name                  | Reactome Gene Sets                          |
|----------|----------------------------|---------------------------------------------|
| 336425   | aldoaa, aldolase a, fructose-bisphosphate, a | R-DRE-168256, R-DRE-168249                  |
| 114428   | ARF1, ADP-ribosylation factor 1 | R-DRE-168256                                |
| 415204   | ARPC3, actin related protein 2/3 complex, subunit 3 | R-DRE-168256, R-DRE-168249                  |
| 767630   | ATPase phospholipid transporting 8A1 | R-DRE-168256, R-DRE-168249                  |
| 192322   | CALM2B, calmodulin 2b, (phosphorylase kinase, delta) | R-DRE-168256, R-DRE-168249                  |
| 334527   | CAP1, CAP, adenylate cyclase-associated protein 1 | R-DRE-168256, R-DRE-168249                  |
| 394037   | CCT8, chaperonin containing TCP1, subunit 8 (theta) | R-DRE-168256, R-DRE-168249                  |
| 567192   | CD59, CD59 molecule (CD59 blood group) | R-DRE-168256, R-DRE-168249                  |
| 767754   | CENPE, centromere protein E | R-DRE-168256                                |
| 336381   | COFL1, coactosin-like F-actin binding protein 1 | R-DRE-168256, R-DRE-168249                  |
| 100134935 | CSF3R, colony stimulating factor 3 receptor (granulocyte) | R-DRE-168256, R-DRE-1280215, R-DRE-449147   |
| 30265    | CTNNB1, catenin (cadherin-associated protein), beta 1 | R-DRE-168256                                |
| 100002946 | CXCL8A, chemokine (C-X-C motif) ligand 8a | R-DRE-168256                                |
| 336613   | CYFIP1, cytoplasmic FMR1 interacting protein 1 | R-DRE-168256, R-DRE-168249                  |
| 324089   | DET1, DET1 partner of COP1 | R-DRE-168256                                |
| 114446   | DLG1, discs, large homolog 1 (Drosophila) | R-DRE-168256, R-DRE-1280215, R-DRE-449147   |
| 100005297 | EPGN, epithelial mitogen homolog (mouse) | R-DRE-168256, R-DRE-1280215, R-DRE-449147   |
| 562999   | HECD2, HECT domain containing | R-DRE-168256                                |
| 30573    | HSPO9AB1, heat shock protein 90, alpha (cytosolic), class B member 1 | R-DRE-168256, R-DRE-168249                  |
| 563560   | IKBKB, inhibitor of nuclear factor kappa B kinase subunit beta | R-DRE-168256, R-DRE-168249, R-DRE-1280215   |
| Gene ID | Gene   | Name                                         | Reactome Gene Sets                                      |
|--------|--------|----------------------------------------------|--------------------------------------------------------|
| 560193 | il34   | interleukin 34                               | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 403013 | irf9   | interferon regulatory factor 9               | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 561370 | jak3   | Janus kinase 3 (a protein tyrosine kinase, leukocyte) | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 555969 | lpcat1 | lysophosphatidylcholine acyltransferase 1     | R-DRE-168256, R-DRE-168249                             |
| 399480 | mapk3  | mitogen-activated protein kinase 3            | R-DRE-168256, R-DRE-168249, R-DRE-1280215, R-DRE-449147 |
| 100537196 | mapkap1 | MAPK associated protein 1                  | R-DRE-168256                                           |
| 373081 | mvp    | major vault protein                          | R-DRE-168256, R-DRE-168249                             |
| 569779 | myo10  | myosin X                                     | R-DRE-168256, R-DRE-168249                             |
| 557335 | nlrx1  | NLR family member X1                        | R-DRE-168256, R-DRE-168249                             |
| 60658  | nos1   | nitric oxide synthase 1 (neuronal)           | R-DRE-168256, R-DRE-168249                             |
| 404036 | nos2a  | nitric oxide synthase 2a, inducible          | R-DRE-168256, R-DRE-168249                             |
| 796461 | nrg1   | neuregulin 1                                 | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 554967 | psmd1  | proteasome 26S subunit, non-ATPase 1         | R-DRE-168256, R-DRE-168249                             |
| 373104 | rab14  | RAB14, member RAS oncogene family            | R-DRE-168256, R-DRE-168249                             |
| 323197 | racgap1 | Rac GTPase activating protein 1              | R-DRE-168256                                           |
| 554089 | s100z  | S100 calcium binding protein Z               | R-DRE-168256, R-DRE-168249                             |
| 402992 | scamp1 | secretory carrier membrane protein 1         | R-DRE-168256, R-DRE-168249                             |
| 793290 | spred2b| sprouty related EVH1 domain containing 2b    | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 368519 | stat4  | signal transducer and activator of transcription 4 | R-DRE-168256, R-DRE-1280215, R-DRE-449147              |
| 492489 | sugt1  | SGT1 homolog, MIS12 kinetochore complex assembly cochaperone | R-DRE-168256, R-DRE-168249                             |
| 405769 | syk    | spleen tyrosine kinase                       | R-DRE-168256, R-DRE-168249, R-DRE-1280215, R-DRE-449147 |
## Discussion

RNA-seq compared the gene expression from uninfected control with AGD-infected gill prior to (4, 7 dpi) and after (14, 16 dpi) the appearance of AGD lesions. A total of 29.7K genes identified in naive Atlantic salmon as being altered following inoculation with *P. perurans*. No gill pathology was observed macroscopically in fish sampled up to 14 dpi, however diagnostics qPCR detected the presence of *P. perurans* on 5/6 and 6/6 fish at 4 and 7 dpi, respectively, with 17.3K genes showing altered expression during this time, demonstrating the sensitivity of this method of detection. Up-regulated immune genes were identified at 4, 14 and 16 dpi with down-regulated immune genes identified at 7 dpi.

At 4, 14 and 16 dpi, three genes CD79b, KIT proto-oncogene, receptor tyrosine kinase a (kita) and thymocyte selection associated family member 2 (themis2) were consistently up-regulated. CD79, as CD79a/CD79b heterodimers (α/β), form part of the B-cell antigen receptor (BCR) with membrane immunoglobulin molecules. The BCR complex plays a crucial role in B cell development and antibody production following antigen exposure. On activation, the B-cell receptor ‘signalosome’ initiates multiple signalling cascades that involves kinases, GTPases, and transcription factors resulting in changes in cell metabolism, gene expression, and cytoskeletal organisation\textsuperscript{18}.

| Gene ID   | Gene | Name                                                                 | Reactome Gene Sets                                                                 |
|-----------|------|----------------------------------------------------------------------|----------------------------------------------------------------------------------|
| 100333043 | tap1 | transporter 1, ATP-binding cassette, sub-family B (MDR/TAP)          | R-DRE-168256                                                                     |
| 692289    | tbk1 | TANK-binding kinase 1                                               | R-DRE-168256, R-DRE-168249, R-DRE-1280215, R-DRE-449147                           |
| 100333821 | tnfrsf11a | TNF receptor superfamily, member 11a, NFKB activator              | R-DRE-168256, R-DRE-1280215                                                      |
| 564279    | tnip2 | TNFAIP3 interacting protein 2                                       | R-DRE-168256, R-DRE-168249, R-DRE-1280215                                       |
| 100331669 | tpp2 | tripeptidyl peptidase 2                                             | R-DRE-168256                                                                     |
| 406335    | uba1 | ubiquitin-like modifier activating enzyme 1                         | R-DRE-168256                                                                     |
| 393934    | ube2d2 | ubiquitin-conjugating enzyme E2D 2 (UBC4/5 homolog, yeast)         | R-DRE-168256, R-DRE-168249                                                       |
| 406807    | ube2na | ubiquitin-conjugating enzyme E2Na                                  | R-DRE-168256, R-DRE-168249, R-DRE-1280215, R-DRE-449147                         |
| 321056    | zgc:63569 | zgc:63569                                                             | R-DRE-168256, R-DRE-168249                                                      |
| 393844    | znrf1 | zinc and ring finger 1                                               | R-DRE-168256                                                                     |

Table 4 Note: R-DRE-168256: immune system, R-DRE-168249: innate immune system, R-DRE-1280215: cytokine signalling in the immune system, R-DRE-449147: signalling by interleukins
Kita is a tyrosine-protein kinase that acts as cell-surface receptor for stem cell factor (Scf) and plays an essential role in the regulation of cell survival and proliferation, hematopoiesis, stem cell maintenance, gametogenesis, mast cell development, migration and function, and in melanogenesis.\textsuperscript{19}

Themis2 is a gene which encodes a protein that plays a regulatory role in both positive and negative T-cell selection during late thymocyte development. The protein functions through T-cell antigen receptor (TCR) signalling, and is necessary for proper lineage commitment and maturation of T-cells. Themis2 plays a role in the in macrophage inflammatory response, promoting LPS-induced TNF production.

Five genes were found to be up-regulated only at 14 dpi: CD99, high mobility group b3b (hmgb3b) interleukin-34 (IL-34), Lyn and tissue factor pathway inhibitor 2 (tfip2). CD99 has been described as a costimulatory molecule on T cells.\textsuperscript{20} High mobility group (HMG) proteins have roles in the nucleus and mitochondria as architectural DNA binding proteins, and in the cytoplasm as signalling regulators, and in the extracellular milieu as inflammatory cytokines.\textsuperscript{21} The molecular function of HMGB3b is DNA binding and it is involved in biological processes that include regulation of transcription by RNA polymerase II and positive regulation of the innate immune response. IL-34, binding to the colony stimulating factor 1 (Csf1), increases growth or survival of monocytes. In fish as in mammals, monocytes, macrophages, and neutrophils are the main phagocytic cells.\textsuperscript{22} Lyn is a Src tyrosine kinase which is also involved in the formation of a B-cell receptor 'signalosome'. Lyn also interacts with Themis2 and lyn activation has been reported to reduce the hypersecretion of mucus and MUC5AC in airway inflammation.\textsuperscript{24} Excessive mucus production in the gills is a hallmark of AGD with substantial up-regulation of the secreted MUC5 detected in clinical AGD.\textsuperscript{12} Specialized epithelial (goblet) cells are the major source of MUC5AC, which can be induced by MMP9 through the activation of the epidermal growth factor receptor (EGFR) and mitogen-activated protein kinase 3/2 MAPK 3/2(ERK1/2) cascade.\textsuperscript{25}

MAPKs are a superfamily of serine/threonine protein kinases that transduce a variety of external signals, leading to an array of cellular responses that include growth, differentiation, apoptosis, and host defence response.

The up-regulation of Lyn is only evident from 14 dpi, when mucoid patches were first identified on the gills. Tfpi2 is a serine protease inhibitor and is thought to play a role in the regulation of plasmin-mediated matrix remodelling and in the activation of matrix metalloproteinases (MMPs) including MMP-1 and MMP-13, and to a lesser extent of MMP-2 and MMP-9.\textsuperscript{27}

Six genes were up-regulated only at the later times of 14 and 16 dpi, and included Bruton agammaglobulinemia tyrosine kinase (Btk), complement component 1, q subcomponent, C chain (c1qc), cyclic GMP-AMP synthase (cgas), dual adaptor of phosphotyrosine and 3-phosphoinositides (dapp1), high mobility group box 1a (hmgb1a) and IKAROS family zinc finger 1 (ikzf1) (Table 3). Btk is expressed in cells with hematopoietic lineage, with the exception of T lymphocytes and natural killer cells, and is involved in a multiple immune signalling pathways. Activated by BCR aggregation, btk constitutes a
major component of the B-cell receptor signalosome complex \(^{18}\) and plays a role in B-cell development and mature B-cell activation \(^{28}\).

C1qC was up-regulated suggesting the classical complement pathway was activated. Composed of C1, C4, and C2 components reacting in this order, the classical pathway primarily recognizes antibodies in immune complexes. However, an increased expression of immunoglobulins were not observed in this study indicating that C1q is potentially interacting with other acute phase molecules.

Cgas is a cytosolic DNA sensor that activates a type-I interferon response \(^{29}\), DAPP is a B-cell-associated adapter that regulates B-cell antigen receptor (BCR) \(^{30}\) and ikzf1, a member of the Ikaros family of proteins are involved in lymphocyte development, including a wide range of processes, such as apoptosis, cell cycle arrest, proliferation, and differentiation \(^{31}\). Btk phosphorylates Ikaros at unique sites within the DNA binding domain, augmenting the nuclear localization and sequence-specific DNA binding activity of the transcription factor function of Ikaros \(^{32}\).

Analysis of the top 100 DEG lists at 7 dpi found evidence of the activation of complement and the acute phase response, with the down-regulation of chemokines, interleukins and interferon-inducible proteins and mucins.

The down-regulation of the immune response at 7 dpi identified by the reactome gene set R-DRE-168256 (Table 2) identified 51 immune-related down-regulated genes (Table 4) with 14 genes associated with R-DRE-449147: signalling by interleukins (Table 2). Further analysis of the DEGs lists at 7 dpi identified 9 interleukin (IL) genes represented by 14 individual transcripts of which 8 genes were down-regulated (IL-1\(^\beta\), IL-8, IL-11, IL-12\(^\beta\), IL-15, IL-17F, IL-18 and IL-34) and only one found to be up-regulated IL-27\(^\beta\) (Supp. Table S3).

IL-1\(^\beta\) has been reported as the hallmark of late stage ADG infection \(^{15}\) with up-regulation demonstrated in the gills with numerous lesions, fused lamellae and epithelial cells hyperplasia. The expression of IL-1\(^\beta\) has been associated with larger AGD-lesions, often showing greater mucous cell hyperplasia, where the mucous cells as the possible source of the IL-1\(^\beta\) \(^{11}\). Macroscopically, the progression of ADG in the current study reached gill score 1, the earliest stage in the Taylor AGD gill scoring system from 1–5. The expression of IL-1B has been associated with advance AGD lesions \(^{14}\) which may explain why Il-1B was not found to be up-regulated in the current study. The interleukins IL-1\(^\beta\) and IL-18 promote Th1 and Th17 responses \(^{33}\).

The IL-17 family (IL-17A-F) signals through their correspondent receptors and activate downstream pathways that include NF-\(\kappa\)B, MAPKs and C/EBPs to induce the expression of antimicrobial peptides, cytokines and chemokines \(^{34}\). IL-17, is a key cytokine produced by Th17 cells and is involved in the inflammatory and neutrophil response. A recent study \(^{10}\) reported the expression of IL-17A/F1b and IL-17D to be significantly down-regulated in comparison to the negative control in gills from fish inoculated with a high concentration of \(P.\ perurans\) trophozoites (5000 amoeba /L). In the current study fish were
inoculated with 2750 amoeba/L, and IL-17F was down-regulated at all 4 time points. IL-17F is mainly involved in mucosal host defence mechanisms\(^{35}\).

IL-27\(^\beta\), was the only interleukin significantly induced at 7 dpi. IL-27 is composed of two non-covalently linked subunits, IL-27p28 (p28) and IL-27\(^\beta\), also called Epstein-Barr-virus-induced molecule 3 (EBI3)\(^{36}\). These subunits exhibit structural and sequence homology to IL-12 subunits and IL-6\(^{37}\). IL-27 is unique in that although it induces Th1 differentiation, the same cytokine suppresses immune responses. IL-27 can antagonise the development of the Th17-cell response and limit Th-17 driven inflammation\(^{38}\) which are critical for host defence against bacterial, fungal and viral infections at mucosal surfaces\(^{39}\).

Of the down-regulated immune genes in reactome R-DRE-168256 at 7 dpi (Table 4), 8 genes were found to participate in the Nod-like receptor signalling pathway. Nod-like receptors (NLRs) can initiate or regulate host defence pathways through formation of signalling platforms that subsequently trigger the activation of inflammatory caspases and NF-kB\(^{40}\). Genes found to participate in the Nod-like receptor signalling pathway included cxcl8, hsp90ab1, ikbkb, irf9, mapk3, nlrx1, sugt1, and tbk1. Nod-like receptors (NLRs) sense pathogen-associated molecular patterns (PAMPs) (pathogens/foreign) or damage-associated molecular patterns (cells/self)\(^{41}\). Lrx1 is a dsRNA receptor\(^{42}\) and was identified as being down-regulated in our data set at all 4 time points (4, 7, 14 and 16 dpi).

Genes were also identified with involvement in B-cell receptor (BCR) signalling (ikbkb, mapk3, sky), T-cell receptor (TCR) signalling (dlg1, ikbkb, mapk3) and in the differentiation of Th1/Th2 (ikbkb, jak3, mapk3, stat4) and Th17 cells (hsp90ab1, ikbkb, jak3, mapk3).

Interestingly, all of these signalling pathways had 2 genes in common: inhibitor of nuclear factor kappa B kinase subunit beta (ikbkb) and mitogen-activated protein kinase 3 (mapk3).

The transcription factor Nuclear Factor-kappa beta (NF-\(\kappa\beta\), prior to activation, is held in the cytoplasm by the attachment of inhibitor kappa beta (Ik\(\beta\)) and the formation of the Ik\(\beta\)/NF-\(\kappa\beta\) complex.

Ikbkb is a gene which encodes the enzyme Ik\(\beta\) kinase beta (IKKB) which can phosphorylate 2 serine residues on the inhibitor in the Ik\(\beta\)/NF-\(\kappa\beta\) complex, leading to the dissociation and degradation of the Ik\(\beta\) inhibitor and the subsequent activation of NF-\(\kappa\beta\)\(^{43}\). The dissociated NF-\(\kappa\beta\) can translocate into the nucleus and activate the transcription of hundreds of genes involved in immune response, growth control, or protection against apoptosis. IKKB is critical for cytokine production via NF-kB activation. The initial innate immune response is under the control of IKKB and culminates in a successful humoral response that is dependent on IKK\(\alpha\)\(^{44}\). Mapk3 acts upstream of IKKB in the canonical NF-\(\kappa\beta\) activation pathway\(^{45}\).

This is the first study exploring the transcriptomic response of Atlantic salmon to \(P.\) perurans in a controlled environment. The data presented show that the host response of Atlantic salmon is activated in advance of any clinical symptoms developing on gill tissue of fish inoculated with \(P.\) perurans. Of particular interest is the immune suppression brought about through the downregulation of 2 key genes,
ikbkb and mapk3/ERK1 resulting in the continued inhibition of NF-κβ by Iκβ in the cytoplasm. Pathogens have previously been reported to developed strategies to circumvent the activation of the NF-κβ activation, by preventing the inhibitor, Iκβ, from being ubiquitinated and therefore preventing its degradation, causing NF-κβ to remain sequestered in the cell cytoplasm and therefore inactive\textsuperscript{46}. Indeed, some viruses encode virulence factors, for example vaccinia viral protein B14, that target IKKβ to inhibit NF-κβ-mediated antiviral immune response\textsuperscript{47}, suggesting that virulence factors associated with \textit{P. perurans} potentially have some of immunomodulatory effect on their host. The present study provides the initial discovery and description of genes showing differential expression during early-phase ADG exposure/infection, which provides the basis for future, more in-depth studies of AGD-related immune response pathways in Atlantic salmon.

**Methods**

**Fish husbandry**

Post-smolt Atlantic salmon (70g) were distributed into 4 circular black 1000 L tanks connected to recirculating aquaculture systems at a stocking density of 3.6 kg m\textsuperscript{-3}, water temperature 12°C, artificial seawater (30PSU), and 14h/8h light/dark cycle. Fish were fed a commercial salmon diet at 1% body weight per day. The \textit{in-vivo} fish trial was carried out according to the ARRIVE guidelines for animal research\textsuperscript{48}. This project was authorised by the Health Products Regulatory Authority (HPRA), authorisation number AE19137/P001, in compliance with Directive 2010/ 63/EU transposed into Irish law by S.I. No 543 of 2012.

\textit{Paramoeba perurans} isolation and culture

\textit{P. perurans} trophozoites were collected by gill swabbing from AGD infected Atlantic salmon on a commercial Irish farm. Amoebae were cultured on marine yeast agar plates (MYA; 0.01% malt, 0.01% yeast, 2% Bacto Agar), 16°C overlaid with 7 ml sterile seawater\textsuperscript{49}, and sub-cultured weekly by transferring free-floating cells to fresh MYA plates. Confirmation of \textit{P. perurans} identity was performed using qPCR as previously described by Downes (2015)\textsuperscript{5}.

\textit{Paramoeba perurans} challenge

After an acclimatisation period of 6 days, 90 fish (45 x 2) were challenged with \textit{P. perurans} (2750 amoebae/L) in 300 L for 4 h with oxygen saturation, fish behaviour and welfare closely monitored. Controls, 90 fish (45 x 2) were also held at 300 L for 4 h. Following holding in challenge or control tanks, fish were placed into new tanks containing 1000 L seawater.

**Sample collection**

Fish were euthanised by overdose of anaesthetic (400 mg L\textsuperscript{-1} tricaine methane sulfonate).
Gill tissues, following perfusion with phosphate buffered saline (1x PBS), were collected from six fish at 0 dpi (pre-ADG challenge), 4, 7, 14 and 16 dpi. The 2nd left gill arch was collected from individual fish, the arch cartilage, immersed in RNAlater® (Ambion Inc, Austin, Texas), stored at 4°C overnight, followed by final storage at -80°C.

**Disease progression**

Disease progression was monitored using gill scoring carried out on euthanised fish (Taylor, 2009). Whole gill samples were taken for qPCR and histology to confirm the presence of *P. perurans*. For histological analyses, samples were fixed in 10% neutral buffered formalin, routinely processed, embedded in paraffin wax blocks, sectioned (3–5 µm), and stained with haematoxylin and eosin (H&E).

**Total RNA extraction**

Total RNA was extracted from 30mg gill tissue using the RNeasy® Plus Mini Kit (Qiagen) according to manufacturer’s instructions. RNA was quantified using the Qubit® RNA Assay Kit in Qubit® 2.0 Flurometer (Life Technologies, CA, USA). RNA integrity (RIN) was assessed using the RNA Nano 6000 Assay Kit of the Bioanalyzer 2100 system (Agilent Technologies, CA, USA). Total RNA with RIN ≥ 8.0 or higher were used for RNA-seq.

**Library construction and transcriptome sequencing**

Total RNA from 6 individual fish from at each of the 5 time points; 0, 4, 7, 14, and 16 dpi was used for the construction of 5 sequencing libraries generated using NEBNext® Ultra™ RNA Library Prep Kit for Illumina® (NEB, USA) according to manufacturer’s instructions. Index codes were added to attribute sequences to each sample. The clustering of the index-coded samples was performed on a cBot Cluster Generation System using HiSeq PE Cluster Kit cBot-HS (Illumina) according to the manufacturer’s instructions. After cluster generation, library preparations were sequenced on an Illumina Hiseq platform and 125 bp/150 bp paired-end reads were generated. FastqQc was utilised for quality assessment of reads from each sample (Version 0.11.8) and Multiqc (Version 1.7) was used to visualise all FastQc results. Trimmomatic (v0.36) was used to trim paired reads in FASTQ files, using default parameters for paired-end mode and a minimum read length of 50bp.

Read mapping to the Atlantic salmon (*Salmo salar*) reference genome

The *Salmo salar* genome (https://www.ncbi.nlm.nih.gov/assembly/GCF_000233375.1) was used to map the reads (supplementary Table S1 online). Mapping was implemented using HiSat2 (version 2.1.0) using default parameters and paired-end mode. Counts were generated using featureCounts (v1.6.0) using the default parameters for paired-end reads. RNA-seq specific QC, sample correlation and visualisation were implemented using Seqmonk (Version 1.45.1). Correlation matrices, PCA plots, t-SNE plots and similarity trees were generated to visualise the relationships between all samples, as well as samples by time point.

**Differential expression analysis**
Differential expression analysis was performed on biological replicates (n = 6) from 4, 7, 14 and 16 dpi using the DESeq2 (Version 1.24.0) where each time point was compared to time 0, pre-AGD samples (Fig. 1). DESeq R provide statistical routines for determining differential expression in digital gene expression data using a model based on the negative binomial distribution. The resulting P-values were adjusted using the Benjamin and Hochberg’s approach for controlling the false discovery rate. Genes with an adjusted p-value (FDR adjusted) < 0.05 found by DESeq R were assigned as differentially expressed.

Eight gene lists were generated: up- and down-regulated genes at 4, 7, 14 and 16 dpi.

**Protein-Protein Interaction (PPI)**

PPI was carried out using STRING (version 11.0, https://string-db.org) on the 8 gene lists generated from the pairwise comparisons (4, 7, 14 and 16 dpi, up- and down-regulated genes) using the default settings and *Danio rerio* selected as the species of interest. Significant pathway enrichments and functional information for genes identified in the Immune System (R-DRE-168256) was carried out using STRING against the *Homo sapiens* database.

**Pathway and Process Enrichment**

Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway mapping was performed using Metascape (metascape.org). The zebrafish (*Danio rerio*) database was used to determine GO enrichment. Pathway and process enrichment analysis was performed using the following ontology sources: KEGG Functional Sets, KEGG Pathway, GO Biological Processes, GO Cellular Components, GO Molecular Functions, KEGG Structural Complexes and Reactome Gene Sets.

A user-supplied list of 5824 genes was used as the enrichment background. Terms with a p-value < 0.01, a minimum count of 3, and an enrichment factor > 1.5 are collected and grouped into clusters based on their membership similarities.

**Validation of RNA-seq data using Real-time PCR**

Real-time PCR validation of the RNA-seq data from 6 individual fish at each time point was performed using 48.48 Dynamic Array™ Integrated Fluidic Circuit (IFC) chips on the Biomark HD system. Ten genes identified as being differentially regulated in the gill RNA-seq data were selected for qPCR validation, including cathelicidin 1 (*cath-1*), c-type lectin receptor A (*clr-a*), C-C motif chemokine 4 (*cxc4*), immunoresponsive 1 homolog/ (irg1/acod1), interleukin-8 (*il-8*), interleukin-17F (*il-17*), leukocyte cell-derived chemotaxin-2 (*lect2*), nitric oxide synthase 2, inducible (*nos2*), pentraxin-related protein (*ptx3*), and serum amyloid P-like (*sap*). Primers were designed using PrimerQuest (Integrated DNA Technologies, https://eu.idtdna.com/) (supplementary Table S2 online).

Reverse transcription was carried out using the GoScript (Promega) kit as per manufacturer’s instructions. A pre-amplification step was adopted for the multiplex amplification of the target genes using a MiniAmp Plus PCR machine (Applied Biosystems), as per manufacturer’s recommended protocols (PN 100–5875 C1, Fluidigm). The pre-amplified cDNA was treated with Exonuclease I to remove unincorporated primers prior to running on a Biomark HD, as per manufacturer’s recommended protocols (PN 100–9791 B1,
Fluidigm). PCR amplification efficiency (E) was calculated for each gene of interest and the housekeeping gene by the generation of standard curves using 10-fold serial dilutions of the cDNA template (standard curve: $R^2 > 0.980$, amplification efficiency range 90–105%). Melt curve analysis was used to confirm the amplification of single, PCR products.

**Statistical analysis**

For pathway and process enrichment analyses, p-values < 0.01 were calculated based on the accumulative hypergeometric distribution, and q-values calculated using the Benjamin-Hochberg correction for multiple testing using Metascape (metascape.org). Kappa scores were used as the similarity metric when performing hierarchical clustering on the enriched terms, and sub-trees with a similarity of > 0.3 were considered a cluster. The most statistically significant term within a cluster was chosen to represent the cluster. For qPCR analysis, the fold change of each gene at each time point was analysed relative to the T0 control using an un-paired t-test with differences considered significant at $p < 0.05$. Ggplot2_3.2.1 in R studio Version 3.5.1 was used to generate the Spearman correlation data. Plotly.py version 4.0.0 in Python 3.7.3 was used to graph the RNA-seq and qPCR data validation data.

**Declarations**

**Data Availability Statement**

All data supporting this study are included in the results section, the supplementary material section or openly available in public databases.

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**Competing interests**

The author(s) declare no competing interests.

**Author contributions**

E.M., I.O.C. L.M., J.C. conceived the project.

A.T. and L.M. contributed to the design of the experiments.

A.T., G.M., L.G., L.P., performed the experiments, contributed to the collection and analysis of data.

A.T., L.G., L.P. conducted statistical analysis.

A.T. and E.M. wrote the paper. All authors edited the paper.
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Figure 4

(A): Enriched GO terms in the gill at 4 dpi for up-regulated DEGs. (B): Enriched GO terms in the gill at 4 dpi for down-regulated DEGs.