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Convergent Evolution of Mucosal Immune Responses at the Buccal Cavity of Teleost Fish

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SUMMARY
The buccal mucosa (BM) is a critical first line of defense in terrestrial animals. To gain further insights into the evolutionary origins and primordial roles of BM in teleosts here we show that rainbow trout, a teleost fish, contains a diffuse mucosal associated lymphoid tissue (MALT) within its buccal cavity. Upon parasite infection, a fish immunoglobulin specialized in mucosal immunity (sIgT) was induced to a high degree, and parasite-specific sIgT responses were mainly detected in the buccal mucus. Moreover, we show that the trout buccal microbiota is prevalently coated with sIgT. Overall our findings revealed that the MALT is present in the BM of a non-tetrapod species. As fish IgT and mucus-producing cells are evolutionarily unrelated to mammalian IgA and salivary glands, respectively, our findings indicate that mucosal immune responses in the BM of teleost fish and tetrapods evolved through a process of convergent evolution.

INTRODUCTION
The buccal cavity (BC) of vertebrates is the gateway for both the gastrointestinal and respiratory tracts and is considered a critical mucosal surface in tetrapod species (Winning and Townsend, 2000; Squier and Kremer, 2001; Abbate et al., 2006). Microbes from air, water, and food pose continuous challenges to the homeostasis of the BC (Walker, 2004), and thus, vertebrates have evolved efficient innate and adaptive immune strategies to protect this critical surface. In tetrapod species, secretory IgA (sIgA) is the main humoral component involved in adaptive immune responses against oral pathogens (Brandtzaeg, 2013). Moreover, orally produced sIgA is also involved in the control and homeostasis of the buccal microbiota. sIgA is the most abundant immunoglobulin class in the saliva of mammals (Marotte and Lavoie, 1998). It is worth noting that saliva is produced only by mammals, birds, and reptiles, and it is in mammals where it is known to have a very important digestive function (Pedersen et al., 2002; Dawes et al., 2015), whereas in birds and reptiles this role is significantly less marked. Interestingly, amphibians are known to contain both mucus-producing cells as well as intermaxillary salivary glands (Latney and Clayton, 2014), although the digestive and adaptive immune roles of their putative saliva and mucosal secretions have been ill investigated. In contrast to all tetrapod species, teleost fish lack salivary glands in their BC, which is instead populated with abundant mucus-secreting cells that produce the mucus that coats their buccal epithelium (Yashpal et al., 2007).

In mammals, some mucosal regions within the BC are covered by a keratinized stratified epithelium (gingival, hard palate, outer lips), whereas other areas, including the ventral side of the tongue, the floor of the mouth, the inner surface of the lips, and cheeks, are lined by a non-keratinized stratified epithelium (Squier and Kremer, 2001). In contrast, the entire buccal epithelium of fish is non-keratinized. Interestingly, the non-keratinized buccal areas of mammals resemble the overall structure of the fish buccal mucosa (BM) as both contain two main layers, an outer layer of stratified squamous epithelium and an underlying layer of dense connective tissue (lamina propria) (Winning and Townsend, 2000; Squier and Kremer, 2001; Abbate et al., 2006). Mammalian sIgA found in the saliva is produced by plasma cells (PCs) localized around the salivary glands (Brandtzaeg, 2013). Upon secretion by PCs, sIgA is actively transported by the polymeric immunoglobulin receptor (pIgR) expressed by parenchymal cells within these glands (Carpenter et al., 2004; Brandtzaeg, 2013). In mammals, the salivary gland within BM is considered a mucosal effector site where IgA-producing plasma cells are derived from mucosal inductive sites localized in the...
nasopharynx-associated lymphoid tissue and gut-associated lymphoid tissue (i.e., tonsils and Peyer patches) (Jackson et al., 1981; Brandtzaeg, 2007). Whether non-tetrapod species have evolved mucosal adaptive immune responses in the BM is at this point unknown. Since many aquatic environments harbor much higher concentrations of microbes than that found in air, it is reasonable to hypothesize that fish must have evolved an effective mucosal immune system to protect their BC.

Within non-tetrapods, bony and cartilaginous fish represent the earliest vertebrates containing immunoglobulin (Ig). In contrast to mammals that contain five major Ig classes, only three Ig isotypes have been identified in teleosts, IgM, IgD, and IgT/IgZ. IgM is the best characterized teleost Ig isotype both at the molecular and functional levels, and it is the most abundant Ig class in plasma (Salinas et al., 2011). Moreover, IgM represents the prevalent Ig in systemic immune responses (Salinas et al., 2011). Like IgM, IgD is an ancient Ig class that has been found in most jawed vertebrates (Ramirez-Gomez et al., 2012). However, the immune function of fish IgD remains unknown, although secreted IgD has been found coating a small portion of the fish microbiota (Xu et al., 2016) and may function as an innate pattern recognition molecule (Edholm et al., 2010). IgT (also called IgZ in some species) has been described in all studied teleost fish except for medaka and catfish (Danilova et al., 2005; Hansen et al., 2005; Fillatreau et al., 2013). We have previously shown that IgT plays a major role in teleost mucosal immunity, akin to that of IgA tetrapods (Zhang et al., 2010; Ramsey et al., 2010). This discovery broke the old paradigm that mucosal immunoglobulins were present only in tetrapod species. More specifically, we have demonstrated that, upon infection, IgT is the main Ig induced in several mucosal surfaces, including the gut, gills, nose, and skin (Zhang et al., 2010; Xu et al., 2013, 2016; Tacchi et al., 2014; Yu et al., 2018). Significantly, we also found that similar to the role of slgA in mammals, slgT is the prevalent Ig coating the microbiota in all fish mucosal areas (Zhang et al., 2010; Tacchi et al., 2014; Xu et al., 2013, 2016).

To gain further insights into the evolutionary origins and primordial roles of buccal adaptive immune responses in vertebrates, here, we investigated the presence and immune roles of a buccal MALT in the BC of a teleost fish, the rainbow trout (Oncorhynchus mykiss). Our findings reveal a well-defined diffuse MALT in the trout’s BC, and we demonstrate its key role in inducing strong local innate and adaptive immune responses upon infection with Ichthyophthirius multifiliis (Ich) parasite. Furthermore, we show that, in addition to being the prevalent local Ig induced upon infection, slgT is also the main slg recognizing and coating the trout buccal microbiota. Overall, our findings indicate the presence of a bona fide MALT in the BC of a non-tetrapod species as well as its involvement in both the control of pathogens and recognition of microbiota.

RESULTS

Teleost BM Shares the Typical Features of a MALT
To understand the histological organization of teleost BM (Figures S1A–S1D), paraffin sections of BMs obtained from five different families (Figure S2), Salmonidae, Percichthyidae, Synbranchidae, Siluridae, and Channidae, were stained with both hematoxylin and eosin (H&E) (Figures 1A–1E) and Alcian blue (AB) (Figures 1F and S3A–S3D). We observed that the BM of Japanese sea bass (Lateolabrax japonicus), Asian swamp eel (Monopterus albus), Southern catfish (Silurus meridionalis), and Snakehead (Channa argus) contained intraepithelial lymphocytes and lamina propria leukocytes (Figures 1A–1E), with a large number of mucus-producing cells in the buccal epithelium (Figures 1F and S3A–S3D). These results of distribution and structure in the BMs from all five species resemble those of other mucosal tissues. Moreover, using reverse transcription quantitative real-time PCR (qPCR), we measured the levels of expression of gene markers for the main myeloid and lymphoid cell types in the BM from the control adult rainbow trout. We then compared them to those in the head kidney, skin, and muscle. We found that consistently high levels of expression of most immune markers were detected in the BM, head kidney, and skin, indicating an unrecognized immunological function of the BM in rainbow trout (Figure 1G). The abundance of two main B cell subsets (IgM+ and IgT+ B cells) in the BM was analyzed by flow cytometry (Figure 1H). We found that, similar to the gut, skin, gills, and nose (Zhang et al., 2010; Xu et al., 2013, 2016; Tacchi et al., 2014), IgT+ B cells make up ~52.53% of the total B cells in the BM of rainbow trout, whereas ~47.47% of the total B cells are IgM+ (Figure 1I). In contrast, only ~29.24% of IgT+ B cells were detected in the head kidney (Figure 1I). Mucosal Igs have been previously reported to be transported across the mucosal epithelium via polymeric Ig receptors (pIgRs). Here, using a polyclonal anti-trout pIgR antibody, a large portion of the epithelial cells of the BM were stained by immunofluorescence and found to be located in the apical areas of the mucosal epithelium of the trout (Figure 1J; isotype-matched control antibodies,
Figure 1. General Organization of Teleost BM

(A–E) Hematoxylin and eosin stains of BMs obtained from five different teleost families, including rainbow trout (Oncorhynchus mykiss) (A), Japanese sea bass (Lateolabrax japonicus) (B), Asian swamp eel (Monopterus albus) (C), Southern catfish (Silurus meridionalis) (D) and Snakehead (Channa argus) (E). BC, buccal cavity; BE, buccal epithelium; LP, lamina propria; BS, buccal submucosa; CL, cartilage layer. Scale bar, 50 μm.

(F) AB stain of the BM of a control adult rainbow trout (Oncorhynchus mykiss). Black triangles indicate lymphocytes. Red arrows indicate mucous cells. The red asterisk denotes taste bud in BM. BC, buccal cavity; BE, buccal epithelium; LP, lamina propria. Scale bar, 50 μm.

(G) Heatmap illustrates results from quantitative real-time PCR of mRNAs for selected immune markers in trout head kidney, BM, skin, and muscle (n = 6). Data are expressed as mean Ct values ± SEM.

(H) Flow cytometry analysis of head kidney (Left) and BM (Right) leukocytes stained with anti-IgM and anti-IgT antibodies. Numbers in outlined boxes indicate the percentage of IgM⁺ (Top Left) and IgT⁺ (Bottom Right) B cells in the lymphocyte gate, respectively.

(I) Frequency (Mean ± SEM) of IgM⁺ and IgT⁺ B cells among total B cells present in trout head kidney and BM (n = 12).

(J) Immunofluorescence staining for pIgR (green) in a paraffinic section of trout BM (n = 9). Nuclei are stained with DAPI (blue) (isotype-matched control antibody staining is shown in Figure S4A). Scale bar, 50 μm.

(K and L) Immunoblot and densitometric analysis of the concentration of IgT, IgM, and IgD in buccal mucus (K) and serum (L) (n = 12).

(M and N) Ratio of IgT to IgM concentration (M) and IgD to IgM concentration (N) in buccal mucus and serum (n = 12).

Data in K–N are representative of at least three independent experiments (Mean ± SEM).
After filtering by the Oncorhynchus mykiss genes upregulated and 2,997 and 998 genes downregulated at days 14 and 28, respectively (Figure 3C). A total of 2232 and 1393 pressed genes (DEGs) were identified as immune-related genes, as shown in the histogram (Figure 3D). To further investigate the DEGs of the BM that were involved in responding to Ich infection among the four groups, KEGG pathway analysis was conducted. Interestingly, we found that pathways associated with immune response, signal molecules, infectious disease, and metabolism were all overrepresented in the differentially expressed set of genes (Tables S2 and S3). Importantly, we identified a significant modification in the expression of genes (Figure S7) involved in innate immunity (Figure 3E, left; Table S4) and adaptive immunity (Figure 3E, right; Table S4) on both days 14 and 28 following Ich infection, with 2232 and 1393 genes upregulated and 2,997 and 998 genes downregulated at days 14 and 28, respectively (Figure 3C). To understand whether the different trout immunoglobulins were in monomeric or polymeric form in the buccal mucus, we collected and processed buccal mucus of rainbow trout and loaded it into a gel filtration column. A large portion of IgT in the buccal mucus was found in polymeric form, as it eluted at a fraction similar to that of trout IgM, a tetrameric Ig (Figure S5A). In contrast, a small portion of IgT in the buccal mucus was eluted in monomeric form. Interestingly, by SDS-PAGE under non-reducing conditions, polymeric IgT (pIgT) in buccal mucus migrated in the same position as monomeric IgT, indicating that pIgT subunits are associated by non-covalent interactions (Figure S5B, left). Moreover, immunoblot analysis further confirmed the presence of IgT, IgM, or IgD in these bacteria (Figure 2D). Interestingly, similar to the results previously reported for trout skin microbiota, we found that more than 50% of total IgT present in the buccal mucus was found coating bacteria, whereas only ~20% of IgM and ~17% of IgD was being used for bacterial coating (Figure 2E).
Response and Proliferation of B cells in Trout BM after Ich Parasite Infection

Using immunofluorescence microscopy, we observed few IgT+ and IgM+ B cells in the buccal epithelium of control fish (Figure 4A, left; isotype-matched control antibodies, Figure S4B). Interestingly, a moderate increase in the number of IgT+ B cells was observed in the buccal epithelium of trout from the infected group (28 dpi) (Figure 4A, middle). Notably, a large number of IgT+ B cells accumulated in the buccal epithelium of survivor fish (75 dpi) when compared with those of control fish (Figure 4A, right). Cell counts of the stained sections described in Figure 4A showed that the IgT+ B cells increased ~3-fold and ~8-fold in the infected and survivor fish, respectively (Figure 4B). However, the abundance of IgM+ B cells did not change significantly in the infected and survivor fish when compared with the controls (Figures 4A and 4B).

Next, we investigated whether the increase of IgT+ B cells observed in the BM of survivor fish was derived from the process of local IgT+ B cell proliferation or due to an infiltration of these cells from systemic lymphoid organs. To do so, we measured the in vivo proliferative responses of IgT+ and IgM+ B cells stained with 5-Ethynyl-2'-deoxyuridine (EdU), a thymidine analogue that incorporates into DNA during cell division (Salic and Mitchison, 2008). By immunofluorescence microscopy analysis, we observed a significant increase in the proliferation of EdU+ IgT+ B cells in survivor fish (~6.35 ± 0.34%) when compared...
with that of the control fish ($\sim 2.62 \pm 0.06\%$) (Figures 4C–4E). However, no difference was detected in the percentage of EdU$^+$ IgM$^+$ B cells between control fish and survivor fish (Figures 4C–4E). Similarly, by flow cytometry, we found a significant increase in the percentage of EdU$^+$ IgT$^+$ B cells in the BM of survivor fish ($\sim 12.11 \pm 1.32\%$ in total IgT$^+$ B cells) when compared with that of control fish ($\sim 4.01 \pm 0.34\%$ in total IgT$^+$ B cells) (Figures 4F–4H). On the contrary, we did not observe any significant difference in the percentage of EdU$^+$ IgM$^+$ B cells between control and survivor fish (Figures 4F–4H). Interestingly, a large increase in the percentage of EdU$^+$ IgM$^+$ B cells in the head kidney of survivor fish were detected when compared with that of control fish, whereas the percentage of EdU$^+$ IgT$^+$ B cells did not show a significant difference between the two groups (Figures 4I–4K).

**Ig Responses in Trout BM after Ich Parasite Infection**

To investigate whether parasite-specific Igs were produced in trout after Ich parasite challenge, we measured the Igs concentration and the capacity of Igs from buccal mucus and serum to bind to the
Figure 4. Increases and Proliferative Responses of IgT+ B cells in the BM of Trout Infected with Ich

(A) Two different DIC images of immunofluorescence staining on paraffinic sections of BM from uninfected control fish (left), 28 days infected fish (middle), survivor fish (right), and enlarged images of the areas outlined, stained for IgT (green) and IgM (red); nuclei are stained with DAPI (blue). BC, buccal cavity; BE, buccal epithelium; LP, lamina propria. Scale bar, 20 μm. Data are representative of at least three different independent experiments (n = 12 per group).

(B) IgT+ and IgM+ B cells in paraffinic sections of BM from uninfected control fish, infected fish, and survivor fish (n = 12), counted in 25 fields (original magnification, ×40).

(C and D) Immunofluorescence analysis of EdU incorporation by IgT+ or IgM+ B cells in the BM of control (C) and survivor fish (D). Paraffinic sections of BM were stained for EdU (magenta), trout IgT (green), trout IgM (red), and nuclei (blue) detection (n = 9 per group). White arrowheads point to cells double stained for EdU and IgT. BC, buccal cavity; BE, buccal epithelium; LP, lamina propria. Scale bars, 20 μm.

(E) Percentage of EdU+ cells from the total BM IgT+ or IgM+ B cell populations in control or survivor fish counted from C and D.

(F and G) Representative flow cytometry dot plot showing proliferation of IgT+ B cells (F) and IgM+ B cells (G) in BM leukocytes of control and survivor fish (n = 12 per group).

(H) Percentage of EdU+ cells from the total BM IgT+ or IgM+ B cell populations in control or survivor fish (n = 12). (I and J) Representative flow cytometry dot plot showing proliferation of IgT+ B cells (I) and IgM+ B cells (J) in head kidney leukocytes of control and survivor fish (n = 12 per group). The percentage of lymphocytes representing proliferative B cells (EdU+) is shown in each dot plot.

(K) Percentage of EdU+ cells from the total head kidney IgT+ or IgM+ B cell populations in control or survivor fish (n = 12 per group).

*p < 0.05, ***p < 0.001 (one-way ANOVA with Bonferroni correction). Data in B, E, H, and K are representative of at least three independent experiments (Mean ± SEM).
parasite. Immunoblot analysis showed that the IgT concentration in the buccal mucus from infected and survivor fish increased by ~2- and ~8-fold, respectively, when compared with control fish, whereas the concentration of IgM and IgD did not change significantly in any fish groups (Figure 5A). In contrast, only ~2- and 3-fold increases of serum IgT concentration were observed in infected and survivor fish, respectively, whereas the concentration of serum IgM increased by ~5-fold in both the infected and survivor groups when compared with control fish (Figure 5E). However, in both infected and survivor fish, the concentration of IgD did not change significantly in either the buccal mucus or serum (Figures 5A and 5E). By a pull-down assay, we found a significant increase in parasite-specific IgT binding in up to 1/40 diluted buccal mucus of infected and survivor fish, in which we detected ~3.8-fold and ~3.7-fold binding increases, respectively, when compared with that of the control fish (Figures 5B–5D). Conversely, in serum, parasite-specific IgT binding was detected only in 1/10 dilution in the same Figure 5. Immunoglobulin Responses in the Buccal Mucus and Serum from Infected and Survivor Fish

(A) Concentration of IgT, IgM, and IgD in buccal mucus of control, infected, and survivor fish (n = 12 per group).
(B) Immunoblot analysis of IgT-, IgM-, and IgD-specific binding to Ich in buccal mucus (dilution 1:2) from infected and survivor fish.
(C and D) IgT-, IgM-, and IgD-specific binding to Ich in dilutions of buccal mucus from infected (C) and survivor (D) fish, evaluated by densitometric analysis of immunoblots and presented as relative values to those of control fish (n = 9 per group).
(E) Concentration of IgT, IgM, and IgD in serum of control, infected, and survivor fish (n = 12 per group).
(F) Immunoblot analysis of IgT-, IgM-, and IgD-specific binding to Ich in serum (dilution 1:10) from infected and survivor fish (n = 12 per group).
(G and H) IgT-, IgM-, and IgD-specific binding to Ich in dilutions of serum from infected (G) and survivor (H) fish, evaluated by densitometric analysis of immunoblots and presented as relative values to those of control fish (n = 9 per group).
*p < 0.05, **p < 0.01, and ***p < 0.001 (unpaired Student’s t test). Data in A, C, D, E, G, and H are representative of at least three independent experiments (Mean ± SEM).

The substantial increase of proliferating IgT+ B cells and high parasite-specific IgT responses occurred in the BM of survivor fish, suggesting that specific IgT might be locally generated in the BM of trout. To further test this hypothesis, we measured the parasite-specific Igs titers from the medium of cultured BM, head kidney, and spleen explants from control and survivor fish (Figure S8). Importantly, we found a significant increase in parasite-specific IgT binding in up to 1/10 diluted medium (~3.6-fold) of cultured BM explants of survivor fish, whereas parasite-specific IgM binding was observed only at the 1/2 dilution in the same
medium (Figures S8A and S8D). In contrast, predominant parasite-specific IgM binding was observed in up to 1/40 dilutions in the medium of head kidney and spleen explants, and parasite-specific IgT binding was detected in up to 1/10 dilutions in the same medium (Figures S8B, S8C, S8E, and S8F). Interestingly, negligible parasite-specific IgD binding was detected in the medium of cultured BM, head kidney, and spleen explants from survivor fish (Figures S8A–S8F).

The high expression of local parasite-specific IgT and large increases in the number of IgT+ B cells in the BM of trout after Ich parasite challenge led us to hypothesize a dominant role of IgT in buccal immunity. At 28 dpi, infected fish showed small white dots on the buccal surface, and using immunofluorescence microscopy, Ich trophonts were easily detected in the buccal epithelium of these fish using an anti-Ich antibody (Figures 6A and 6B; prebleed control antibodies, Figure S4C). Interestingly, we found that most parasites in the BM were intensely stained with IgT, whereas only some parasites were slightly recognized by IgM and nearly no parasites were coated with IgD (Figures 6A and 6B). Interestingly, we found that the levels of IgT coating on Ich parasites located inside of the buccal epithelium differed from those on the surface of the buccal epithelium (Figures 6C and 6D). The lower percentage of low (12%), medium (8%), and high levels (2%) of IgT coating on Ich parasites within the buccal epithelium than those (low, 27%; medium, 33%; high, 26%) located on the surface of the buccal epithelium (Figure 6D), respectively, suggests that IgT plays a key role in forcing the Ich parasite to exit from the epithelium of trout (Figure 6E).

plgR in Trout BM

In mammals, sIgA can be transepithelially transported by plgR into the BM (Brandtzaeg, 2013). In trout, we have previously reported that tSC, the secretory component of trout plgR (tpIgR), is associated with sIgT in the gut, gills, skin, and nose (Zhang et al., 2010; Xu et al., 2013, 2016; Tacchi et al., 2014). Importantly, using immunofluorescence microscopy, most tpIgR-containing cells were observed in the buccal epithelium layer of adult control rainbow trout using anti-tpIgR antibody (Figure 1I). Therefore, we hypothesized that tSC plays a key role in the transport of sIgT into the BM of trout. By immunoblot analysis, we detected tSC in the buccal mucus but not in the serum (Figure S9A). To determine whether buccal mucus sIgT was associated with tSC, using antibodies against tSC (trout plgR) and IgT, we carried out co-immunoprecipitation assays in buccal mucus from survivor fish. Our results showed that antibodies against trout IgT were able to co-immunoprecipitate tSC in the buccal mucus (Figure S9B). Moreover, sIgT in the buccal mucus could also be immunoprecipitated by anti-plgR antibody (Figure S9C). Using immunofluorescence microscopy, we observed that most plgR-containing cells were located in the buccal epithelium of control trout, as shown also in Figure 1I. Critically, some of those plgR-containing cells were also positively stained with anti-IgT antibody, thus supporting further a role of plgR in the transport of sIgT into the BM (Figure S9D).

**DISCUSSION**

Mucosal immunoglobulins (slgs), especially slgA responses in the saliva of the BM of mammals have been extensively reported (Brandtzaeg, 2007, 2013). However, nothing is known with regards to the evolution and roles of slgs and B cells at the BM of early vertebrates. In this study, we first show that the trout BC contains a MALT characterized by an epithelium layer containing a higher percentage of IgT+ B cells than IgM+ B cells, similar to what we have previously reported in the fish gut, skin, gills, and nose (Zhang et al., 2010; Xu et al., 2013, 2016; Tacchi et al., 2014). Importantly, using immunofluorescence microscopy, most tpIgR-containing cells were observed in the buccal epithelium layer of adult control rainbow trout using anti-tpIgR antibody (Figure 1I). Therefore, we hypothesized that tSC plays a key role in the transport of sIgT into the BM of trout. By immunoblot analysis, we detected tSC in the buccal mucus but not in the serum (Figure S9A). To determine whether buccal mucus sIgT was associated with tSC, using antibodies against tSC (trout plgR) and IgT, we carried out co-immunoprecipitation assays in buccal mucus from survivor fish. Our results showed that antibodies against trout IgT were able to co-immunoprecipitate tSC in the buccal mucus (Figure S9B). Moreover, slgT in the buccal mucus could also be immunoprecipitated by anti-plgR antibody (Figure S9C). Using immunofluorescence microscopy, we observed that most plgR-containing cells were located in the buccal epithelium of control trout, as shown also in Figure 1I. Critically, some of those plgR-containing cells were also positively stained with anti-IgT antibody, thus supporting further a role of plgR in the transport of sIgT into the BM (Figure S9D).

In contrast, trout sIgD was in monomeric form in the buccal mucus, similar to the finding of slgA in the saliva from humans (Brandtzaeg, 2013). In contrast, trout sIgD was in monomeric form in the buccal mucus, as previously found in the gills and nose (Xu et al., 2016; Yu et al., 2018). In line with the descriptions of other sources of mucus in teleosts (Xu et al., 2014), all subunits of polymeric slgT in trout buccal mucus were associated by non-covalent interactions. It is worth pointing out that, in agreement with the finding that the ratio of IgA/IgG in saliva is much higher than that in serum in mammals (Brandtzaeg, 2004), we found that the ratio of IgT/IgM in the buccal mucus was 25-fold higher than in the serum. Thus, the predominance of IgT+ B cells and the high IgT/IgM ratio in the trout BM indicate a potential role for sIgT in mucosal buccal immune responses. Moreover, similar to the transport mechanism for slgA in the BM from mammals through the plgR
Figure 6. Parasites Are Predominantly Coated by IgT in the Buccal Epithelium of Infected Trout

(A and B) Four different microscope images of slides showing immunofluorescence staining of Ich parasites in BM paraffinic-sections from trout after 28 days of infection with Ich (n = 6). From left to right: Ich (magenta), IgM (red), and IgT (green) with nuclei stained with DAPI (blue) (A); From left to right: Ich (magenta), IgD (red), and IgT (green) with nuclei stained with DAPI (blue) (B). DIC images showing merged staining (prebleed control and anti-Ich antibodies are shown in Figure S4C). Scale bars, 50 µm.

(C) The different levels of Ich parasites coated by IgT (green) in the inside or surface of BM were divided into four main categories. -, no coating; +, slight level of coating; ++, high level of coating, and ++++, strong level of coating. Buccal epithelium (BE) and lamina propria (LP) are shown. Scale bars, 50 µm.

(D) Percentage of different levels of Ich parasites coated by IgT in the inside (n = 50) or surface (n = 70) of BM.

(E) Proposed model of Ich parasites coated by IgT in the epithelium of BM. Buccal cavity (BC), buccal epithelium (BE), and lamina propria (LP) are shown. Data are representative of at least three different independent experiments.
ever, no studies on IgT+ B cellslocalsegregations have carried out of antiguts and skin mucosal areas. The result is in agreement with previously reported findings in trout gut, skin, gills, and nose microbiota (Zhang et al., 2010; Xu et al., 2013, 2016; Tacchi et al., 2014). Interestingly, it has been reported that salivary slgA predominantly coats the surface of bacterial pathogens, such as Streptococcus mutans, Actinobacillus actinomycetemcomitans, and Porphyromonas gingivalis, which are strongly associated with oral diseases in mammals (Marcotte and Lavoie, 1998; Nogueira et al., 2005; Mikuls et al., 2009). Hence, to gain insight into the role of slgT in the homeostasis of BM, future studies are needed to ascertain the type of buccal microbiota species coated by slgT.

Here, we show a key involvement of buccal slgT in the immune response against Ich, a trout mucosal pathogen. Interestingly, the capacity of Ich to invade the BM of fish had never been appreciated to date. Following Ich infection, the upregulation of both innate and adaptive immune genes was detected in the trout BM, thus showing the involvement of teleost BM in immunity. It is worth noting that we found that B cell markers (i.e., IgT, IgM, CD22) but not T cells markers are significantly upregulated after infection with Ich. This may indicate that B cells but not T cells play a key role against Ich infection. Alternatively, it is possible that T cell responses were absent in the two time points used for transcriptome analysis, although we cannot exclude the possibility that T cells may still be involved in the immune response against Ich.

Moreover, we found a large accumulation of IgT+ but not IgM+ B cells appearing in the buccal epithelium of infected and survivor fish, whereas a few scattered cells could also be observed in the lamina propria. In contrast, slgA-secreting cells are localized for the most part in the lamina propria of salivary glands in mammals (Deslauriers et al., 1985; Brandtzaeg, 2007, 2013). Importantly, these findings are in agreement with the increased concentration of IgT but not IgM or IgD at the protein level in the buccal mucus of the same individual, thus indicating that large increases in the concentration of IgT were produced by the accumulation of IgT+ B cells in the buccal epithelium. Moreover, high parasite-specific IgT titers were detected in buccal mucus, whereas predominant parasite-specific IgM responses were particularly detected in serum. Thus, our findings in the teleost BM reinforce the notion that IgT and IgM responses are specialized in mucosal and systemic areas, respectively (Zhang et al., 2010; Xu et al., 2013, 2016; Yu et al., 2018). However, previous studies showed that IgT is also involved in immune responses in trout spleen upon systemic viral infection (Castro et al., 2013). Interestingly, we found that the parasite-specific IgT titers in buccal mucus were higher than those previously reported in skin mucus (Xu et al., 2013) but lower than those found in gill and nasal mucus (Xu et al., 2016; Yu et al., 2018), suggesting that the degree of the immune response differs depending on the mucosal surfaces. In addition, in this study we found significant proliferative IgT+ B cell responses in the BM of trout, similar to what we have previously reported in the fish gill and nose (Xu et al., 2016; Yu et al., 2018). These results suggest that the accumulation of IgT+ B cells in these mucosal surfaces after infection is due to local proliferation, although this remains to be fully demonstrated. However, no studies on IgT+ B cell local proliferation have been carried out so far in gut and skin mucosal areas. Thus, future studies are needed to investigate whether similar IgT+ B cells proliferative responses are locally observed in the skin and gut of trout upon parasite infection. It is clear, however, that important commonalities are observed in the immune responses thus far studied in the gut, skin, gill, nose, and buccal mucosa, all of which are summarized in Figure S10. Thus, our data strongly suggest that the observed parasite-specific IgT responses in the BM were induced locally as we detected significant proliferative responses of IgT+ B cells in the BM upon parasite infection, and supernatants from BM explants of survivor fish contained significant parasite-specific IgT titers. Although these data suggest that IgT-specific responses are induced locally in the BM, at this point we cannot exclude the possibility that many of the BM IgT+ B cells have not proliferated locally and that have instead been transferred through blood circulation into the BM after proliferating elsewhere. Further studies are warranted to analyze this important point. In line with what we found in the gill and nose MALTs (Xu et al., 2016; Yu et al., 2018), our data suggest that the trout BM would act both as inductive and effector site of IgT responses. In contrast, mammalian BC appears to act only as an effector site (Jackson et al., 1981; Brandtzaeg, 2007; Novak et al., 2008). In that
regard, slgA is the predominant Ig isotype in human saliva (Brandtzaeg, 2007, 2013), and a dramatic increase of IgA secretion as well as IgA-positive cells in the salivary gland occurs following infection with pathogens, including the HIV virus (Lu and Jacobson, 2007), the bacteria S. mutans (Colombo et al., 2016), and the parasite toxoplasma gondii (Loyola et al., 2010). Thus, from an evolutionary viewpoint, our findings indicate a conserved role of mucosal Igs (i.e., IgT, IgA) in the control of pathogens at the BM in aquatic and terrestrial vertebrates.

Here we found a larger ratio of high-intensity IgT coating on the Ich parasites located on the surface of BM when compared with that of Ich inside the buccal epithelium. Thus, it is conceivable that the strong parasite-specific IgT responses elicited in the BM after infection force the BM parasites inside the epithelium to exit it (Wang and Dickerson, 2002). In line with this hypothesis, previous studies have demonstrated that passive immunization of channel catfish (Ictalurus punctatus) using mouse monoclonal antibodies specific to Ich immobilization antigens contributes to the parasite clearance or exit from the host (Clark et al., 1996; Clark and Dickerson, 1997). However, we cannot exclude the possibility that rather than occurring inside the BM epithelium, the high coating of the parasite by specific IgT occurs outside of the BM epithelium, by parasite-specific IgT present in the BM mucus (i.e., specific IgT would be generated upon infection of fish by the parasite). If coating occurs via IgT present in the mucus outside the BM epithelium, this IgT could then be involved in the immobilization of the parasite, thus preventing it from invading the epithelium. Moreover, we cannot exclude that factors other than IgT may force the parasite to exit the BM epithelium. For example, complement might be activated by IgT bound to the parasite, which in turn might elicit the exit response, or contribute to such response. Future studies are needed to investigate the specific role of IgT coating as well as other immune factors in forcing the exit of Ich from the BM epithelium, or in preventing its invasion.

In conclusion, our findings show the presence of a previously unrecognized bona fide MALT in the BM of a non-tetrapod species and its involvement in both the control of pathogens and recognition of microbiota. Significantly, these data indicate that mucosal adaptive immune responses evolved both in tetrapod and non-tetrapod species through a process of convergent evolution, as fish IgT and mucus-producing cells are evolutionary unrelated to mammalian IgA and salivary glands, respectively. It is in this aspect that fish and mammals have evolved different fascinating strategies in the way by which their immunoglobulin-containing fluids are produced and secreted in the BM (Figure 7). On the one hand, mammalian slgA produced by lamina propria-plasma cells is transported inside the salivary glands via plgR-expressing parenchymal cells; the slgA-containing saliva within the salivary gland is thereafter secreted into the outer layer of the BM epithelium (Figure 7A). In contrast, fish slgT produced by intraepithelial IgT+ B cells is transported via plgR-expressing epithelial cells into the outer layer of the buccal epithelium where it mixes with mucus derived from mucus-secreting cells (Figure 7B). Thus, different molecules (slgT versus slgA) and cell types/glands (mucus-secreting cells versus salivary glands) of fish and mammals utilize different but functionally analogous strategies to coat the outer layer of the BM epithelium with different slg-containing fluids (mucus versus saliva) with the same goal (the control of pathogens and microbiota). Interestingly, and based on our data, it would appear that the main role of mucus-based buccal fluids in lower vertebrates is immune defense and mucosal homeostasis, whereas throughout evolutionary time, the saliva-based buccal fluids of tetrapods have gained an important role in digestion. Future work is required to further address this attractive. Finally, since we find that slgT responses are locally produced in the fish BM, from a practical level our findings may have important implications for the design of future fish vaccines that stimulate mucosal BM responses.

Limitations of the Study
This study shows that a well-defined diffuse MALT is present in the trout’s BC, which can produce strong innate and adaptive immune responses to the parasitic infection. Moreover, we provide evidence that specific IgT is the main player involved in the buccal adaptive immunity. However, there are limitations to our study due to some experimental constraints. For instance, even though the upregulated expression of B cell and T cell makers in teleost BM reveal that both of them are involved in the buccal immunity against Ich, we did not address the interaction between B cells and CD4-T cells in teleost BM during pathogenic infection, because of the lack of anti-trout CD4 mAb. In addition, this study shows local proliferative IgT+ B cell responses and pathogen-specific IgT production in the BM of a fish species, but we cannot rule out the possibility that, on antigen uptake, loaded BM APCs may migrate into central secondary lymphoid organs (that is, spleen or head kidney) where the resulting activated IgT+ B cells may then home into the BM. Thus, further experiments will be required to address those aspects conclusively.
Figure 7. Simplified Scheme of the Analogous Strategies of Mammals and Fish in the Production and Secretion of Immunoglobulin-containing Fluids in their BM

(A, lower) In mammals, the BM contains numerous salivary glands, which produce and secrete saliva into the salivary layer (SL) via secretory ducts. Localized aggregations of IgA+ plasma cells are commonly found in the lamina propria (LP) of the BM. (A, upper) Mucosal immunoglobulin (sIgA) containing the joining (J) chain is produced by local IgA+ plasma cells in the LP and transported inside salivary gland via plgR also termed as (membrane secretory component [mSC]) expressed basolaterally on parenchymal cells. Thereafter, sIgA mixes with saliva in the salivary gland and the IgA-containing saliva is secreted into the SL through ductal system.

(B, lower) Teleost BM is instead populated with abundant mucus-secreting cells, which produce mucus, which is secreted directly into the mucous layer (ML). IgT-secreting cells are found scattered mainly in the buccal epithelium (BE) where they increase in significant numbers upon infection. (B, upper) Mucosal IgT (sIgT) is secreted by intraepithelial IgT-secreting cells and transported via plgR-expressing epithelial cells directly into the ML where it mixes with mucus derived from mucus-secreting cells. Finally, the sIgT-containing mucus and sIgA-containing saliva from fish and mammals, respectively, preserve BC homeostasis by maintaining the establishment of a healthy microbiota and at the same time, by fighting potential pathogens.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2019.08.034.

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AUTHOR CONTRIBUTIONS
Y.-Y.Y. and Z.X. designed research; Y.-Y.Y. and W.-G.K. performed most of the experiments; H.-Y.X., X.-T.Z., S.D., and F.D. contributed to the immunofluorescence analysis; L.-G.D. contributed to RNA-seq analysis; Z.-Y.H. and J.-F.C. contributed to flow analysis; G.-M.Y., W.Y., K.-F.M., and X.L. contributed to western blot analysis; Y.F., X.-Z.Z., Y.-A.Z. contributed to real-time analysis. J.O.S. provided the anti-IgT, IgM, -IgD, and -plgR antibodies; Y.-Y.Y., Z.X., and J.O.S. wrote the paper.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Convergent Evolution of Mucosal Immune Responses at the Buccal Cavity of Teleost Fish

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Supplemental Information

Convergent evolution of the mucosal immune response in the buccal mucosa of teleost fish

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Figure S1. An overview of trout BM. Related to Figure 1

Anatomy of rainbow trout (A), BM (B), and paraffin sections of buccal upper mucosa (C) and lower mucosa (D), stained with Haematoxylin/eosin. The black dotted boxes represent the sampling site for paraffin sections. BC, buccal cavity; BE, buccal epithelium; LP, lamina propria; BS, buccal submucosa; CL, cartilage layer. Scale bar, 50 μm.
Figure S2. Candidate fish species from five different families were selected to understand the general organization of teleost BM. Related to Figure 1

Figure S3. Abundant mucous cells in teleost BM. Related to Figure 1

AB staining of the BM of a control adult Japanese seabass (*L. japonicus*) (A), Asian swamp eel (*M. albus*) (B), Southern catfish (*S. meridionalis*) (C), and Snakehead (*C. argus*) (D). Red triangles indicate mucus cells. Scale bar, 50 μm. BC, buccal cavity; BE, buccal epithelium; LP, lamina propria; BS, buccal submucosa.
Figure S4. Isotype control staining for anti-IgT, anti-IgM, anti-pIgR and anti-Ich antibodies in trout BM paraffin-sections. Related to Figure 1

Differential interference contrast images of buccal paraffin-sections from control fish (A and B) and infected fish, with merged staining of isotype control antibodies for anti-trout pIgR pAb (green, A); or anti-trout IgT pAb (green) and anti-trout IgM (red, B) mAb; or anti-trout Ich pAb (magenta, C). Nuclei were stained with DAPI (blue, A-C). BC, buccal cavity; BE, buccal epithelium; LP, lamina propria. Scale bars, 50 µm. Data are representative of three independent experiments.
Figure S5. Protein characterization of buccal mucus immunoglobulins. Related to Figure 1

(A) Fractionation of buccal mucus (~0.5 mL) by gel filtration (upper) followed by immunoblot analysis of the fractions with anti-trout IgM, anti-trout IgD-specific mAbs or anti-trout IgT-specific pAbs (lower). (B) SDS-PAGE of gel-filtration fractions (4–15%) corresponding to elution volumes of 8.5 and 11.5 mL under non-reducing conditions followed by immunoblot analysis with anti-trout IgM-, anti-trout IgD-specific mAbs or anti-trout IgT-specific pAbs.

Figure S6. Staining of trout buccal bacteria with isotype control antibodies for anti-IgT, anti-IgM and anti-IgD mAbs. Related to Figure 2

Differential interference contrast images (DIC) of buccal bacteria stained with a DAPI-Hoeschst solution (blue), isotype control antibodies for anti-trout IgT (green), for anti-trout IgM (red), or for anti-trout IgD (magenta) mAbs, and merging isotype control antibodies for IgT, IgM and IgD staining. Scale bar, 10 µm. Upper and lower panels display two different samples, representative of at least three independent experiments.
Figure S7. A heatmap with clustering using the RPKM (Reads Per Kilobase per Million mapped reads) of genes present in Table S4 (C14d, day 14 control group; C28d, day 28 control group; E14d, day 14 exposed to Ich group; E28d, day 28 exposed to Ich group). Related to Figure 3

Pheatmap package of R (version 3.4.4) was used to picture the heat map, and ‘single’ method was used to cluster values. The values were scaled in the row direction.
Figure S8. Local IgT-, IgM- and IgD-specific specific responses in BM explants of survivor fish. Related to Figure 5

The BM, head kidney, and spleen explants (~25 mg each) from control and survivor fish were cultured for 7 days. Immunoblot analysis of IgT-, IgM- and IgD-specific binding to Ich in the culture medium of BM (A), head kidney (B) and spleen (C) (dilution 1:2) from control and survivor fish. (D-F) IgT-, IgM- and IgD-specific binding to Ich in dilutions of culture medium from BM (D), head kidney (E) and spleen (F) from control and survivor fish, measured by densitometric analysis of immunoblots and presented as relative values to those of control fish (n = 9 per group).

*p < 0.05, **p < 0.01 and ***p < 0.001 (unpaired Student’s t-test). Data are representative of at least three independent experiments (Mean ± SEM).
**Figure S9. Trout pIgR associates with buccal sIgT. Related to Figure 1**

(A) SDS-PAGE under reducing conditions of trout serum and buccal mucus, followed by immunoblot analysis using anti-trout pIgR antibody. (B) Co-immunoprecipitation (CoIP) of buccal mucus with anti-trout IgT antibody, followed by immunoblot analysis (IB) under reducing conditions (pIgR detection, upper) or non-reducing conditions (IgT detection, lower). (C) CoIP of buccal mucus with rabbit anti-trout pIgR followed by IB under non-reducing conditions (IgT detection, upper) and reducing conditions (pIgR detection, Lower). IgG purified from rabbit’s serum before immunization (Prebleed) served as negative control for rabbit anti-trout pIgR and rabbit anti-trout IgT, respectively (left lane on each panel for B and C). (D) Immunofluorescence staining for pIgR with IgT in BM paraffin-sections of survivor fish. Differential interference contrast images of BM paraffin-sections were stained with anti-trout IgT (green), anti-trout pIgR (magenta) and DAPI for nuclei (blue) (n = 6). (isotype-matched control antibodies for anti-pIgR in Figure S4A). (E) Enlarged sections of the areas outlined in D without DIC showing some pIgR/IgT colocalization (white triangles). Scale bars, 20 μm. BC, buccal cavity; BE, buccal epithelium. Data are representative of at least three independent experiments.
|               | Polymorphic IgT | Monomeric IgT | IgT specific IgM | IgM specific IgG | Accumulation of IgT B cell | Accumulation of IgM B cell | Proliferation of IgT B cell | Proliferation of IgM B cell | Microsomes coated with IgT | Microsomes coated with IgM | Microsomes coated with IgG | The transport of IgG is associated to tight |
|---------------|----------------|---------------|------------------|------------------|-----------------------------|-----------------------------|----------------------------|----------------------------|---------------------------|---------------------------|---------------------------|----------------------------------|
| Gut M.        | +++            | +             | +++              | +                | +++                         | -                          | ?                          | ?                          | +++                       | ++                       | ?                         | +++                               |
| Skin M.       | +++            | ++            | ++               | -                | +++                         | -                          | ?                          | ?                          | +++                       | +                        | ?                         | +++                               |
| Gill M.       | +++            | ++            | +++              | +                | +++                         | -                          | +                          | +++                        | +++                       | +                        | +++                       | +                                |
| Nose M.       | +++            | ++            | +++              | +                | +++                         | -                          | ++                         | ++                         | +                        | ++                       | ?                         | +++                               |
| Buccal M.     | +++            | ++            | +++              | +                | +++                         | -                          | ++                         | +++                        | +                        | ++                       | ?                         | ++                                |
| Serum/Head kidney | –            | +++          | +++              | –                | ++                         | –                          | –                          | –                          | –                        | –                        | –                         | –                                |

**Figure S10.** Comparative analysis immune responses in different mucosa and systemic organs of teleost fish. Related to Figure 5

+, low level; ++, medium level; ++++, high level; ++++, strong level; -, no detection; ?, unknown, M., mucosa
| Gene | GenBank accession no. | Forward primer | Reverse primer |
|------|----------------------|----------------|---------------|
| MHCII | DQ246664.1 | GGTGAGTTGTTGGATAAC | AGCGTTAGGCTTCATAGA |
| CCL19 | XM_021602563.1 | GCTGCCACTCTCTCATCTCT | CTTGCTTTTCCCTTATG |
| IgD | JN173049.1 | CAGGAGGAAATGCTCGACATCA | CCTAAGGAGCTCGTGGTGG |
| pIgR | FJ940682.1 | AGAACGGTTGAGTGGTGCTA | AAGCCTTGGTCAGGTG |
| IgM | OMU04616 | AAGAAAGCCTACAAGAGGAGA | CGTCAACAAAGCAGCAGCA |
| MPO | GBTD01119227 | GGACGAGATCTCAGAAGCAA | ATCCACGGGCTAAACCTTT |
| FceRIα | AI69533.1 | TACTCAAACCTCTCATCTCT | CTTGCTTTTCCCTTATG |
| GCSA | J6196901 | TGGTCAGAGAGACGATAGATCC | GAGGTACTTGTTTGTGGCAT |
| CD4-1 | AY973028 | TCGACAGCTGTCCTTCTCT | CGTTATAGGGTTACGCACTTT |
| MPEG1 | GBTD01065710 | CGCGACGCTTCCTCTCTC | CGTTATAGGGTTACGCACTTT |
| CD4-2β | AY89932 | TGGTCAGAGAGACGATAGATCC | GAGGTACTTGTTTGTGGCAT |
| CD3γ | GBTD01057626 | GAAACTGCTTCGATGGCTTC | GAGGTACTTGTTTGTGGCAT |
| Lck | AY973033 | CTCCTTATGTCCTCAGAAGCAA | ATCCACGGGCTAAACCTTT |
| CD8a | AF178053 | TGGTCAGAGAGACGATAGATCC | GAGGTACTTGTTTGTGGCAT |
| TRα | OMU50991 | CAGGAGGAAATGCTCGACATCA | CGTCAACAAAGCAGCAGCA |
| MCSFRA | AB091826 | ATGGTTCAGGCGAAGCAA | ATCCACGGGCTAAACCTTT |
| RORγ1 | NM_001199827.1 | AAGACCTTTCAAGGACTTTTGTTTG | GGGAAAGCTGGACACACCATCTT |
| CD4-2α | AY772711 | CTTGACAGAGGCAAGCCAGAAGG | GGGAAAGCTGGACACACCATCTT |
| Gata3 | NM_001195792.1 | ACAGAAAGCTGTCCTGTTG | GGGAAAGCTGGACACACCATCTT |
| CD8b | AY563420 | CACCTGTCCTCAGAAGCAA | ATCCACGGGCTAAACCTTT |
| lag3 | XM_021590439.1 | TGGTCAGAGAGACGATAGATCC | GAGGTACTTGTTTGTGGCAT |
| Lck1 | AY973032 | TCGCTACAGGAGGGCTCTT | AGGTTGGGCGAAGGGTTT |
| CD40L | NM_001124666.1 | CACCTGTCCTCAGAAGCAA | ATCCACGGGCTAAACCTTT |
| IL-6 | CCV01624.1 | ATTTCATCGTTCTCAGACG | ACTACCTCAAGGAGCC |
| M-CSFR | NM_001124739.1 | CCTCCTCCCTCAACACTT | CTTGCTTTTCCCTTATGG |
| C3-1 | L24433 | GAGATGGCCCTCAAGGATAGA | ACCGATGATGGCTGATATTT |
| IFNAR | AGO14285.1 | CAGAGCCTGTCAGAAGCAA | ATCCACGGGCTAAACCTTT |
| C1QL2 | XM_021624859.1 | GTCTACTGACACACAGG | CATTCTTGGTCAACACAC |
| IgG | AY870264 | CAGACAAGACGCCTCCCTACTT | GAGTCAATAAGGAGACACAC |
| CD22 | XM_021625667.1 | TGAAGGATACAGGCGAAG | GAGTCAATAAGGAGACACAC |
| C7-1 | NM_001124618.1 | TGGTTCATCAGCCACAGTG | TAGCCTGTAACCTGACATAG |
| IL-10 | NM_0011245099.1 | CAGACAAGACGCCTCCCTACTT | GAGTCAATAAGGAGACACAC |
| IL-11 | NM_001124382.1 | CAGACAAGACGCCTCCCTACTT | GAGTCAATAAGGAGACACAC |
STAT1   NP_001118179.1   GACCAGCGAACCCAAGAACCTGAA   CACAAAGCCCGAGGATGCAACCAT
C1QBP   XM_021617398.1   CCGCAGTCCGAATTCTTA   GCTTTGCTACCTTCGATAT
EF-1alpha   NM_001124339.1   CAACGATATCCGTGCATGG   ACAGCGAAACGACCAAGAGG
CATH-D   XM_021617398.1   CCGCAGTCCGAATTCTTA   GCTTTGCTACCTTCGATAT
CCL-19*   KF683302.1   GTTTCCTGCCACTTCAA   GCCACCCACTTGCTCTTTG
IL-1β*   NM_001124347.2   TGATGTAATGGTGCTTCTGA   GATGGTGGAAGGTGTAAGG
CLCE4E*   XM_021562202.1   GCCAGCAACCCCTTACCAC   CACCACACCCTCAATCC
PIP5K*   XM_021585241.1   TCCATCGGCCTGCTCTCAT   TCCTCTCAACACCCACTTT
VWF*   XM_021580906.1   AGTGATGAAAGGTTGAGG   GTTGCCTGTTAGAAGGTCGT
HP-1*   XM_021595153.1   CGGAGGAGITGGAAGGC   GCAGGCAGAGCCAGGCAGC
CCL-13*   XR_002472294.1   CAGAACAACCTCCAGTAGC   ATCGTCGTTTAGAAGGTCGT
SAA*   XM_021607573.1   TTGTTCTGACCTCGTTG   CCTGGCAGACATAGTT
IgM*   EF467980.1   GCTATGGGATGAACTGG   TACCCTGAAATGACTGG
MHC II*   XM_021556605.1   AATGGCGACTGGCAGCT   GCCGATGGCTATCTTA
pIgR*   XM_021599266.1   TGTACACCTCCGATTTCTCA   CAGGGCAGGTTTCTGATTT

*Indicates the isoform specific primers used for validating the differentially expressed genes identified by RNA-Seq.
Table S2. Pathways involving in response to Ich (14 d). Related to Figure 3

| KEGG pathway | Description                                | Input number | Background number | q-value     |
|--------------|--------------------------------------------|--------------|-------------------|-------------|
| ko04974      | Protein digestion and absorption           | 82           | 228               | 3.19E-19    |
| ko04512      | ECM-receptor interaction                   | 52           | 179               | 4.39E-08    |
| ko04060      | Cytokine-cytokine receptor interaction     | 101          | 469               | 8.00E-08    |
| ko00330      | Arginine and proline metabolism            | 34           | 102               | 5.74E-07    |
| ko04510      | Focal adhesion                             | 87           | 466               | 4.33E-04    |
| ko05144      | Malaria                                    | 26           | 91                | 5.53E-04    |
| ko00680      | Methane metabolism                         | 17           | 48                | 6.40E-04    |
| ko04640      | Hematopoietic cell lineage                 | 33           | 134               | 7.49E-04    |
| ko05217      | Basal cell carcinoma                       | 34           | 149               | 2.49E-03    |
| ko05414      | Dilated cardiomyopathy (DCM)               | 44           | 213               | 2.96E-03    |
| ko00010      | Glycolysis / Gluconeogenesis                | 27           | 116               | 7.12E-03    |
| ko04360      | Axon guidance                              | 78           | 459               | 9.20E-03    |
| ko04151      | PI3K-Akt signaling pathway                 | 123          | 802               | 1.92E-02    |
| ko05020      | Prion diseases                             | 16           | 60                | 2.02E-02    |
| ko04978      | Mineral absorption                         | 22           | 97                | 2.74E-02    |
| ko00220      | Arginine biosynthesis                      | 13           | 46                | 2.87E-02    |
| ko05146      | Amoebiasis                                 | 36           | 188               | 2.87E-02    |
| ko00051      | Fructose and mannose metabolism            | 17           | 69                | 2.89E-02    |
| ko04514      | Cell adhesion molecules (CAMs)             | 58           | 345               | 3.59E-02    |
| ko05205      | Proteoglycans in cancer                    | 78           | 489               | 3.59E-02    |
| ko04657      | IL-17 signaling pathway                    | 33           | 175               | 4.33E-02    |
| ko04260      | Cardiac muscle contraction                 | 35           | 189               | 4.39E-02    |
| KEGG pathway | Description                              | Input number | Background number | q-value     |
|-------------|------------------------------------------|--------------|-------------------|-------------|
| ko00010     | Glycolysis / Gluconeogenesis             | 23           | 116               | 1.80E-06    |
| ko04610     | Complement and coagulation cascades       | 23           | 131               | 9.77E-06    |
| ko04514     | Cell adhesion molecules (CAMs)            | 41           | 345               | 1.01E-05    |
| ko04640     | Hematopoietic cell lineage               | 22           | 134               | 3.12E-05    |
| ko05150     | Staphylococcus aureus infection          | 14           | 61                | 4.88E-05    |
| ko05414     | Dilated cardiomyopathy (DCM)             | 28           | 213               | 6.89E-05    |
| ko04260     | Cardiac muscle contraction               | 25           | 189               | 1.64E-04    |
| ko05410     | Hypertrophic cardiomyopathy (HCM)        | 26           | 201               | 1.64E-04    |
| ko04512     | ECM-receptor interaction                  | 24           | 179               | 1.76E-04    |
| ko00680     | Methane metabolism                       | 11           | 48                | 3.59E-04    |
| ko04261     | Adrenergic signaling in cardiomyocytes   | 35           | 366               | 2.49E-03    |
| ko04530     | Tight junction                           | 39           | 441               | 4.95E-03    |
| ko05143     | African trypanosomiasis                  | 12           | 77                | 6.10E-03    |
| ko04971     | Gastric acid secretion                   | 20           | 178               | 7.48E-03    |
| ko04970     | Salivary secretion                       | 19           | 177               | 1.57E-02    |
| ko00030     | Pentose phosphate pathway                | 8            | 47                | 2.55E-02    |
| ko05020     | Prion diseases                           | 9            | 60                | 3.00E-02    |
| ko04060     | Cytokine-cytokine receptor interaction   | 37           | 469               | 3.15E-02    |
| ko04978     | Mineral absorption                       | 12           | 97                | 3.15E-02    |
| ko04974     | Protein digestion and absorption         | 21           | 228               | 4.10E-02    |
| ko04976     | Bile secretion                           | 16           | 159               | 4.44E-02    |
| ko05144     | Malaria                                  | 11           | 91                | 4.44E-02    |
Table S4. List of selected mRNAs, grouped according to functional classes (shown in bold), found to be up- and down-regulated by buccal infection with Ich (14 d and 28 d). Related to Figure 3.
Transparent Methods

Fish maintenance

Adult rainbow trout (triploid female fish, mean weight = 200–300 g) used for oral bacteria isolation and routine histology and juvenile rainbow trout (triploid female fish, mean weight = 20–30 g) used in infection trials were obtained from a fish farm in Shiyan (Hubei, China), and maintained in aquarium tanks with a water recirculation system including thermostatic temperature control and extensive biofiltration. Fish were acclimatized for at least 2 weeks at 15 °C and fed daily with commercial trout pellets at a rate of 0.5–1% biomass during the whole experiment periods. The feeding was terminated 48 h prior sacrifice. Japanese sea bass (Lateolabrax japonicus), Asian swamp eel (Monopterus albus), Southern catfish (Silurus meridionalis) and Snakehead (Channa argus) were purchased from aquatic product market in Wuhan (Hubei, China). Animal procedures were approved by the Animal Experiment Committee of Huazhong Agricultural University.

Ich parasite isolation and infection

For Ich parasite isolation, the method was described previously with slight modification (Yu et al., 2018). Briefly, heavily infected rainbow trout were anaesthetized with an overdose of MS-222 and placed in a beaker with water to allow trophonts and tomonts to exit the host. Fish were removed 4 h later, while the trophonts and tomonts were left in the water at 15 °C for 24 h to allow tomocyst formation and subsequent theront release. For Ich infection, two types of challenges were performed. In the first group, fish were exposed to a single dose of ~5,000 theronts per fish for 3 hours, and then migrated into the aquarium containing new aquatic water. Tissue samples including BM, head kidney and spleen were taken 0.5, 1, 4, 7, 14, 21, 28, and 75 days after infection. Moreover, fluids (serum and buccal mucus) were taken after 28 days (infected fish). In the second group, fish were monthly exposed for 75 days period (survivor fish) with the same dose. Fish samples were taken two weeks
after the last challenge. Both experiments were performed at least three independent times. As a control (mock infected), the same number of fishes were maintained in a similar tank but without parasites.

**Collection of serum, buccal mucus and bacteria**

For sampling, trout were anesthetized with MS-222, and serum was collected by centrifugation for 10 min at 4 °C, 5000 g and stored at -80 °C prior to use (Xu et al., 2016). To obtain the buccal mucus, briefly, fish BM tissue was excised and rinsed with PBS three times to remove the remaining blood. Thereafter BM tissue was incubated for 12 h at 4 °C, with slightly shaking in protease inhibitor buffer (1× PBS, containing 1× protease inhibitor cocktail [Roche], 1 mM phenylmethylsulfonyl fluoride [Sigma]; pH 7.2) at a ratio of 250 mg of BM tissue per mL of buffer.

The suspension (buccal mucus) was collected into an Eppendorf tube, and then vigorously vortexed and centrifuged at 400 g for 10 min at 4 °C to remove trout cells. To separate buccal bacteria from mucus, the cell-free supernatant was centrifuged at 10,000 g for 10 min at 4 °C. The resulting supernatant (containing buccal mucus) was harvested, filtered with 0.45 μm syringe filter (Millipore) and stored at 4 °C prior to use. The pellet (containing buccal bacteria) was washed three times with 1 mL of cold PBS (pH 7.2) and resuspended for further analysis.

**Gel filtration**

Gel filtration were performed to analyze the monomeric or polymeric state of Igs in trout buccal mucus using a Superdex-200 FPLC column (GE Healthcare) as presented previously for gut mucus (Zhang et al., 2010). The column was previously equilibrated with cold PBS (pH 7.2), and protein fractions were eluted at 0.5 mL/min with PBS using a fast protein LC instrument with ÄKTApurifier systems (GE Healthcare). Identification of IgM, IgD and IgT in the eluted fractions was performed by western blot using anti-IgM, anti-IgD and anti-IgT antibodies, respectively. A standard curve was generated by plotting the elution volume of the standard proteins in a Gel Filtration Standard (Bio-
Rad) against their known molecular weight, which was then used to determine the molecular weight of the eluted IgT, IgM and IgD by their elution volume.

**Isolation of trout head kidney and BM leucocytes**

To isolate trout head kidney and BM leucocytes, we modified the existing protocol as explained by us (Yu et al., 2018). Briefly, we anaesthetized the rainbow trout with MS-222 and collected the blood from the caudal vein. Trout head kidneys were removed aseptically and pressed through a 100-μm nylon mesh and suspended in Dulbecco’s modified eagle medium (DMEM, supplemented with 5% FBS, 100 U/mL penicillin and 100 μg/mL streptomycin). Then the BM was taken and washed with cold PBS to avoid blood contamination. Thereafter, the BM was cut into small pieces (approximately 0.1 cm²) in DMEM and then mechanically disaggregated on a 100-μm cell shredder on the ice. The cell fraction was collected, and the aforementioned procedure was repeated four times. The non-disaggregated BM tissue pieces were treated with PBS (containing 0.37 mg/mL EDTA and 0.14 mg/mL dithiothreitol DTT) for 30 min followed by enzymatic digestion with collagenase (Invitrogen, 0.15 mg/mL in PBS) for 30 h at 20 °C with continuous shaking, and mechanically disaggregated on a 100-μm cell shredder and the cell fraction was collected. Subsequently, the cell fractions from the above BM tissue treatments were pooled and passed through a 100-μm nylon mesh. Finally, the resulting cell fractions were washed three times in fresh DMEM and layered over a 51/34% discontinuous Percoll gradient. After 30 min of centrifugation at 400 g, cells lying at the interface of the gradient were collected and washed with DMEM medium.

**SDS-PAGE and western blot**

Serum and buccal mucus samples were resolved on 4–15% SDS-PAGE Ready Gel (Bio-Rad) under non-reducing and/or reducing conditions. For western blot analysis, the gels were transferred onto PVDF membranes (Bio-Rad). Thereafter, the membranes were blocked with 8% skim milk and
incubated with anti-trout IgT (rabbit polyclone antibody [pAb]) anti-trout IgM (mouse monoclonal antibody [mAb]) or biotinylated anti-trout IgD (mouse mAb) antibodies followed by incubation with peroxidase-conjugated anti-rabbit, anti-mouse IgG (Invitrogen) or streptavidin (Invitrogen).

Immunoreactivity was detected with an enhanced chemiluminescent reagent (Advanta) and scanned by GE Amersham Imager 600 Imaging System (GE Healthcare). The captured gel images were analyzed by ImageQuant TL software (GE Healthcare). Thereafter, the concentration of IgM, IgD and IgT were determined by plotting the obtained signal strength values on a standard curve generated for each blot using known amounts of purified trout IgM, IgD or IgT.

**Flow cytometry**

For flow cytometry analysis, leukocytes suspensions of trout head kidney and BM were double-stained with monoclonal mouse anti-trout IgT and anti-trout IgM (1 μg/mL each) on ice for 45 min. After washing three times, APC-goat anti-mouse IgG2b and PE-goat anti-mouse IgG1 (1 μg/mL each, BD Biosciences) were added and incubated for 45 min at 4 °C to detect IgT+ and IgM+ B-cells, respectively. Buccal bacteria were stained with mouse anti-trout IgM (1 μg/mL), anti-trout IgD (1 μg/mL), anti-trout IgT (2 μg/mL) or their respective isotype controls (1 μg/mL) at 4 °C for 1 h with continuous agitation. After washing three times, Alexa Fluor 488-goat anti-mouse IgG1, and Alexa Fluor 488-goat anti-mouse IgG2b were added respectively, and incubated for 1 h at 4 °C. To discriminate bacteria from debris, buccal bacteria were labelled with BacLight Red bacterial stain (Invitrogen), following the manufacturer’s instructions. After washing three times, analysis of stained leucocytes or bacteria was performed with a CytoFLEX flow cytometer (Beckman coulter) and analyzed by FlowJo software (Tree Star).

**Histology, light microscopy and immunofluorescence microscopy studies**
The BMs of control adult rainbow trout, southern catfish, Japanese seabass, and Chinese sturgeon, as well as the infected rainbow trout was dissected and processed for routine histology. All the BMs were fixed in 4% neutral buffered formalin, embedded in paraffin, sectioned, and stained with H&E and AB (Yu et al., 2018; Yashpal et al., 2007). Images were acquired in microscope (Olympus) using the Axiovision software. For the detection of Ich parasite as well as IgT⁺ and IgM⁺ B-cells, sections were stained with polyclonal rabbit anti-trout IgT (pAb; 0.49 μg/mL) and monoclonal mouse anti-trout IgM (IgG1 isotype; 1 μg/mL) for 2 h at 37 °C. After washing three times, sections were stained with Alexa Fluor 488-conjugated AffiniPure Goat anti-rabbit IgG and Cy3-conjugated AffiniPure Goat anti-mouse IgG (Jackson ImmunoResearch Laboratories Inc.) at 2.5 μg/mL each for 40 min at room temperature to detect IgT⁺ and IgM⁺ B-cells, respectively. After washing three times with PBS, mouse anti-Ich polyclonal antibody (1 μg/mL) was added and incubated at 4 °C for 6 h. After washing three times, Alexa Fluor 647-goat anti-mouse IgG (Jackson ImmunoResearch Laboratories Inc.) with 5 μg/mL were added and incubated at room temperature for 40 min to detect Ich parasite. For detection of trout buccal pIgR, we used the same methodology described to stain trout skin pIgR by using our rabbit anti-pIgR antibody (Zhang et al., 2010). As isotype controls, the rabbit IgG prebleed and the mouse-IgG1 isotype antibodies were labelled with the same antibody labelling kits and used at the same concentrations. All sections were stained with DAPI (4’, 6-diamidino-2-phenylindole; 1 μg/mL: Invitrogen) before mounting. For visualization of coating of buccal bacteria with IgT, IgM and IgD, the bacteria were firstly double-stained with rabbit anti-trout IgT and mouse anti-trout IgM (1 μg/mL each), or isotype controls (the rabbit IgG and the mouse-IgG1 (1 μg/mL each) at 4 °C for 2 h with continuous agitation. After washing three times, the secondary antibodies Alexa Fluor 488-conjugated AffiniPure Goat anti-rabbit IgG or Cy3-conjugated AffiniPure Goat anti-mouse IgG (Jackson ImmunoResearch Laboratories Inc.) at 2.5 μg/mL each were added and
incubated for 30 min at 4 °C. After washing as described above, biotin-labelled mouse anti-IgD antibody (1 μg/mL) was added and incubated at 4 °C for 2 h, after washing three times, Alexa Fluor 647-conjugated Streptavidin (Jackson ImmunoResearch Laboratories Inc.) with 5 μg/mL were added and incubated at 4 °C for 30 min. Before mounting, bacteria were stained with a mixed solution of DAPI and Hoechst 33342 dye (5 μg/mL; Molecular Probes). Stained bacteria were cytopspinned on glass slides and mounted with fluorescent microscopy mounting solution. All images were acquired and analyzed using an Olympus BX53 fluorescence microscope (Olympus) and the iVision-Mac scientific imaging processing software (Olympus).

**Proliferation of B-cells in the BM of trout**

For the proliferation of B-cells studies, we modified the methodology as previously reported by us (Xu et al., 2016; Yu et al., 2018). Briefly, control and survivor fish (~30 g) were anaesthetized with MS-222 and intravenous injected with 200 μg EdU (Invitrogen). After 24 h, leucocytes from BM and head kidney were isolated as described above. Subsequently, cells were incubated with mAb mouse anti-trout IgM and anti-trout IgT (1 μg/mL each) on ice for 1 h. After washing three times with DMEM medium, Alexa Fluor 488-goat anti-mouse IgG (Invitrogen) was used as secondary antibody to detect IgM+ or IgT+ B-cells. After 30 min incubation on ice, cells were washed three times and then fixed with 4% neutral buffered formalin for 15 min. EdU+ cell detection was performed according to the manufacturer’s instructions (Click-iT EdU Alexa Fluor 647 Flow Cytometry Assay Kit, Invitrogen). Cells were thereafter analyzed in a CytoFLEX flow cytometer (Beckman coulter) and FlowJo software (Tree Star). For immunofluorescence analysis, the paraffin sections of BM were incubated with rabbit anti-trout IgT (pAb; 1 μg/mL) and mouse anti-trout IgM (IgG1 isotype; 1 μg/mL) at 4°C for 45 min. After washing with PBS, paraffin sections were incubated with Alexa Fluor 488-conjugated AffiniPure Goat anti-rabbit IgG or Cy3-conjugated AffiniPure Goat anti-
mouse IgG (Jackson ImmunoResearch Laboratories Inc.) at 2.5 μg/mL each at room temperature for 45 min. Stained cells were fixed with 4% neutral buffered formalin and EdU+ cell detection was performed according to the manufacturer’s instructions (Click-iT EdU Alexa Fluor 647 Imaging Kit, Invitrogen). Cell nuclei were stained with DAPI (1 μg/mL) before mounting with fluorescent microscopy mounting solution. Images were acquired and analyzed using an Olympus BX53 fluorescence microscope (Olympus) and the iVision-Mac scientific imaging processing software (Olympus).

**Tissue explants culture**

Control and survivor fish were anaesthetized with an overdose of MS-222, and blood was removed from the caudal vein to avoid blood content in the collected organs. Thereafter, spleen, head kidney, and BM were collected. Approximately 20 mg of each tissue was isolated and submerged in 70% ethanol for 1 min to eliminate possible bacteria on their surface and then washed twice with PBS. Subsequently, tissues were transferred into a 24-well plate and cultured with 200 μL DMEM medium (Invitrogen), supplemented with 10% FBS, 100 U/mL penicillin, 100 μg/mL streptomycin, 200 μg/mL amphotericin B and 250 μg/mL gentamycin sulfate, with 5% CO₂ at 17 °C. After 7 days, supernatants were harvested, centrifuged and stored at 4 °C prior to use at the same day.

**Binding of trout immunoglobulins to Ich**

To assess whether infected and survivor fish had generated pathogen-specific immunoglobulins, we measured the capacity of IgT, IgM and IgD from serum, buccal mucus or tissue (BM, spleen and head kidney) explant supernatants to bind to Ich using a pull-down assay as described previously (Yu et al., 2018). Initially, approximately 100 tomonts were pre-incubated with a solution of 0.5% BSA in PBS (pH 7.2) at 4 °C for 2 h. Subsequently, tomonts were incubated with diluted fluids samples (buccal mucus, serum, or tissue explant supernatants) separately from infected, survivor, or control
fish at 4 °C for 4 h with continuous shaking in a 300 μL volume. After incubation, the tommonts were washed three times with PBS and bound proteins were eluted with 2× Laemmli Sample Buffer (Bio-Rad) and boiled for 5 min at 95 °C. The eluted material was resolved on 4–15% SDS-PAGE Ready Gel under non-reducing conditions, and the presence of IgT, IgM or IgD was detected by western blotting using the anti-trout IgT, IgM or IgD antibody as described above.

**Co-immunoprecipitation studies**

We followed the same strategy to detect the association of pIgR to IgT in gill mucus as we previously described (Xu et al., 2016). To detect whether polymeric trout IgT present in the buccal mucus were associated to a secretory component-like molecule derived from tSC, we performed co-immunoprecipitation analysis using anti-trout IgT (pAb) antibody with the goal to potentially co-immunoprecipitate the secretory component of trout (tSC). To this end, 10 μg of anti-IgT antibody were incubated with 100 μL of trout buccal mucus. As control for these studies, the same amount of rabbit control IgG (purified from the prebleed serum of the rabbit) were used as negative controls for anti-IgT. After overnight incubation at 4 °C, 20 μL of protein G Agarose (Invitrogen) was added into each reaction mixture and incubated for 1 hour at 4 °C. Thereafter, the beads were washed five times with cold PBS, and subsequently bound proteins eluted in 2× Laemmli Sample Buffer (Bio-Rad).

The eluted material was resolved by SDS-PAGE on 4–15% Tris-HCl Gradient ReadyGels (Bio-Rad) under reducing (for tSC detection) or non-reducing (for IgT detection) conditions. Western blot was performed with anti-pIgR or anti-IgT antibody as described above.

**RNA isolation and quantitative real-time PCR (qPCR) analysis**

Total RNA was extracted by homogenization in 1 mL TRIZol (Invitrogen) using steel beads and shaking (60 HZ for 1 min) following the manufacturer’s instructions. A spectrophotometry (NanoPhotometer NP 80 Touch) was used to quantitate the extracted RNA and agarose gel.
electrophoresis was used to determine the integrity of the RNA. To normalize gene expression levels, equivalent amounts of the total RNA (1000 ng) of each sample were used for cDNA synthesis with the SuperScript first-strand synthesis system for qPCR (Yeasen) in a 20 μL reaction volume. The synthesized cDNA was diluted 4 times and then used as a template for qPCR analysis. The qPCRs were performed on a 7500 qPCR system (Applied Biosystems) using the EvaGreen 2× qPCR Master mix (Yeasen). All samples were performed following conditions: 95 °C for 5 min, followed by 40 cycles at 95 °C for 10 s and at 58 °C for 30 s. A dissociation protocol was carried out after thermos cycling to confirm a band of the correct size was amplified. Trout housekeeping gene elongation factor 1α (EF1α) was used as control gene for normalization of expression. Primer sequences can be found in Table S1. The relative expression level of the genes was determined using the Pfaffl method (Pfaffl et al., 2001).

**RNA-Seq libraries and RNA-seq analysis**

The RNA-Seq libraries from twelve samples were generated according as a previous study (Abyzov et al., 2012). Briefly, polyadenylated RNA fragments were purified by a Dynabeads mRNA Purification Kit, fragmented with RNA fragmentation buffer, and reverse transcribed into first-strand cDNA using random hexamer and Superscript II reverse transcriptase, followed by second-strand cDNA synthesis using RNaseH and DNA polymerase I. The resulting cDNA was end-repaired, and a single “A” was added at the 3’ ends and a unique identifier (UID) was labelled at the 5’ ends before ligation to Illumina paired-end sequencing adaptors. PCR-amplified using Phusion High-Fidelity master mix and Illumina primers with the condition of 98 °C for 60 s, 15 cycles of 98 °C for 10 s, and 65 °C for 75 s, and concluding with 65 °C for 5 min.

All RNA-seq data were generated by Illumina paired-end sequencing with read length 150 bp. Reads were mapped to the *Oncorhynchus mykiss* genome using STAR with default parameters.
(Dobin et al., 2013). The mapped reads were analyzed by featureCounts (Liao et al., 2014). Differential expression was estimated with edgeR package (Robinson et al., 2010). We excluded the genes with low expression (CPM [count-per-million] < 1 in nine or more samples) from downstream analysis. The resulting genes were considered as differentially expressed genes (DEGs) if FDR ≤ 0.05 and |log₂ (fold-change)| ≥ 1. Pathway analysis of significantly differentially expressed genes was conducted with DAVID using all the expressed genes as background (Huang et al., 2009).

**Statistics**

An unpaired Student’s $t$-test and one-way ANOVA with Bonferroni correction (Prism version 6.0; GraphPad) were used for analysis of differences between groups. p values of 0.05 or less were considered statistically significant.

**Availability of data and material**

The NCBI Sequence Read Archive (SRA) accession number for the data reported in this manuscript is PRJNA560142.

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