Leptin Gene Protects Against Cold Stress in Antarctic Toothfish

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Leptin is a cytokine-like peptide, predominantly biosynthesized in adipose tissue, which plays an important role in regulating food intake, energy balance and reproduction in mammals. However, how it may have been modified to enable life in the chronic cold is unclear. Here, we identified a leptin-a gene (lep-a) in the cold-adapted and neutrally buoyant Antarctic toothfish Dissostichus mawsoni that encodes a polypeptide carrying four α-helices and two cysteine residues forming in-chain disulfide bonds, structures shared by most vertebrate leptins. Quantitative RT-PCR confirmed that mRNA levels of the leptin-a gene of D. mawsoni (DM-lep-a) were highest in muscle, followed by kidney and liver; detection levels were low in the gill, brain, intestine, and ovary tissues. Compared with leptin-a genes of fishes living in warmer waters, DM-lep-a underwent rapid evolution and was subjected to positive selection. Over-expression of DM-lep-a in the zebrafish cell line ZFL resulted in signal accumulation in the cytoplasm and significantly increased cell proliferation both at the normal culture temperature and under cold treatment. DM-lep-a over-expression also reduced apoptosis under low-temperature stress and activated the STAT3 signaling pathway, in turn upregulating the anti-apoptotic proteins bcl2l1, bcl2a, myca and mdm2 while downregulating the pro-apoptotic baxa, p53 and caspase-3. These results demonstrate that DM-lep-a, through STAT3 signaling, plays a protective role in cold stress by preventing apoptotic damage. Our study reveals a new role of lep-a in polar fish.

Keywords: leptin-a, positive selection, polar fish, STAT3 signaling, p53, Dissostichus mawsoni

INTRODUCTION

The obesity gene was first discovered on chromosome 6 in morbidly obese mice in 1994; the gene contains a 501-bp open reading frame (ORF) which encodes a polypeptide called leptin (Zhang et al., 1994). The leptin gene has subsequently proven to be highly conserved in other mammals, with similarities of up to 96% between rats and mice and 83% among rats, mice, and humans (Murakami and Shima, 1995; Sasaki et al., 1996; Konfortov et al., 1999). Studies on fish leptin were initiated more recently. In 2005, with reference to human and murine leptin, a leptin homolog was cloned in the Japanese pufferfish (Takifugu rubripes) via comparative genomics, promoting studies on ichthyic leptin (Kurokawa et al., 2005). However, the sequence similarity among bony fishes is relatively low. Further, fishes harbor multiple leptin homologs owing to gene duplication. For example, Japanese medaka (Oryzias latipes) harbors lep-a and lep-b, which are located on chromosomes 6 and 23, respectively, (Kurokawa and Murashita, 2009). Sequence alignment has
revealed that the sequence similarity between lep-a and lep-b in Japanese medaka is only 16.5%. Similarly, zebrafish (Danio rerio) also have two homologous genes, lepa and lepb (Gorissen et al., 2009), and the amino acid sequence similarity between the two is also low at 24%. The T. rubripes, Salvelinus alpinus (Arctic char), and Pelteobagrus fulvidraco genomes carry the leptin-a gene but not leptin-b (González et al., 2000; Kurokawa et al., 2005; Frøiland et al., 2010; Choi et al., 2014). In contrast, in Cyprinus carpio (common carp) and Salmo salar (Atlantic salmon), leptin-a is duplicated (Huising et al., 2006; Rønnestad et al., 2010). This suggests that the functions of leptins in fish may be more complicated than in mammals.

Recent studies indicate that leptin genes undergo rapid evolution and are subject to positive selection in many animal clades, including seals (Hammond et al., 2005; Yu et al., 2011), cetaceans (Yu et al., 2011), pikas (Ochotona spp.; Yang et al., 2008, 2011), heterotherm bats (Yuan et al., 2011), and primates (Bennet et al., 2002; Siltberg and Liberles, 2002; Gaucher et al., 2003; Berglund et al., 2005; Yang et al., 2011). These findings suggest that selection on leptin may be an important contribution to environmental adaptation. However, the question of whether the leptin gene of fishes living in extreme cold environments has also undergone evolutionary selection is of interests, but has been not reported.

Dissostichus mawsoni, belonging to the Antarctic notothenioids, has adapted to the extremely low temperature of the ocean surrounding Antarctica. It has extensive lipid (mostly triglyceride) deposits under the skin and in the musculature (Chen et al., 2019), compared to other fish species in the same depth of water. As leptin plays a central role in the regulation of lipid metabolism, we speculate that the leptin of D. mawsoni has undergone adaptive functional evolution, in addition to regulating fat deposition, it also plays an important role in cold temperature adaptation. Recent studies have shown that leptin can inhibit cell apoptosis by activating the STAT3 signaling pathway, leading to increased expression of anti-apoptotic genes and a decrease of pro-apoptotic gene expression (McGaffin et al., 2009). Furthermore, it has been shown in many cell types that STAT3 activation is associated with cell survival and proliferation (Kanda et al., 2004; Yu et al., 2014; Banerjee and Resat, 2016). There is also evidence that leptin exerts its anti-apoptotic effect by reducing the level of p53 (Toro et al., 2014). In this study, a leptin-a gene (lepa) from D. mawsoni was cloned and transfected to a fish-liver cell line. We revealed that D. mawsoni leptin-a provides cells with potent protection from the stresses of low-temperatures.

**MATERIALS AND METHODS**

**Cloning and Tissue Expression Analysis of the leptin-a Gene of D. mawsoni**

The fish specimens used in the present study were the same as those used in our previous study (Chen et al., 2019). Total RNA was extracted from the liver, using TRIzol (Invitrogen, Carlsbad, CA, United States), tested for purity via agarose gel electrophoresis, and spectrophotometrically quantified (NanoDrop2000, Thermo Fisher, CA, United States). One microgram of total RNA was reverse-transcribed into single-stranded cDNA, using the Prime Script RT Kit (Takara, Japan). Based on the cDNA library of D. mawsoni sequencing results previously obtained by our laboratory, primers lepaF/lepaR (Table 1) for cloning were designed by screening the sequence in comparison with its homologs. PCR was performed using rTaq polymerase (TaKaRa, Japan) according to the following protocol: 95°C for 5 min, 34 cycles of 95°C for 30 s, 58°C for 30 s, 72°C for 30 s. For tissue expression analysis, a FastStart Universal SYBR Green Master Kit (Roche, Germany) and LightCycler 480 (Roche, Germany) were used for qPCR with 18S as the reference gene (Lauth et al., 2002). The 18S primer was as used in the previous study. Thermal cycling was performed using an initial denaturation step of 95°C for 3 min followed by 40 cycles of 95°C for 10 s, annealing temperature for 20 s, and 72°C for 20 s. The relative tissue expression levels of lepa were calculated by the $2^{-\Delta\Delta C_T}$ method. The primer sequences are shown in **Table 1**.

**Selective Pressure Analysis**

To test the selective pressure of the lepa gene of D. mawsoni, we compared its sequence with those six fish species (L. japonicus, L. maculatus, S. chuatsi, E. coioides, O. latipes, and D. rerio) from relatively warm waters (more than 10°C). Sequences were aligned using MEGA 6 (Tamura et al., 2013), which generated the phylogenetic tree for lepa genes via the maximum-likelihood method. Two likelihood-ratio tests (LRTs) were performed by means of the branch and site models of codon evolution using the codeml program (Yang, 2007) from the PAML package (Yang, 1997). Significance ($P$ value < 0.05) of the compared LRTs was conducted by the chi2 program from the PAML package.

**Construction of DM-lepa Expression Vector**

For lepa over-expression, the CDS sequence of lepa was cloned into the constructed pTol2-actin-egfp plasmid (the promoter was β-actin from zebrafish) to produce pTol2-actin-lepa-egfp. Due to the lack of a specific lepa antibody, we also introduced the His-tag before the lepa termination codon to produce the LEPA-HIS fusion protein. In this way, we could use commercial HIS antibodies to detect LEPA protein expression for convenience. The expression plasmids then were transferred into E. coli DH-5α competent cells, plated on culture dish containing 100 µg mL⁻¹ kanamycin (Sangon, Shanghai, China) and cultured in a 37°C incubator for 12 h before analysis via individual bacterial colony

**Table 1** | Primers for D. mawsoni lepa cloning and tissue expression.

| Primer name | Primer sequence (5'-3') | Application |
|-------------|-------------------------|-------------|
| lepaF | GGGTTTGGAGAAGGAGGTAGAG | Cloning |
| lepaR | TGTTGGCAGCAAGCTTACATA | qPCR |
| lepaF | GATCGACCTTGAAGGCACT | Cloning |
| lepaR | CTCTGATGTTGTTGAG | Reference gene |
| 18SqF | CTGGATCCGCGAGAGGAAGG | Cloning |
| 18SqR | CCTCTCCTGGATTGTTACG | qPCR |
PCR. The positive clones obtained via sequencing underwent expansion culturing, and plasmids were extracted using a plasmid extraction kit (Promega, Wisconsin, United States) in accordance with the manufacturer's instructions and stored at −20°C until use.

**Culture and Plasmid Transfection of the Zebrafish Liver Cells**
The commercially available zebrafish liver (ZFL) cell line (American Type Culture Collection, ATCC, VA, United States) was cultured in culture medium containing 89% DMEM/F-12, 10% FBS, and 1% penicillin/streptomycin at 28°C and 5% CO₂ (i.e., conditions for optimum growth). Lonza Nucleofector (Lonza, Germany) was used for transfection at 70–80% confluence. G418 was added to the transfected ZFL cells after expansion culturing to select for successfully transfected cells. The concentration of G418 for the first week was 1,000 mg·L⁻¹ and 500 mg·L⁻¹ thereafter; these were the concentrations at which most of the non-transfected cells died. The G418 was then discarded, and the successfully transfected cells highlighted with green fluorescence.

**Protein Expression and Subcellular Localization of DM-lepa in Zebrafish Liver Cells**
The screened ZFL cells were cultured under optimal growth conditions for 48–72 h and then mixed with RIPA Lysis and Extraction Buffer (Thermo, CA, United States) and 1 mmol·L⁻¹ PMSF for protein extraction. The protein concentration was determined using a BCA Kit (Thermo, CA, United States) in accordance with the manufacturer's instructions, and proteins (30 µg per sample) were separated via SDS-PAGE and electro-transferred to a polyvinylidene difluoride nitrocellulose membrane (Millipore, Germany). Antibodies used in the assays included commercial HIS primary antibody (1:1000, Abcam, United Kingdom), β-actin (1:5000, Hua An, Hangzhou, China), and goat anti-mouse secondary antibody (1:5000, Hua An, Hangzhou, China). The proteins were visualized by enhanced chemiluminescence detection reagents (Beyotime, China).

To identify the localization of these proteins, ZFL cells were cultured in 24-well microporous plates (Corning, NY, United States) and fixed using 4% paraformaldehyde at ambient temperature for 30 min. Cells were then washed twice with PBS and blocked with treating buffer (0.1% Triton X-100, 1% BSA) for 30 min. Thereafter, the cells were counterstained with 2 µg·mL⁻¹ of 4,6-diamidino-2-phenylindole (DAPI, Sigma, St. Louis, MO, United States) for 5 min and washed thrice with water for 5 min each time. An inverted fluorescence microscope (ZEISS, Germany) was used to examine the ZFL cells and to obtain photographs.

**Cell Viability Assays by Cell Counting Kit-8**
A Cell Counting Kit-8 (CCK-8, Beyotime, China) was used to measure cell proliferation according to the manufacturer's protocol. The over-expression lepa group and control group cells were inoculated into a 96-well plate (Corning, NY, United States), and 10 µL CCK-8 reagent was added to each well at the time of harvest. After that, the cells were incubated at 28°C and 18°C for 0, 24, 48, 72, and 96 h, respectively. The low treatment temperature (18°C) refers to our previous research (Chen et al., 2017). At the indicated time points, the absorbance at 450 nm was measured to determine the cell viability using the microplate reader (Synergy H4 Hybrid Reader, BioTek, United States). The data are representative of three independent experiments in triplicate.

**ROS Detection by Flow Cytometry**
The generation of intracellular ROS was determined using a fluorescein-labeled dye, 2',7'-dichlorofluorescein diacetate (DCFH-DA, Beyotime, China), following the manufacturer's protocol. Briefly, the DCFH-DA fluorescent probe was added to cells (5 × 10⁴ cells/mL) followed by incubation for 20 min at their respective temperatures. Finally, cells were washed by PBS and then analyzed by a BD Accuri C6 flow cytometer (BD Biosciences, United States). FlowJo software (FlowJo, Ashland, OR, United States) was used to analyze the data.

**Western Blot Analysis of Signal-Transduction Proteins**
Total cellular proteins of each sample were separated by SDS-PAGE electrophoresis, and western blots were conducted as described in our previous study. After blocking, the membranes were overlayed with primary antibodies for cleaved caspase-3 (1:1000, Hua An, Hangzhou, China), STAT3 (1:1000, Hua An, Hangzhou, China), and p53 (1:2000, Hua An, Hangzhou, China) overnight at 4°C, then incubated with secondary antibody at 37°C for 1 h. Proteins were visualized by enhanced chemiluminescence detection reagents (Beyotime, China). The endogenous control used was β-actin.

**Analysis by qRT-PCR of Genes Involved in Signal-Transduction Pathway**
Total RNA was extracted from cells using TRIzol (Invitrogen, United States) reagent according to the manufacturer's instructions. The methods for RNA quantity, reverse transcription and real-time PCR were as mentioned above.
Supplementary Table 1 presents the primer sets used. Relative expression levels were calculated using the $2^{-\Delta \Delta CT}$ method, with β-actin as the reference gene for normalization.

**Sequence Analysis of the DM-lepa Gene**

ORFfinder\(^1\) was used to predict the ORFs and encoded amino acid sequence of DM-lepa. The NCBI BLASTP tool was used to search for the homologous gene sequences of leptin in different species, including *Lateolabrax japonicus* (Gene Bank accession number: AH857698.1), *Lateolabrax maculatus* (Gene Bank accession number: QFQ5150.1), *Siniperca chuatsi* (Gene Bank accession number: AHH86062.1), *Epinephelus coioides* (Gene Bank accession number: AMR58943), *O. latipes* (Gene Bank accession number: NP_001122048.1), and *Homo sapiens* (Gene Bank accession number: NP_001098190.2). The conserved motifs of the LEPA protein were analyzed using MEME\(^2\). The online tool ProtParam\(^3\) by ExPasy was used to predict the molecular weight of the DM-LEPA protein, its theoretical isoelectric point, instability index, aliphatic index, and grand average of hydropathicity at 0.092, suggesting a relatively low sequence similarity (<60%) among the fish species, although two cysteine residues forming disulfide bonds were conserved in vertebrates (Figure 1B). Prediction of the 3D structure also illustrated the structural similarity among fish and higher vertebrates, all of which carried four α-helices and irregular corners to form a hollow barrel structure (Figure 1C).

**Expression Patterns of the DM-lepa mRNA**

Levels of the DM-lepa mRNA were analyzed via quantitative RT-PCR, normalized relative to 18S mRNA of *D. mawsoni*. Transcription of DM-lepa mRNA was highest in muscle and lowest in intestine tissues. The next highest levels were observed in kidney and liver while weak expression was found in gill, brain, and ovary (Figure 2).

**Analysis of Selective Pressure**

Test of positive selection for the DM-lepa gene was carried out using the *codeml* program in the PAML package with the maximum likelihood codon model. The phylogenetic tree was constructed by MEGA 6 using the maximum likelihood model (Figure 3). The PAML branch model calculated the LRT test statistics to be $2\Delta \ell = 18.359215$, $p = 0.000018$, and df = 1 (Supplementary Table 2). The ω ratio for the DM-lepa branch ($\omega_1 = 1.71507$) was significantly higher than the fish species from warmer waters ($\omega_0 = 0.27320$). Furthermore, estimations of the PAML branch-site models identified 5 sites (43N, 96K, 103F, 108T, and 142R) under positive selection (Figure 3 and Supplementary Table 3), suggesting that DM-lepa underwent a rapid evolution.

**Identify and Molecular-Structure Analysis of the DM-lepa Gene**

The coding sequence of the *leptin-a* gene in *D. mawsoni* has a 501-bp ORF to encode 167 amino acid residues (Figure 1A). The initial 20 amino acid residues at its N terminus function as the signal peptide (Figure 1B). The relative molecular weight was determined to be at about 16 kDa, the theoretical isoelectric point at 6.73, the instability index at 51.50, the aliphatic index at 99.82 and the grand average of hydropathicity at $\sim 0.092$, suggesting a hydrophilic protein. Multiple sequence alignments indicated that lepa of *D. mawsoni* shared a similarity of 71.43% with human leptin, as well as 55.42, 56.02, 57.23, 58.43, 38.55, and 18.87% with *L. japonicus*, *L. maculatus*, *S. chuatsi*, *E. coioides*, *O. latipes*, and *D. rerio*, respectively, indicating a relatively low sequence similarity among the fish species, although two cysteine residues forming disulfide bonds were conserved in vertebrates (Figure 1B). The coding sequence of the DM-lepa gene was highest in muscle and lowest in intestine tissues. The next highest levels were observed in kidney and liver while weak expression was found in gill, brain, and ovary (Figure 2).

**Results**

Identification and Molecular-Structure Analysis of the DM-lepa Gene

The solvent accessibility of the mature sequence of DM-lepa was analyzed using TMHMM Server 2.0\(^5\) to predict the transmembrane domain. NetPhos 2.0\(^6\) was used to predict the Ser, Thr, and Tyr phosphorylation sites. NetNGlyc 1.0 Server\(^7\) was used to predict glycosylation sites. The SWISS-MODEL server\(^8\) was used to analyze higher-order protein structure.

Statistical Analyses

GraphPad Prism 8 (GraphPad Software, United States) was used for statistical analyses. All experiments were performed at least in triplicate. All data were expressed as the mean ± SD. Comparisons between two groups were performed using Student’s t-test, and comparisons among multiple groups were performed using two-way ANOVA. $P < 0.05$ were considered statistically significant. One asterisk, two asterisks and three asterisks indicate $< 0.05$, $P < 0.01$ and $P < 0.001$, respectively.

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**Over-Expression and Subcellular Localization of DM-lepa in Zebrafish Liver Cells**

Since the *D. mawsoni* population occupies extremely cold temperatures, temperature was expected to be the most influential factor in determining evolution of DM-lepa. We investigated *in vitro* the functions of DM-lepa, because of the difficulty of artificial breeding. Owing to the lack of a commercial antibody specific to DM-LEPA, we constructed the pTol2-lepa-His-egfp expression vector (Figure 4A) to help detect the counterparts of DM-LEPA from the total ZFL cell lysates using a commercial anti-His tag monoclonal antibody. We eliminated the stop codon and His tag in the expression vector (Figure 4B) and transfected ZFL cells (Figure 4C) to identify the subcellular localization of DM-LEPA. Western-blot analysis of the total protein in ZFL cells indicated that the molecular weight of the mature protein at 16.04 kDa was consistent with predictions (Figure 4D). Detection of green fluorescent proteins...
**FIGURE 1** | Characterization of *D. mawsoni* lepa. (A) Coding sequence and encoded amino acid sequence. The box and the asterisk indicates the start codon, the stop codon, respectively. (B) Alignment of *leptin-a* homologs among the vertebrates. The signal peptide and conserved function domains are indicated with the boxes. Identical amino acid residues are shown in black and the amino acids with >50% similarity shown in gray. Two asterisks indicate the conserved cysteine residues, which form a single disulfide bridge. (C) Predicted tertiary structure in different species. Four α-helices are displayed in brown, blue, dark green, and green helices.
Accordingly, we determined the cell regulation of cell proliferation and apoptosis (McGaffin et al., 1998; Fernández-Riejos et al., 2008; Mattioli et al., 2009). Our investigation of ROS generation indicated that over-expression of DM-lepa had no effect on the intracellular level of ROS at the normal culture temperature, while significantly decreasing ROS generation under cold treatment (Figures 5E,F,G); this demonstrates that over-expression of DM-lepa decreased apoptosis and stimulated cell proliferation under cold temperature.

**DM-lepa Over-Expression Activates the STAT3 Signal Pathway**

To gain further insight into the role of DM-lepa in facilitating proliferation and survival of ZFL cells, we focused on JAK-STAT3 pivotal signal cascades that were reported to regulate proliferation and apoptosis of cells (Takeda et al., 1998; Fernández-Riejos et al., 2008; Mattioli et al., 2009). The immunoblot analysis indicated that over-expression of leptin stimulated STAT3 expression in both normal conditions and under cold stress (Figure 6A). Using transcription of SOCS3/socs3a mRNA as a marker of the leptin stimulation involved in the JAK2/STAT3 pathway, the qPCR analysis confirmed significant upregulation of SOCS3/socs3a expression in over-expression cells (Figure 6B). We also checked the mRNA levels of related genes, Bcl-xl, Bcl-2, and c-myc, which were reported to be the target genes of STATs when apoptosis was suppressed by the activation of STAT3 (Yu et al., 2009; Abad et al., 2014). The quantified expression levels indicated that over-expression of DM-lepa enhanced the expression of c-myc/myca both at normal culture temperature and under cold conditions (Figure 6C). The expression of Bcl-xl/bcl2l1 did not show much difference in normal culture temperature (data not shown), but under cold temperatures it was significantly higher in DM-lepa cells than the control (Figure 6D). Bcl2 and Bax are important members of the Bcl2 protein family (Simchi et al., 2020) and Bax was identified as a Bcl2-interacting protein that opposed Bcl2 and promoted cell death (Oltval et al., 1993). Examination of Bcl2/bcl2a and Bax/baxa transcription showed that over-expression of DM-lepa significantly up-regulated the expression of the anti-apoptotic gene Bcl2/bcl2a (Figure 6E) while down-regulating Bax/baxa (Figure 6F) expression. These data indicate that DM-lepa was associated with upregulating anti-apoptotic genes and inhibiting pro-apoptotic genes via stimulating the STAT3 signal, thus benefiting cell proliferation and survival.

**DM-lepa Over-Expression Decreased p53 Level by Enhancing mdm2 Expression in Zebrafish Liver Cells Under Cold Temperature**

To confirm the p53 signaling pathway involved in apoptosis, a p53-specific antibody was used to evaluate the levels of the
corresponding protein in over-expression group cells. As shown in Figure 7A, over-expression of DM-lepa in ZFL cells decreased p53 levels under cold stress, which was probably negatively regulated by MDM-2, an E3 ubiquitin ligase, via a negative feedback loop that is essential to determining cell survival (Wade et al., 2013). Due to the lack of specific antibodies for zebrafish MDM-2, we checked the mRNA expression levels of MDM-2/ mdm2. As seen in Figure 7B, compared with the control group, over-expression of DM-lepa significantly increased the MDM-2/ mdm2 mRNA level under cold conditions. These results accorded with leptin suppressing the activity of p53 and inhibiting p53-mediated apoptosis by upregulating MDM-2/ mdm2 expression under cold temperature.

**DISCUSSION**

Leptin is the primary adipogenesis inhibitor produced by mammalian adipocytes (Friedman and Halaas, 1998). It acts on the central nervous system in the brain to regulate...
energy intake and consumption (Barb, 1999). In addition to regulating fat mass, mammalian leptin also plays a role in glucose homeostasis regulation (Morton, 2007) and reproduction (González et al., 2000). As opposed to mammalian leptin, data on the function of ichthyic leptins are so far preliminary. In this study, we cloned and identified the \textit{leptin-a} gene from...
**D. mawsoni**, a large slow-growing fish inhabiting the deep ocean at 300–2,500 m off the Antarctic coast where the temperature is perennially sub-zero. The amino acid sequence similarity between the DM-lepa mature peptide and lepa in other teleosts is relatively low, with only an 18.87% similarity between the DM-lepa mature peptide and zebrafish lepa, in accord with a previous study reporting that the primary structure of teleost leptin is not conserved. However, DM-lepa was predicted to contain four conserved antiparallel α-helices, forming a 4-helix structure consistent with mammals and most fish species. The two conserved disulfide bonds are potentially important in maintaining leptin structure and function. Mutagenic studies have reported that the disulfide bond formed by the two Cys residues is essential for leptin secretion (Keren et al., 2004). In teleosts, leptin homologs are expressed in many tissues, including the liver. For example, leptin-a expression is found to be highest in the liver in many fish species, such as *T. rubripes* (Kurokawa et al., 2005), *O. mykiss* (Murashita et al., 2008), *D. rerio* (Gorissen et al., 2009), and *O. latipes* (Kurokawa and Murashita, 2009). By contrast, lepa in *D. mawsoni* was highest in muscle. This might relate to the extensive lipid deposits under the skin and in the musculature of *D. mawsoni*.
of reduced ROS content in DM-lepa survival and proliferation in the ZFL cell line is consistent with our new knowledge of leptin in regulating cell fate.

In a previous study from our laboratory, it was found that the over-expression of calmodulin from Antarctic notothenioid fish increases cold tolerance in tobacco (Yang et al., 2013). Another study showed that elevated LINE activity from an Antarctic notothenioid fish *D. mawsoni* increases the number of viable cells in cold temperatures (Chen et al., 2017). In this study, over-expression of DM-lepa significantly increased survival and proliferation in ZFL cells, when stimulated by extreme low temperature. This suggests that DM-lepa has a new function as a cell protector at cold temperature and significantly broadened our new knowledge of leptin in regulating cell fate.

It has been reported that leptin activates human peripheral blood B-cells and maintains B-cell homeostasis by inhibiting apoptosis, inducing proliferation, and prolonging survival (Lam et al., 2010; Agrawal et al., 2011). In lean mice, the addition of mouse recombinant leptin significantly increased tracheal epithelial cell proliferation (Tsuchiya et al., 1999). Both lean and obese leptin-deficient mice exhibited increased cardiac apoptosis compared with wild-type mice (McGaffin et al., 2009). Our discovery that the fish leptin has beneficial effects of survival and proliferation in the ZFL cell line is consistent with many previous findings in mammals. However, the finding of reduced ROS content in DM-lepa transfected cells in cold stress from this study is novel and implies attenuation of cold stress-induced cellular damage; the underlying mechanism is an interesting avenue for further study. Leptin modulates cell function through activation of the Janus kinase (JAK)-STAT system. The activation of STAT1 or 3 by leptin is believed to involve cell proliferation in the liver (Takahashi et al., 1997; Wang et al., 1997). Leptin’s effects on cell apoptosis are likely mediated through STAT3 to increase anti-apoptotic *bcl-2* and *survivin* gene expression and reduces caspase-3 activity (McGaffin et al., 2009). Inhibition of STAT3 signaling in tumor cells increases the apoptotic rate (Morgan and Macdonald, 2019), and loss of active STAT3 has a significant impact on both cervical cancer cell proliferation and survival (Lis et al., 2017). The downstream targets of STAT3 include factors of anti-apoptotic (*bcl-2, bcl-XL*), pro-apoptotic (*baxa*) and proliferative genes (*c-Myc*, Kumar et al., 2016; Baek et al., 2017). Our results clearly show that DM-lepa over-expression markedly inhibited Bax/baxa expression but increased Bcl-2 expression under cold conditions, eventually resulting in an increased Bcl-2/Bax ratio and cell survival rate.

As the guardian of the genome, p53 is sensitive to environmental factors and is easily activated by a variety of stress signals, especially in response to temperature changes in aquatic organisms (Qian et al., 2020). Li et al. (2018) demonstrated that upregulation of p53 expression in response to low-temperature stress can cause tail malformation of the zebrafish (Li et al., 2018). Qian et al. (2020) reported that genes involved in the p53 signaling pathway were largely affected in the large yellow croaker response to cold stress (Qian et al., 2020). Similarly, p53 mRNA expression was significantly up-regulated in the muscle tissue of *D. rerio* under low-temperature stress (Li et al., 2018). The accumulation of p53 is prevalent in fish responses to cold stress, but over-expression of p53 leads to apoptosis (Chen et al., 1996; Sharp et al., 2014; Armstrong et al., 2017). Over-expression of DM-lepa attenuated the expression of p53, suggesting reduced genotoxic stress, corresponding with a lowered ROS level. It has been reported that leptin protects human trophoblasts from serum deprivation-induced cell death by decreased expression of p53 and increased level of MDM-2 (Toro et al., 2014). Over-expression of DM-lepa significantly increased survival and proliferation in ZFL cells, when stimulated by extreme low temperature. This suggests that DM-lepa has a new function as a cell protector at cold temperature and significantly broadened our new knowledge of leptin in regulating cell fate.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material; further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

The animal study was reviewed and approved by Shanghai Ocean University. Written informed consent was obtained from the owners for the participation of their animals in this study.
AUTHOR CONTRIBUTIONS

YW was responsible for the experimental design and completed the data analysis and wrote the manuscript. HW and LH performed the experiment. LC helped perform the analysis with constructive discussions. All authors contributed to finalizing and approving the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys. 2021.740806/full#supplementary-material

Supplementary Table 1 | Primers for genes involved in signal transduction pathway.

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