Isotocin controls ion regulation through regulating ionocyte progenitor differentiation and proliferation

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Abstract The present study using zebrafish as a model explores the role of isotocin, a homolog of oxytocin, in controlling ion regulatory mechanisms. Double-deionized water treatment for 24 h significantly stimulated isotocin mRNA expression in zebrafish embryos. Whole-body Cl⁻, Ca²⁺, and Na⁺ contents, mRNA expressions of ion transporters and ionocyte-differentiation related transcription factors, and the number of skin ionocytes decreased in isotocin morphants. In contrast, overexpression of isotocin caused an increase in ionocyte numbers. Isotocin morpholino caused significant suppression of foxi3a mRNA expression, while isotocin cRNA stimulated foxi3a mRNA expressions at the tail-bud stage of zebrafish embryos. The density of P63 (an epidermal stem cell marker)-positive cells was downregulated by isotocin morpholinos and was upregulated by isotocin cRNA. Taken together, isotocin stimulates the proliferation of epidermal stem cells and differentiation of ionocyte progenitors by regulating the P63 and Foxi3a transcription factors, consequently enhancing the functional activities of ionocytes.

Keywords Isotocin · Ionocyte · Zebrafish · Ion · Differentiation

Introduction

Oxytocin and vasopressin are structurally related neurohypophysial hormones in mammals [1, 2]. Isotocin and vasotocin are teleost homologues of mammalian oxytocin and vasopressin, respectively [1, 2]. According to a study on Fugu genomic organization, vasotocin/isotocin locus suggests that the two genes have evolved from a common ancestor through tandem duplication, and that this locus may have undergone a localized reorganization during vertebrate evolution [2]. Both oxytocin and isotocin stimulate contractions of smooth muscles and influence several behavioral and physiological processes such as social, sexual, and maternal behaviors, learning, memory, and parturition [3–5]. As for other physiological events, oxytocin and isotocin were demonstrated to regulate body fluids. Several studies suggested that dehydration or salt-loading resulted in increases in the transcription of the oxytocin gene and the proportion of neurons expressing oxytocin [6]. Oxytocin was assumed to directly act on the right atrium to stimulate atrial natriuretic peptide release, which acts to reduce the water, sodium, and adipose loads on the circulatory system, thereby reducing blood pressure [7].

Isotocin (isotocin-neurophysin, imp) was also proposed to be associated with the internal osmotic and ionic homeostasis in fish. Transfer from fresh water to seawater stimulated the plasma concentration of isotocin, suggesting
its involvement in osmoregulation of teleostean fish [8–11]. On the contrary, isotocin receptor was not well known to date. Isotocin receptor is expressed in many tissues including brain, gill, liver, spleen, heart, bladder, muscle, and kidney in teleost fish [12]. A study in eel indicated that upper esophageal sphincter muscle possesses isotocin receptor, which causes muscle relaxation and controls drinking behavior by enhancing cAMP production [13]. Studies of both mammals and fish suggested that oxytocin and isotocin are associated with hydromineral homeostasis of body fluids; however, our understanding of the physiological roles of oxytocin and isotocin in vertebrate ion- and osmoregulation is still fragmentary and needs further elucidation. The mechanisms behind these regulatory functions, including the target cells of oxytocin/isotocin, are still unclear.

Recently, oxytocin was suggested to be a growth and cell differentiation factor [14]. Oxytocin stimulates the proliferation of thymocytes [15] and enhances myoepithelial cell differentiation and proliferation in the mouse mammary gland [16]. The carboxyl terminally extended oxytocin precursor peptide induces embryonic stem cell-derived cardiomyogenesis and increases spontaneous beating activity [17]. During bone formation, oxytocin stimulates the differentiation of osteoblasts [18]. On the other hand, ionocytes in the skin/gills are the main cells responsible for fish ionoregulatory mechanisms, and the regulation of ionocyte functions was proposed to be mediated by the proliferation and differentiation of ionocytes [19, 20]. At least three subtypes of ionocytes were found in zebrafish skin/gills: Na⁺-K⁺-ATPase-rich cell (NaRC), H⁺-ATPase-rich cell (HRC), and Na⁺-Cl⁻ cotransporter cell (NCC), which are responsible for calcium uptake, acid/base balance, and chloride uptake, respectively [21–24]. In addition, several transcription factors that are related to ionocyte differentiation have been identified. The interaction between forkhead box transcription factors, Foxi3a and Foxi3b, was demonstrated to control NaRC and HRC differentiation [25–28]. Recently, another transcription factor, glial cell missing homolog 2 (Gcm2), was identified to specifically influence differentiation and maturation of HR cells [27, 29]. When zebrafish embryos are acclimated to acidic environments, the number of HR cells increases, causing enhancement of the acid-secreting function [30]. Similarly, the number of epithelial calcium channel (ECaC)-expressing cells (a subtype of NaRC) and ecac messenger (m)RNA expression were elevated after low-Ca²⁺ treatments [21]. Taking those results into consideration, we hypothesized that isotocin may modulate the proliferation and differentiation of ionocytes to control ionoregulatory mechanisms.

Zebrafish (Danio rerio) have a similar body plan and essentially the same organ systems as humans [31], and the functions of zebrafish genes can be easily evaluated in vivo by overexpression and antisense morpholino oligonucleotide-mediated knockdown experiments in embryos and larvae. The ionoregulatory mechanisms of zebrafish were recently well studied. According to serial molecular physiological studies, a working model of ionoregulatory mechanisms in the skin and gills of zebrafish, and the molecular physiological approaches to understand fish ion regulation and the controlling pathways were accordingly established [19, 32, 33]. Therefore, zebrafish are suitable to provide a platform to study the hormone control of ionoregulatory mechanisms. In the present study, we used zebrafish as a model to investigate the roles of isotocin on ionoregulatory mechanisms. Isotocin mRNA expression was induced by hypotonic stimulation. Knockdown of isotocin retarded whole-body ion contents, ionocyte-related gene expressions, and ionocyte densities; while overexpression of isotocin caused an increase of ionocyte cell numbers. Moreover, isotocin morphants showed lower foxi3a (a transcriptional factor controlling ionocyte differentiation) [25–28, 34] expression level at the tail-bud stage and a decrease in P63 (an epidermal stem cell marker) [35, 36] expression; while injection of isotocin capped-mRNA stimulated foxi3a and P63 expressions, suggesting that isotocin is involved in the differentiation and proliferation of ionocyte progenitors. Our results elucidate a molecular/cellular pathway of isotocin control in ionoregulatory mechanisms and provide new insights into this issue.

Materials and methods

Experimental fish

The AB strain of zebrafish (Danio rerio) were obtained from the stock of the Institute of Cellular and Organismic Biology, Academia Sinica, and mature fish were raised in a circulating system at 28°C under an illumination cycle of 14/10 h of a light/dark photoperiod. Embryos were also allowed to develop at 28°C. For hypotonic treatments, embryos were incubated in double-deionized water (Milli-RO60; Millipore, Billerica, MA, USA) for 72 h (for measurement of Ca²⁺ and Na⁺ contents) or incubated in local tap water for 72 h and then transferred to double-deionized water for another 24 h (for detection of the itnp mRNA expression level). Experiments were performed in accordance with guidelines of the Academia Sinica Institutional Animal Care and Utilization Committee (approval no. RFiZOOHP2007086).

Preparation of total RNA

In order to obtain a sufficient quantity of RNA, 30 embryos or adult zebrafish tissues dissected from six individuals...
were pooled as a sample. Samples were homogenized in 0.8 ml Trizol Reagent (Invitrogen, Carlsbad, CA). After chloroform extraction, the total RNA samples were purified and treated with DNaseI to remove the genomic DNA using an RNeasy Mini Kit (Qiagen, Huntsville, AL). The quantity and quality of total RNA were, respectively, assessed by Nanodrop spectrophotometry (ND-1000, NanoDrop Technology, Wilmington, DE, USA) and agarose gel electrophoresis.

Reverse-transcription polymerase chain reaction (RT-PCR) analysis

For cDNA synthesis, 5 µg of total RNA was reverse-transcribed in a final volume of 20 µl containing 0.5 mM dNTPs, 2.5 µM oligo(dT)20, 250 ng of random primers, 5 mM dithiothreitol, 40 units of an RNase inhibitor, and 200 units of PowerScript reverse transcriptase (Invitrogen) for 30 min at 50°C, followed by incubation at 70°C for 15 min. Then, 20 units Escherichia coli RNase H (Invitrogen) was added to remove the remnant RNA in a 20-min incubation at 37°C. For PCR amplification, 1 µl of cDNA was used as a template in a 25-µl final reaction volume containing 0.25 mM dNTP, 1.25 units of Gen-Tag polymerase (Genemark, Taipei, Taiwan), and 0.2 µM of each primer. The primer sets are shown in Table 1.

Quantitative (q)RT-PCR

mRNA expression levels of ionocyte-related genes, such as forkhead box I3a (foxi3a, NM_198917), Na⁺-K⁺ ATPase, beta 1b polypeptide (atp1b1b, NM_131671), epithelial Ca²⁺ channel (trpv6, NM_001001849), H⁺-ATPase subunit A (atp6v1a, NM_201135), the Na⁺-Cl⁻ cotransporter (slc12a10.2, NM_001045001), Na⁺/H⁺ exchanger 3b (nhe3b, EF591980.1), and isotocin (itnp, AY069956) were measured by a qRT-PCR with a Roche Lightcycler 480 (Roche, Penzberg, Germany). The final volume in a well was 10 µl, and contained 5 µl of 2× SYBR green master mix (Roche), 3.2 ng of cDNA, and 50 nM of primer pairs. The standard curve of each gene was checked in the linear range with β-actin, elongation factor 1-alpha (EF1a), and ribosomal protein L13a (RPL13a) as internal controls. For the qRT-PCR experiments, normalization using three reference genes showed similar patterns; therefore, only data normalized to β-actin are shown in the present study. Primers were designed using Primer Express 2.0 software (Applied Biosystems, Wellesley, MA, USA). The primer sets are given in Table 1.

Whole-body Na⁺, Ca²⁺, and Cl⁻ contents

Fifteen zebrafish larvae were briefly rinsed in deionized water and then pooled as one sample. HNO₃ at 13.1 N was added to the samples for digestion at 60°C overnight. Digested solutions were diluted with double-deionized water, and the total Na⁺ and Ca²⁺ contents were measured by atomic absorption spectrophotometry (Z-8000; Hitachi, Tokyo, Japan). For Cl⁻ content measurements, samples were homogenized with 1 ml deionized water and centrifuged at 14,000 rpm for 30 min. The supernatant was collected, and thereafter Hg(SCN)₄ (0.3 g in 95% ethanol) and NH₄Fe(SO₄)₂·12 H₂O (30 g in 135 ml 6 N HNO₃) solutions were added for the analysis. The Cl⁻ concentration was measured by the ferricyanide method with a double-beam spectrophotometer (model U-2000; Hitachi). Measurements of standard solutions of Na⁺, Ca²⁺, and Cl⁻ from Merck (Darmstadt, Germany) were used to make the standard curves.

### Table 1 Specific primer sequences for RT-PCR and qRT-PCR

| Gene name | Forward primer sequences (5’–3’) | Reverse primer sequences (5’–3’) |
|-----------|----------------------------------|----------------------------------|
| β-actin   | ATTGCTGACAGGATGCGAGAAG          | GATGTTCCAGACTCATGTACTC          |
| imp       | GTCAATCCACCGCCACAG              | GGGTGGCGAGTCGTTG               |
| impr1     | ACGCCCTTCTTCCTTGTCAG            | TATTTTCAGTGCTCTTGACG           |
| impr2     | TCACAGCATCGCCGAGGTTA            | CAAACCCACATAATGATGCACATC       |
| atp6v1a   | GAGGAAACACTGCATTCCCA            | GTCGGGATCATGTCCTGCTG           |
| atp1b1b   | CCAGGGTTTTAATTTGGGCTG          | TGGACTTGGCAGCTAGACACT          |
| gcm2      | TCCCTGGTTGTTGATCTTGGCA          | CACGCTTGAGTCGCCACTTTC          |
| foxi3a    | CTCTCGCTCAATGACTGCTTCA         | TAAACCAGGAGAAGGGCTCCTTG       |
| trpv6     | TCTTTCCTCACCTTCTCCTTCTCTCTC    | ATCAAGAATAGTTACCCGTAGCATTAC   |
| slc12a10.2| GCCCCCAAAGTTTTCAGGGTT          | CAGCCTGACACCTTTGTTTGG         |
| ef1a      | CTGGAGGCGGCGCTCAAACAT          | TGTGGCCTGCTCTGTTG             |
| rpl13a    | CCTCGGTCCTGTTCGCTTGCATT         |                                |
| nhe3b     | TGCAGACAGCCTCTAGC              |                                |
Translational knockdown with antisense morpholino oligonucleotides (MOs)

The morpholino-modified antisense oligonucleotide was purchased from Gene Tools (Philomath, OR, USA). The MO used against itnp begins at −5 and runs to ATG at the +20 nucleotide position (5′-GACAGCAGACC TCCAGACATTTC-3′). The maximal dosage that caused no obvious toxic effects on embryogenesis was used as follows: itnp MO at 1 ng/embryo and mismatched itnp MO (5′-GAGAGCACACCTCGAGACAA TAATC-3′) at 1 ng/embryo. The MOs were prepared with 1× Danieu solution (58 mM NaCl, 0.7 mM KCl, 0.4 mM MgSO₄, 0.6 mM Ca(NO₃)₂, and 5.0 mM HEPES; pH 7.6). The MO solution (1 ng/embryo) containing 0.1% phenol red (a visualizing indicator) was injected into zebrafish embryos at the 1–2-cell stage using an IM-300 microinjection system (Narishigi Scientific Instrument Laboratory, Tokyo, Japan). Wild-type without injection was only subjected to itnp protein measurement and morphological observation. In the subsequent knockdown experiments, mismatched-MO injected embryos were used as the control group.

Enzyme-linked immunosorbent assay (ELISA)

The sequences of zebrafish isotocin [Cys-Tyr-Ile-Ser-Asn-Pro-Ile-Gly-(NH₂)] and mammalian oxytocin [Cys-Tyr-Ile-Gln-Asn-Cys-Pro-Leu-Gly-(NH₂)] are highly conserved. In order to obtain sufficient amounts of protein, 25 embryos were pooled as a sample. Isolated sample were disrupted in homogenization buffer (100 mM imidazole, 5 mM EDTA, 200 mM sucrose, and 0.1% sodium deoxycholate; pH 7.6), and then centrifuged at 4°C and 10,000 rpm for 10 min. The supernatants were stored at −70°C until further experiments. The proteins were measured by Oxytocin Enzyme Immunoassay Kit (Assay Designs, Ann Arbor, MI, USA). Cross-reactivity of the antibody with the synthetic zebrafish isotocin (Kelowna International Scientific Inc., Taipei, Taiwan) was conducted to confirm the quantitative capability of the Oxytocin Enzyme Immunoassay Kit for zebrafish isotocin. Samples that contained 150 μg protein were loaded into each well and incubated at 4°C for 24 h, then the contents of the well were emptied and washed with a wash solution three times. After the final washing, any remaining wash buffer was removed, and the pNpp substrate solution was added to each well. The absorbance was measured at 450 nm in a synergy multi-mode plate reader (BioTek Instruments, Winooski, VT, USA) after stopping the enzymatic reaction by adding stop solution to each well. The standard curve for isotocin was constructed using commercial software (BioTek), and the concentration of isotocin in unknown samples was determined by interpolation.

Whole-mount in situ hybridization

Fragments of zapp6vl1a, ztrpv6, zslc12a10.2, and foxi3a were obtained by PCR and inserted into the pGEM-T easy vector (Promega, Madison, WI, USA). The inserted fragments were amplified with the T7 and SP6 primers by PCR, and the products as templates were used for the in vitro transcription with T7 and SP6 RNA polymerase (Roche) in the presence of digoxigenin (DIG)-UTP (Roche) to, respectively, synthesize sense and anti-sense probes. DIG-labeled RNA probes were examined using RNA gels, and a dot blot assay was used to confirm their quality and concentrations. Zebrafish embryos were anesthetized on ice and fixed with 4% paraformaldehyde in a phosphate-buffered saline (PBS; 1.4 mM NaCl, 0.2 mM KCl, 0.1 mM Na₂HPO₄, and 0.002 mM KH₂PO₄; pH 7.4) solution at 4°C overnight. Afterwars, the samples were washed with diethylpyrocarbonate (DEPC)-treated PBST (PBS with 0.1% Tween-20) several times (for 10 min each). After PBST washing, the samples were incubated with hybridization buffer (HyB, 50% formamide, 5× SSC, and 0.1% Tween 20) at 65°C for 5 min and with HyB containing 500 μg/ml yeast tRNA at 65°C for 4 h before hybridization. After overnight hybridization with 100 ng/ml DIG-labeled antisense or sense RNA probes, the embryos were serially washed with 50% formamide-2× SSC (at 65°C for 20 min), 2× SSC (at 65°C for 10 min), 2× SSC (at 65°C for 10 min), 0.2× SSC (at 65°C for 30 min, two times), and PBST at room temperature for 10 min. Afterwards, the embryos were immunoreacted with an alkaline phosphatase-coupled anti-DIG antibody (1:8,000) and stained with nitro blue tetrazolium (NBT) (Roche) and 5-bromo-4-chloro-3-indolyl phosphate (BCIP) (Roche) for the alkaline phosphatase reaction. For the quantification of density, eight areas (85 × 80 μm each) on the yolk sac surface of an embryo were chosen for counting.

Whole-mount immunohistochemistry

Embryos from different developmental stages were fixed in 4% paraformaldehyde in PBST. Samples were incubated with 3% bovine serum albumin (BSA) for 2 h to block nonspecific binding. Samples were then incubated overnight at 4°C with an z5 monoclonal antibody against the x-subunit of the avian Na⁺-K⁺ ATPase (Developmental Studies Hybridoma Bank, University of Iowa, Ames, IA, USA), and a polyclonal antibody against the a subunit of killifish H⁺ ATPase [37]. In addition, monoclonal anti-human P63 (Santa Cruz Biotechnology, Santa Cruz, CA, USA) was used to label zebrafish epidermal stem cells [35].
After washing with PBST for 20 min, samples were further incubated in Alexa Fluor 488 goat anti-rabbit immunoglobulin (IgG) (Molecular Probes, Carlsbad, CA, USA; 1:200 dilution with PBS) and an Alexa Fluor 568 goat anti-mouse IgG antibody (Molecular Probes; 1:200 dilution with PBST) for 2 h at room temperature. Images were acquired with a Leica TCS-SP5 confocal laser scanning microscope (Leica Lasertechnik, Heidelberg, Germany). The method of cell density quantification was described above.

Plasmid construction

To generate the pCS2\(^+\)-imp constructs, the corresponding imp coding region (465 bp) was PCR-amplified with the following pairs of primers: forward, 5'-GAATT CGGTGTCAGCCTTGGTGAATAATGTCTG-3'; reverse, 5'-TCTAGAGGCATTTCATTGGTGGATTCTGGAGGG G-3'; the imp PCR amplicon was cloned into a pGEM-T easy vector (Promega) with EcoRI and XbaI sites and was then subcloned into a pCS2\(^+\) vector at the EcoRI and XbaI sites. The corresponding EcoRI and XbaI sites are underlined.

Capped-mRNA (cRNA) injection

All constructs cloned in the pCS2\(^+\) vectors were linearized with NarI, and cRNA was transcribed using an SP6 message RNA polymerase kit (Ambion, Huntington, UK). Control embryos were injected with 1× Danieu solution. cRNA was injected into embryos at the 1 to 2-cell stage at 1 ng/embryo.

Statistical analysis

Values are presented as the mean ± SD and were compared using Student’s t test or one-way analysis of variance (ANOVA, Tukey’s pair-wise comparison).

Results

Expression of imp and its receptors in adult zebrafish tissues and embryos

mRNA expressions of the zebrafish imp and its two receptors (imp-like 1 and imp-like 2; accession number: FJ556869 and FJ556870, respectively) in various tissues were examined by RT-PCR, with β-actin as the internal control. imp mRNA was abundantly expressed in the brain, gills, muscles, and ovaries, and also expressed in the eyes, fins, heart, kidneys, liver, skin, and spleen (Fig. 1a). Both imp-like 1 and imp-like 2 were expressed in a variety of tissues but with a lower level of imp-like 2 in the eye (Fig. 1a). In zebrafish embryos, imp and the two receptors were first detected at 1 h post-fertilization (hpf), and the expressions were maintained till 72 hpf (Fig. 1b). The RT-PCR analysis was repeated with three different sets of samples, with similar results.

To examine the hypotonic effects on imp expression, 72-hpf embryos were transferred from fresh water to double-deionized water for 24 h, and imp expression levels were measured by qRT-PCR. As shown in Fig. 1c, imp mRNA was significantly upregulated after double-deionized water treatment.
Effects of *itnp* knockdown on *itnp* protein expressions and phenotypes in *itnp* morphants

Figure 1c implies a possible role of isotocin in ion regulation mechanisms in zebrafish. To explore the functional role of isotocin in zebrafish ion regulation, loss- and gain-of-function approaches with *itnp* MO and cRNA were used. To test the specificity and efficiency of the *itnp* MO and cRNA, we compared *itnp* protein concentrations in *itnp* morphants, mismatched-MO (Mis MO)-injected and cRNA-injected embryos by an ELISA. First, we confirmed the quantitative capability of the ELISA kit (Oxytocin Enzyme Immunoassay Kit). Zebrafish isotocin and mammalian oxytocin showed similar competitive binding curves with the kit (Fig. 2a), indicating the cross-reactivity of the antibody and the quantitative capability of the kit for *itnp* protein measurement.

The *itnp* MO was used to knock down the *itnp* protein expression. In preliminary experiments, we tried several dosages of *itnp* MO (1, 2, and 4 ng/embryo), and found higher dosages (2 and 4 ng) caused higher mortality and abnormal development and body shape. In contrast, the morphants injected with 1 ng MO appeared as wild-type (WT) embryos without injection. The mortality, hatching rate, body length, and body shape between morphants and WT embryos at 72 hpf were similar (Fig. 2c). We therefore

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![Diagram](https://via.placeholder.com/150)

*Fig. 2 Effects of knockdown or overexpression of *itnp* on protein expression and phenotypes in zebrafish. a Competitive binding curves of synthetic zebrafish isotocin and mammalian oxytocin with the ELISA (Oxytocin Enzyme Immunoassay Kit) showed a similar pattern. b One to two-cell stage embryos were injected with *itnp* MO (1 ng/embryo), mismatched-MO (Mis MO), and *itnp* cRNA (1 ng/embryo), and the protein expression at 72 hpf was analyzed by ELISA. *itnp* protein expressions were significantly downregulated in the morphants. In contrast, *itnp* cRNA-injected embryos showed twofold expression of *itnp* protein. No significant difference was found between wild-type (WT) without injection and Mis MO group.

*Fig. 2c The morphologies of *itnp* morphants, mismatched-MO-injected embryos, and WT embryos were similar. No significantly different phenotypes were found, suggesting that there were very few side effects in *itnp* morphants. d The whole-body contents of Cl⁻, Ca²⁺, and Na⁺ were significantly lower in *itnp* morphants than in Mis MO-injected embryos. Values are presented as the mean ± SD (n = 5). Different lowercase letters indicate significant differences (one-way ANOVA, Tukey’s pair-wise comparison). Asterisks indicates a significant difference from the Mis MO group (Student’s *t* test, *p* < 0.05). Scale bar 500 μm*
chose the dosage of 1 ng/embryo to do the subsequent experiments. In this situation, effects of developmental perturbation could be excluded or minimized when the physiological responses induced by imp MO were found. One nanogram of the imp MO and Mis MO were, respectively, injected into 1–2-cell-stage embryos. At 72 hpf, itnp protein expression in imp morphants was much lower than those in wild-type and Mis MO-injected zebrafish (Fig. 2b). This result clearly indicated that the imp MO significantly blocked itnp protein synthesis. In addition, isotocin protein expression in imp cRNA-injected embryos was twofold higher than that in the wild-type group (Fig. 2b). The morphology of imp morphants, Mis MO-injected embryos, and WT embryos were similar, and no significant phenotype differences were found (Fig. 2c). However, the Cl\(^{−}\), Ca\(^{2+}\), and Na\(^{+}\) contents were significantly lower in imp morphants than in Mis MO-injected embryos (Fig. 2d).

Effects of knockdown or overexpression of itnp on ionocyte-related gene expressions and ionocyte differentiation

As described above, isotocin appears to be involved in zebrafish ion regulation mechanism. Subsequent loss-of-function experiments were designed to test a hypothesis if isotocin is involved in the development of zebrafish skin/gill ionocytes, which are major cells responsible for ion regulation mechanisms. The mRNA expressions of ionocyte-related genes were downregulated by the imp MO. As shown in Fig. 3a, mRNA expressions of atp6v1a, atp1b1b, trpv6, slc12a10.2, and nhe3b were reduced in imp morphants. In whole-mount in situ hybridization experiments, RNA signals of atp6v1a, trpv6, and slc12a10.2 were also decreased by imp MO knockdown consistent with the qRT-PCR data (Fig. 3b, c). In the whole-mount immunocytochemistry experiments, NaRCs and HRCs were recognized. The cell densities of NaRC and HRC were reduced in itnp morphants compared with those in Mis MO-injected embryos (Fig. 4a–c). In contrast, increased cell densities of NaRC and HRC were detected in itnp-overexpressed embryos compared with those in the control injected with 1× Danieu solution (Fig. 4d–f).

Since decreases in cell number of different ionocyte subtypes and in expression levels of relevant transporters were observed in itnp morphants, we further investigated whether isotocin regulates ionocyte progenitor differentiation. We measured the mRNA expression of foxi3a, which was reported to be involved in ionocyte differentiation in zebrafish during early development stages [25–28]. foxi3a expression level (Fig. 5a, b) and the number of foxi3a-expressing cells (Fig. 5c, e, f) were downregulated by the imp MO and stimulated by itnp cRNA at the tail-bud stage when zebrafish embryos only expressed foxi3a but not ion transporter genes, indicating that isotocin affects ionocyte differentiation. The effects on foxi3a expression level modulated by imp MO injection or imp overexpression were still observed at the later stage, 3 dpf (Fig. 5a, b).

Effects of knockdown or overexpression of itnp on epidermal stem cells in zebrafish embryos

The loss- and gain-of-function approaches were used to further examine if the control pathways of isotocin in ionocyte progenitor differentiation are mediated by regulating epidermal stem cells. P63 is a marker of epithelial stem cells, which were demonstrated to differentiate into skin ionocytes and keratinocytes in zebrafish [35, 36]. To test whether isotocin regulates the number of epidermal stem cells in zebrafish embryos, P63 staining was performed. The density of P63 cells decreased in itnp morphants and, on the contrary, increased in itnp ectopically expressed embryos (Fig. 6). No difference was found between wild-type and mismatched MO-injected embryos (data not shown).

Discussion

The tissue distribution of mammalian oxytocin was reported [38, 39]. Oxytocin is primarily produced in magnocellular and parvocellular neurons of the hypothalamus, which plays an important role in integrating the vertebrate endocrine system [40]. The biological effects of oxytocin on female reproduction are well known [41]. In addition, oxytocin influences several behavioral and physiological processes such as social and maternal behaviors, learning, memory, steroidogenesis, and muscle contraction [3, 4, 42, 43]. In addition to be expressed in the central nervous system, mRNA was also detected in peripheral tissues including the heart, thymus, uterus, testes, adrenal gland, and ovaries [39], consisting with its wide spectrum of central and peripheral effects. In the present study, zebrafish imp mRNA was detected in various tissues and developing embryos by an RT-PCR analysis (Fig. 1). However, most studies in fish revealed imp mRNA expression in the brain [44–47]; only one paper reported that imp mRNA of the elephant shark was expressed in peripheral tissues [48]. Spatial distribution may differ among species, and further research is required to determine the exact functions of isotocin in these tissues. The highest expression level in the brain which is similar in all vertebrates indicates that the hypothalamus is the main site producing the isotocin peptide. The relative higher expression levels in gills, muscles, and ovaries reveal that isotocin regulates many physiological functions in fish such
neuropeptide. Hyperosmolality results in enhanced oxytocin secretion [39, 55]. Injection of hypertonic saline induced activation of neuronal activities in oxytocinergic cells in mouse hypothalamic paraventricular (PVN) and supraoptic nuclei (SON) [56]. After water deprivation, osmolality and plasma sodium were increased accompanied the elevation of oxytocin mRNA in PVN and SON [57]. The concentration of plasma oxytocin was decreased by water drinking, which caused a signal of osmotic dilution [58]. An oxytocin injection caused increases in urinary osmolality and natriuresis in the rat [59]. In response to hyperosmotic conditions, oxytocin stimulates water uptake and simultaneously reduces Na⁺ reabsorption, which result in downregulation of body fluid osmolarity. Likewise, studies on euryhaline teleosts also suggested that isotocin, an ortholog of oxytocin, is associated with hyperosmotic acclimation. In rainbow trout and sea bream, levels of isotocin increased when fish were acclimated to different salinities [60, 61]. Conversely, itnp mRNA levels in the pituitary decreased after tiger puffer were transferred from seawater to 10% seawater [47]. In European sea bass, the short-circuit Cl⁻ current increased when isotocin was added to the medium of cultured gill cells [62]. However, knowledge of the mechanisms behind these physiological phenomena is limited due to a lack of information on target cells of isotocin as well as the way in which isotocin controls the functions of target cells. The present study provides molecular physiological data exploring these issues.

In the present study, zebrafish embryos were treated with double-deionized water, and we surprisingly found that itnp mRNA expressions were stimulated in an ion-deficient environment (Fig. 1c). In euryhaline teleosts, isotocin was previously found to be induced by hypertonic environments [60]. In contrast, the present findings indicate a role of isotocin in a stenohaline freshwater species during acclimation to a hypotonic situation. Loss-of-function experiments supported this notion. Injection of the itnp MO caused lower body ion contents in morphants than those in Mis MO-injected embryos (Fig. 2c). The uptake functions of ions are accomplished by specific ion transporters which are expressed in different types of ionocytes in zebrafish: HRC, NaRC, and NCC [32, 33, 63]. Injection of itnp MO significantly decreased mRNA expression level of relevant transporters in ionocytes (Fig. 3) and downregulated cell densities of ionocytes (Figs. 3, 4), providing convincing and direct evidences that skin/gill ionocytes are target cells of isotocin. HRC, NaRC, and NCC were demonstrated to be similar to mammalian kidney proximal tubular cells, distal convoluted cells, and Ca²⁺-reabsorption cells responsible for Na⁺, Ca²⁺, and Cl⁻ uptake, respectively [32, 33]. Therefore, the present study opens a new window for exploring the roles of isotocin (and oxytocin) in

as reproduction and social behavior [49, 50] and support the view that isotocin is synthesized by a paracrine system operating within peripheral tissues [51–53]. It is noted that the transcripts of the two receptors, impr-like 1 and impr-like 2, were found in all the examined tissues including gill and skin where ionocytes appear, and were initially detected at early developmental stage. The broad expressions of oxytocin/isotocin and its receptors in mammals [39, 54] and fish tissues suggest the conservation and multiple biological actions of these peptides in vertebrates.

In light of body fluid homeostasis, oxytocin was demonstrated to be a modulator like vasopressin, another

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**Fig. 3** Effects of itnp MO on ionocyte-related gene expressions in zebrafish embryos. One to two-cell stage embryos were injected with itnp MO (1 ng/embryo) and a mismatched-MO (Mis MO), respectively, and the mRNA expressions at 72 hpf were analyzed by qRT-PCR. **a** The mRNA expressions of ionocyte-related genes were significantly downregulated by an itnp MO injection. **b** In situ hybridization analysis indicated that the mRNA signals of V-ATPase subunit A (atp6v1a), epithelial Ca²⁺ channel (trpv6), and Na⁺-Cl⁻ cotransporter (slc12a10.2) were lower in itnp morphants. **c** Cell densities of atp6v1a, trpv6, and slc12a10.2-expressing cells in itnp morphants were significantly lower than those in mismatched-MO-injected embryos. qRT-PCR values were normalized to β-actin. Mean ± SD (n = 6). Asterisks indicates a significant difference from the Mis MO group (Student’s t test, p < 0.05). Scale bar 200 μm.
ionoregulatory mechanisms not only in fish but also in mammals.

The numbers and densities of ionocytes are associated with the functional regulation of ion uptake acid/base balance mechanisms in fish during acclimation to environmental changes [19, 32, 33, 64]. The differentiation of ionocytes is regulated by a helix/forkhead box transcription factor, Foxi3a, which functions as a cell fate determinant or cell differentiation regulator [65]. Zebrafish Foxi3a was demonstrated to function as a master regulator of ionocyte differentiation in embryonic stages [25–28, 34]. During zebrafish embryonic development, foxi3a starts to be expressed at around the 90% epiboly to tail-bud stages, and the ionocyte markers, atp1b1b or atp6v1a, are expressed later at the 14–18-somite stages [25]. In the present study, foxi3a expression was significantly suppressed by itnp MO and stimulated by itnp cRNA at the tail-bud stage when skin ionocytes were not terminally differentiated and remained as progenitors (Fig. 5). On the other hand, overexpression of itnp caused increases of NaRC and HRC densities in zebrafish embryos (Fig. 4d–f), similar to the phenotypes of embryos injected with foxi3a cRNA [25]. These data indicate that isotocin affects the differentiation of ionocytes by regulating foxi3a expression. Moreover, P63+ cell densities decreased in itnp morphants and inversely increased in the itnp cRNA-injected embryos (Fig. 6), suggesting that isotocin may also be involved in stimulating the proliferation of epidermal stem cells. Epidermal stem cells are the source of ionocyte and keratinocyte progenitors. Regulation of epidermal stem cells would modulate ionocyte progenitors and then affect the number of ionocytes.

![Fig. 4 Effects of itnp MO and cRNA on cell densities of NaRC and HRC in zebrafish embryos. One to two-cell stage embryos were injected with the itnp MO (1 ng/embryo) (a), a mismatched-MO (Mis MO) (b), 1× Danieau solution (Control) (d), and itnp cRNA (1 ng/embryo) (e), respectively, and NaRCs (red) and HRCs (green) were detected by double immunocytochemistry of Na⁺-K⁺-ATPase (red) and H⁺-ATPase (green) at 72 hpf. Cell densities of NaRCs and HRCs in itnp morphants were significantly lower than those in mismatched-MO-injected embryos (e), while the cell densities in cRNA-injected embryos were significantly higher than those in the control group (f). Mean ± SD (n = 5–6). Asterisks indicate a significant difference (Student’s t test, p < 0.05). Scale bar 100 μm.](attachment:fig4.png)
Similarly, the actions of oxytocin on cell differentiation were reported [66]. The oxytocin/oxytocin receptor system exists in the heart, and activation of the cardiac oxytocin receptor stimulates the release of the atrial natriuretic peptide, which lowers blood pressure and promotes cell growth [67]. In cultured mouse P19 embryonic stem cells, incubation of oxytocin stimulated the production of beating cell colonies, while treatment with an oxytocin antagonist, ([d(CH2)51, Tyr(Me)2, Thr-4,Orn-8,Tyr-NH29] vasotocin) completely inhibited the formation of cardiomyocytes [68]. Oxytocin was also found to be an anabolic bone hormone [18]. Through upregulating BMP-2 and its downstream molecules, Schnurri-2 and -3, Osterix, and ATF-4, expressions, oxytocin stimulated osteoblast differentiation to exhibit the mineralizing phenotype [18]. Taken together, isotocin and oxytocin mediate the proliferation and/or differentiation of cells to regulate physiological responses, and this pathway seems to be conserved among vertebrates, thus indicating a common and important function of these peptides during evolution.

From an osmoregulatory point of view, previous studies proposed that oxytocin and isotocin act as hypertonic hormones to maintain body fluid hydromineral homeostasis; however, molecular mechanisms behind these physiological responses have not been revealed. In contrast, the present study used different approaches to identify target cells of isotocin and discovered a novel mechanism for isotocin to control ion uptake functions. Our data suggest that isotocin plays a role on regulation of the proliferation of epidermal stem cells and differentiation of ionocyte progenitors through mediation by modulating activities of transcription factors, Foxi3a and P63, to increase ionocyte densities and ultimately regulate the functions of ionocytes. However, more experiments will be necessary to further explore the detailed cellular and molecular mechanisms and to see if this function is direct or indirect.
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