Comparative Proteomic Analysis of Erythropoiesis Tissue Head Kidney Among three Antarctic Fish Species

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Research Article

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Abstract

Antarctic icefish is the only known vertebrate species that lacks oxygen-carrying hemoglobin and functional erythrocytes. To reveal the unique hematopoietic process of icefish, we used an integrated approach including tandem mass tag (TMT) labeling and liquid chromatography-tandem mass spectrometry (LC-MS/MS) to quantify the dynamic changes in the head kidney whole proteome of a white-blooded icefish, *Chionodraco hamatus*, compared to those in two other red-blooded Antarctic fish, *Trematomus bernacchii* and *Notothenia coriiceps*. Of the 4,672 identified proteins, in the Antarctic ice fish head kidney, 123 proteins were significantly up-regulated and 95 proteins were down-regulated. The functional grouping of differentially expressed proteins based on KEGG pathway analysis shows that white blood fish and red blood fish have significant differences in erythropoiesis, heme biogenesis, leucocyte and platelet cell development. The proteins involved in the hematopoietic process in icefish showed a clear trend of downregulation of erythroid lineage marker proteins and upregulation of lymphoid and megakaryocytic lineage marker proteins, including CD9, ITGB2, and MTOR, which suggests a shift in hematopoiesis in the icefish head kidney due to the loss of erythrocytes. The results of the present study not only provide basic datasets for the head kidney proteins of Antarctic fishes, but also provide important references for studies on immunity and hematopoiesis in various species.

1. Introduction

The marine environment of the Antarctica is characterized by a long-term icy snow cover on the ocean, which results in the temperature of sea water being constantly maintained at -1.9°C (Gonzalez-Aravena et al. 2016). Because of low temperature, more oxygen can dissolve in cold water, leading to the Southern Ocean becoming an oxygen-rich water region (O’Brien et al. 2020). The modern Southern Antarctic fish fauna comprises over 300 species of fishes (Eastman 2005; Thatje et al. 2008), and among them, the suborder Notothenioids (perciform) is the most abundant endemic fish fauna, which underwent considerable diversification with significant ecological differences. This suborder is divided to six families that include Nototheniidae, Arctedidraconidae, Bovichtidae, Harpagiferidae, Bathydraconidae, and Channichthyidae (Buonocore et al. 2006). To survive in such isolated and harsh cold conditions, these fishes have evolved numerous unique specializations to adapt, particularly antifreeze glycoproteins, which prevent freezing of the body fluids of the fish (Chen et al. 1997). The icefish of the Channichthyidae family, also known as the “white-blooded” fish, are the only species that have developed in this region due to the to lack of hemoglobin and functional active erythrocytes (Ito et al. 1995).

The Antarctic icefish *Chionodraco hamatus* (Perciformes: Channichthyidae) is a benthic fish that lives in the subzero waters of the Antarctic Ocean (Chen et al. 2019; Gerdol et al. 2019). As the only known vertebrate group that lacks hemoglobin (Evans et al. 2021), the blood circulation system of icefish has undergone large compensatory physiological changes. *Trematomus bernacchii* (Perciformes: Nototheniidae) (Della Pelle et al. 2020) and *Notothenia coriiceps* (Perciformes: Nototheniidae) (Cao et al. 2016), two red-blooded notothenioid fishes, share a similar living environment with *Chionodraco hamatus* (CH). The hematopoietic system, especially the immune system of the Antarctic fishes (the
icesh in particular), has undergone a rigorous selection process due to the harsh long-term coldness (Romano et al. 2000).

Head kidney is the main hemopoietic tissue in fish and is also an important immune organ with a cellular composition of erythrocytes and leucocytes, such as macrophages, granulocytes, and B lymphocytes (Romano 1998). In our previous study, we found more than 100 kinds of microRNAs, including miR-152, with high expression in the head kidney of CH, which might ultimately result in the suppression of erythropoiesis in icefish (Chan et al.). The Antarctic fishes also serve as a good candidate to study the evolution of the immune system under extreme cold environments.

Previous studies have shown differences in the histological structure of lymphomyeloid organs of the Antarctic fish CH and *Trematomus bernacchii* (Romano et al. 2000). The transcriptome and microRNA analysis of the primary hematopoietic tissue (head kidney) of CH and two red-blooded revealed the evo-devo mechanisms involving erythrophoeitic suppression through the upregulation of TGF-β signaling (Xu et al. 2015). Unique evolution of hepcidin genes was also noted in icefish as compared to that in other Antarctic fishes. However, little is known about the proteomic profile of the primary hematopoietic and immune-related tissue (head kidney) of icefish and other Antarctic fishes. In the current study, we used an integrated method including TMT labeling and LC-MS/MS to quantify the dynamic variations in the entire proteome (*Chionodraco hamatus*) of head kidney of icefish (CH) and two red-blooded notothenioids *Trematomus bernacchii* (TB) and *Notothenia coriiceps* (NC) with an aim to analyze the proteomic differences between the three Antarctic fishes. The results of the present study not only provide important references for further exploration of the unique hematopoietic process of the white-blooded icefish, but also shed a light on the innate and adaptive immune mechanisms of the Antarctic teleost.

2. Materials And Methods

2.1 Sample preparation

*Chionodraco hamatus* and *Trematomus bernacchii* individuals were gathered locally from Prydz Bay (69°22′ S, 76°22′ E). NC individuals were collected from the water near the Great Wall Station, a Chinese Antarctic research base (62°12′ S, 58°57′ W) located on the Fieldes Peninsula on King George Island. Head kidney tissues were dissected from adult male fish samples and used for total protein extraction. For each species, head kidney tissues from three different adult male fish individuals were dissected and used for total protein extraction. The average body length was approximately 22 cm for CH and NC and approximately 15 cm for TB individuals.

2.2 Protein extracts

The head kidney tissue was ground in liquid nitrogen for 20 minutes and sonicated three times on ice with a high intensity ultrasound processor. Then the samples in triplicate for each group were solubilized and centrifuged at a low-speed instantaneously, and the supernatant and intermediate layer are taken to add cocktail (ThermoFisher Scientific, USA) and 4% SDS (Shanghai Chinese Medicine, China). The
suspensions were disrupted by sonication on ice for 3 min, lysates were incubated on ice for 30 min. Subsequently, lysates were separated by centrifuge at 4°C for 20 minute and collected supernatants. Protein concentration of each sample was quantified by 2-D Quant kit.

Digested trypsin for digestion, take 100µg protein and replenish the volume to 100µl with lysate, add a final concentration of 10 mM DTT (ThermoFisher Scientific, USA) at 37 °C for 60 min, then adding iodoacetamide with an ultimate density of 20 mM (Sigma) at 20 °C for 45 min in the dark. Afterwards 200 mM TEAB was added to each sample for achieve urea density of less than 2 M (Wang et al., 2019). then centrifuged 10,000g for 20 min and collected the precipitation. Solubilized precipitation digested overnight at 37°C using trypsin with the mass ratio of Trypsin to protein 1:50.

2.3 TMT labeling

Finally, the peptides were dried by vacuum pump, labeling TMT reagents was added in advance (Thermo Fisher Scientific, Torrance, CA, USA) per 100 µg peptides, incubated for 2 hours at room temperature. Sample labeling was performed as follows: group CH: 129, group TB: 130, group NC: 131. Add 50ul of ultrapure water and leave it at room temperature for 30 min. Each group labeled samples of the peptide were mixed and dried down in a vacuum, the mixtures sample containing high-performance liquid chromatography (HPLC) were separated on loaded onto UltraPerformance LC (Waters, USA) containing a C18 column in front of a 2.1x 150 mm X Bridge BEH300 (Waters, USA). The HPLC gradient was buffer B (100% ACN) in buffer A (ammonia water, formic acid adjusted pH to 10) started with 0% B and increased to 100% B then followed by 0% B 6 min at a flow rate of 400µl/min total over 80 min. Collect 80 fractions based on peak shape and time and combine them into 18 fractions using rotation vacuum concentrators, the samples then were dissolved in mass spectrometry loading buffer the following analysis.

2.4 LC-MS/MS analysis and database search

The peptides were dissolved in 0.1% FA and separated from the high-performance liquid chromatography (HPLC) analysis were combined with liquid chromatography and tandem mass spectrometry( LC-MS/MS) containing EASY-nLC 1200, Q-Exactive(Thermo, USA)and C18 column (75µm x 25cm, Thermo ,USA). The schemes of gradient elution rise from 4–22% solvent B (0.1% FA in 98% ACN) in 26 min, 22–35% in 8 min, increased to 80% in 3 min, and then held at 80% for the last 3 min.

The MS data using data-dependent acquisition (DDA) model detection top 20, and following a MS measure scan over 350–1800 m/z at a resolution of 70,000 with the fragmentation pattern of high energy collisional dissociation, and, thena full scan at a resolution of 17,500 for MS/MS measured value was conducted and the dynamic exclusion time was 30s.

Use the PD search engine to process the MS/MS data. The *Dissostichus mawsoni*(DM) protein database was used as the common reference protein for set Tandem mass spectra searching (Chen et al. 2019) to study the differential protein expression patterns for the three notothenioids. In this study, DM is a closely related species of Antarctic fish. The laboratory sequenced and released the genome of DM(Chen et al. 2019).
2.5 Bioinformatics Methods

Gene Ontology (GO) annotation of the proteome was derived from DM Protein Database (Chen et al. 2019). Firstly, the KEGG database description of protein is annotated by Kaas. Then use the KEGG online service tool KEGG mapper to map the comment results on the KEGG path database.

Each class of proteins was enriched by domain analysis, and the InterPro database was studied. The two-tailed Fisher’s exact test was used to detect the enrichment degree of differentially expressed proteins for all identified proteins (Ma et al. 2020). GO annotation divides proteins into three categories: cellular compartment, molecular function and biological process. The corrected p-value of < 0.05 was considered to be significant.

3. Results

3.1 Quantitative overview

Reproducibility analysis of three biological replicate experiments was conducted by Pearson's correlation coefficient. For each replicate, relative expression of the proteins of CH, TB and NC was determined with Pearson's correlation coefficient (Fig. 1A). 4822 proteins were identified, among which 4672 proteins were quantified. A comparison of the proteome profile of CH with those of TB and NC showed that 123 proteins were upregulated (quantitative ratio of > 1.2 was considered as upregulation, CH vs. combined) and 95 proteins were downregulated (quantitative ratio of < 1/1.2 was considered as downregulation, CH vs. combined) in CH. The amount of the DEPs is summarized in Fig. 1B.

3.2 Functional classification of DEPs

The 218 DEPs were classified into three categories, namely biological processes, cellular components, and molecular functions, according to GO analysis (p < 0.05) (Fig. 2). These proteins were mainly clustered into 20 GO functional categories, which accounted for nine biological processes, nine cellular components, and two molecular functions (Fig. 2A–C). The most prevalent biological processes were response to inorganic substances, response to metal ions, erythrocyte homeostasis, and myeloid cell homeostasis, which accounted for 55% (Fig. 2A). According to cellular component annotation, the majority of the dysregulated proteins originated from the membrane (29%) and membrane part (15%). Other significant components included the endomembrane system (13%) and the endoplasmic reticulum part (9%) (Fig. 2B). The common molecular functions included gated channel activity and ion-gated channel activity (Fig. 2C).

3.3 GO enrichment of differentially quantified proteins

Figure 3 shows the results of the GO enrichment analysis. The differentially quantified proteins of the upregulated pathways for the three Antarctic fish species are shown in Fig. 3A. The most significantly
enriched cellular components were the endoplasmic reticulum membrane, the endoplasmic reticulum membrane network, and the endomembrane system. The main molecular functions were gated channel activity, ion-gated channel activity, and ion channel activity. The biological processes were mainly enriched in the glycoprotein metabolic process, gated channel activity, and divalent metal ion transport. The differentially quantified proteins of the downregulated pathways for the three Antarctic fish species are shown in Fig. 3B. The most significantly enriched cellular components were the main axon, muscle thin filament tropomyosin, and mitotic spindle. The main molecular functions were peptide binding and amide binding. The biological processes were mainly enriched in embryonic hemopoiesis, cardiac muscle contraction, hemoglobin biosynthetic process, and hemoglobin metabolic process. The GO enrichment analysis showed that the DEPs mainly participated in membrane, binding, and metabolic processes.

3.4. Pathway enrichment analysis of DEPs

KEGG analysis was performed to investigate the enriched pathways for the upregulated and downregulated proteins. The upregulated proteins were mainly mapped to 10 signaling pathways, which included hematopoietic cell lineage, malaria, cell adhesion molecules, protein processing in endoplasmic reticulum, HTLV-I infection, viral myocarditis, and others (Fig. 4). There are significant differences in erythropoiesis, heme biogenesis, and leucocyte and platelet cell development between the white- and red-blooded fishes (Souza et al. 2018). The downregulated proteins were mainly mapped to five signaling pathways: pentose phosphate pathway, porphyrin and chlorophyll metabolism, prion diseases, focal adhesion, and amoebiasis (Fig. 4).

On the basis of the differential protein function determined from the KEGG pathway, white-blooded fish and red-blooded fish showed significant differences in erythropoiesis and platelet cell development. The protein expression pattern of the hematopoietic process of teleosts shows that in icefish head kidney, erythropoiesis-related proteins are downregulated or almost inhibited and the production of megakaryocytic- and lymphopoiesis-related proteins is upregulated. The expression of B lymphocyte production-related proteins CD9 and ITGB2 was significantly upregulated ($p < 0.05$); the platelet production-related protein CD9 was significantly upregulated, and the expression of the myeloblast production-related proteins ITGB2, CD9, and MTOR was also significantly upregulated. This was in contrast to erythropoiesis-related proteins β-Spectrin, tfr1a, and hemoglobin, which were significantly downregulated and almost showed no expression (Fig. 5).
4. Discussion

In the present study, we conducted interspecies comparisons of head kidney proteomes between one icefish (CH) and two red-blooded notothenioids (TB and NC) by using TMT labeling and LC-MS/MS. We compared the expression profiles and described the differences of 123 upregulated proteins and 95 downregulated proteins. The comparison of the expression patterns of the teleost head kidney proteomes showed that the expression of erythrocyte-related proteins such as β-Spectrin, tfr1a, and hemoglobin was significantly downregulated in icefish; however, lymphoid and megakaryocytic lineage marker proteins, including CD9 and ITGB2, were significantly upregulated. An interesting point to note is that the lymphoid and megakaryocytic lineage marker proteins were not affected by the absence of erythrocytes and they were even upregulated in the icefish head kidney (Table 1). It is known that CD9 and ITGB2 play important roles in the development of B cells (Wang et al. 2020; Yang et al. 2020), and B lymphocytes are a subgroup of white blood cells that can differentiate into antibody-secreting cells. CD9 promotes the
recruitment of CD10 (a protease involved in the maturation and migration of B cells (Wang et al. 2020)) in exosomes (Navarro-Hernandez et al. 2020). CD9 is not expressed in primitive or memory B cells in mammals, but is expressed in plasma cell lines (Yoon et al. 2013) and in “innate-like” B lymphocytes (Won and Kearney 2002).

CD9 has also been found in multiple aquatic species, including rainbow trout and Atlantic salmon (Fujiki et al. 2002), red stingray (Zhu et al. 2006), shrimp (Wang et al. 2010), and lamprey (Wu et al. 2012). The fact that CD9 is constitutively expressed in trout primitive B cells indicates that a high proportion of primitive fish B cells is the equivalent of mammalian innate B cells (Zhu et al. 2014). ITGB2 is also an important immune effector that plays a vital role in protecting fish from a variety of pathogens (Wang et al. 2020). CD9 also plays an important role in platelet generation (Kaprielian et al. 1995; Aurora et al. 2018). Consistent with our results, Xu et al. (2015) has been reported that genes related to erythropoiesis and heme biogenesis are downregulated in the kidney of ice fish. While genes related to leukocyte differentiation and development, platelet activation, aggregation, and formation are upregulated (Xu et al., 2015). The proteomics analysis of the total protein in the head kidney of three Antarctic fish further confirmed that this general expression trend may cause a severe reduction in red blood cell differentiation of CH, while B cells and platelet cells are relatively up-regulated.

The major function of erythrocytes is not only to deliver oxygen to the organs, but they are also involved in innate immune responses, as they can capture specific immune complexes such as certain pathogens and bacteria, partly through membrane electrostatic attraction, and then kill them in the liver and spleen by presenting the pathogens to Kupffer cells and antigen-presenting cells (APCs) (Ukidve et al. 2020). In our study, the number of erythrocytes in the icefish was found to be severe reduced probably due to changes in the immune response; however, at present, we are unsure whether this change is more harmful or more beneficial. A comparison of the mucus microbial communities between white-blooded CH and red-blooded TB individuals sampled from the same locations showed much higher levels of bacterial species diversity in the CH mucus samples (data not published). To our knowledge, the fish skin mucus is a viscoelastic, adhesive gel that covers the exposed skin. The skin and mucus layer of fish are regarded as the first line of defense and play an important role in preventing the entry of pathogenic bacteria and similar harmful substances (Guellec et al. 2003; Laura et al. 2012). The low temperature of the surrounding environment, however, results in inhibition of the inflammatory responses (Finn and Nielsen 2010), including phagocytosis and acquired immune responses. Therefore, it remains unclear whether the increase in the number of mucus microorganisms indicates the increased susceptibility of the CH immune system to attack by microorganisms or whether it represents the ways through which the fish improves its acquired immunity; further studies are needed to confirm this aspect.

The present study mainly discusses the DEPs from total proteins of head kidney of three Antarctic fish species by comparing the significant differentially expressed proteins of red-blooded TB and NC and white-blooded CH. This study found 218 DEPs that could be used for further analysis of their related genes to provide a basis for further studies on the occurrence of red blood cells in icefish. Because there are almost no red blood cells in the Antarctic icefish, we found that among the three fish species, the
expression of proteins is important role in B-cell production and platelet cell development in the hematopoietic cell lineage was significantly upregulated CH. The results of this study can serve as good references for further research and exploration of the immunity and hematopoiesis of the Antarctic icefish.

Declarations

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Conflict of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Availability of data and material

All data generated or analysed during this study are included in this article [and its supplementary information files].

Code availability

Not applicable

Authors’ contributions
QHX was responsible for the overall guidance of this article. RNJ and SJH were mainly responsible for the overall experimental process and experimental data analysis involved in this paper; WYZ and SWJ participated in sample collection and processing; WHL and FXW participated in the analysis of experimental data and image processing.

**Ethics approval**

Experimental protocols involving live animals were approved by the Ethics Committee for the Use of Animal Subjects of Shanghai Ocean University.

**Consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

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Figures

Figure 1

(A) Reproducibility analysis. (B) Summary of differentially quantified proteins.

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Figure 2

Functional classification of differentially quantified proteins (CH vs combine). (A) Biological processes; (B) cellular components; (C) molecular functions.
Figure 3

GO-based enrichment analysis of up-regulated proteins (A) and down-regulated proteins (B) (CH vs combine).
Figure 4

KEGG pathway-based enrichment analysis of up-regulated and down-regulated proteins (CH vs combine).
Figure 5

The red arrows indicate the upregulated proteins, in the icefish compared with the red-blooded fish (see Table 1, Supporting information). HSC: haematopoietic stem cell; CMP: common myeloid progenitor; CLP: common lymphoid progenitor; MEP: megakaryocyte/erythroid progenitor; GMP: granulocyte/macrophage progenitor.

Supplementary Files

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- SupplementaryMaterialTable1.docx