

Inter-individual variability in freshwater tolerance is related to transcript level differences in gill and posterior kidney of European sea bass

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1	Inter-individual variability in freshwater tolerance is related to transcript level differences in gill and
2	posterior kidney of European sea bass
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18	

19 Introduction

20 In transitional habitat such as lagoons, fish have to deal with a wide range of changing environmental 21 parameters and therefore a high phenotypic plasticity is beneficial to cope with fluctuating environments. 22 Salinity can rapidly drop in these habitats through freshwater (FW) supplies by rainfalls or rivers. Low-salinity 23 environments can lead to differential distributions of marine species according to their acclimation capacity 24 (Pierce et al., 2012; Wong et al., 1999). Differential habitat distribution has also been observed between 25 individuals within the same species as in stickleback Gasterosteus aculateus and in mummichog, Fundulus heteroclitus, showing a differential capacity to regulate ions in FW (Scott et al., 2004; McCairns and 26 27 Bernatchez, 2009). To maintain hydromineral balance in FW environments, fish have to minimise diffusive ion 28 losses and compensate them by actively (re)absorbing ions at the gill and posterior kidney levels (Hickman and 29 Trump, 1969; Dantzler, 1996). The European sea bass, Dicentrarchus labrax, is known to undertake seasonal 30 migrations in transitional habitats where salinity fluctuates (Kelley, 1988; Barnabé, 1989; Waldman, 1995). 31 Previous studies highlighted intraspecific differences in FW tolerance in this species at different ages (Giffard-32 Mena et al., 2008; L'Honoré et al., 2019; Nebel et al., 2005) with about 30% mortality following FW exposure. 33 Fish that did not tolerate FW could survive when they were rechallenged to SW conditions and no intraspecific 34 variation was observed in SW (Giffard-Mena et al., 2008; L'Honoré et al., 2019). In F. heteroclitus, Scott et al. 35 (2004) concluded that the divergence in osmoregulatory capacities may result in different capacities to absorb 36 and reabsorb Na^+ and Cl^- at gill and/or kidney levels. In Nebel et al. (2005), it has been suggested that the 37 kidney might be responsible for the osmoregulatory failure detected in FW intolerant fish, linked to a low renal 38 Na⁺-K⁺-ATPase (NKA) activity and a lower kidney tubular density. In L'Honoré et al., (2019), authors 39 highlighted that intraspecific variation in FW tolerance of European sea bass is supported by strong differences in nka ala expression in the posterior kidney while no difference was measured at the gill level. Authors also 40 41 showed differences in corticosteroid receptors mRNA levels (gr1, gr2 and mr) with lower expression levels in 42 FW intolerant fish compared to FW tolerant. This suggests an impaired hormonal and stress regulation between 43 both FW-tolerance phenotypes. Pituitary prolactin (PRL) is a key hormone involved in FW acclimation by 44 promoting the maintenance of the hydromineral balance (Hirano 1986; Manzon, 2002; Sakamoto and 45 McCormick, 2006; Breves et al., 2014; Bossus et al., 2017). PRL interacts with PRL receptor (PRLR) to 46 regulate Na⁺ efflux, water permeability and the differentiation of ionocytes expressing Na⁺/Cl⁻ cotransporter 47 (NCC) as shown in tilapia and zebrafish (Breves et al., 2013, 2010; Dharmamba et al., 1967; Dharmamba and 48 Maetz, 1972). Prolactin receptors are cell surface receptors known to be expressed in osmoregulatory organs of many teleost species (Manzon, 2002). As for many fish species like *Takifugu rubripes*, two paralogs of *prlr*were identified in *D. labrax* genome called *prlra* and *prlrb* (Lee et al., 2006; Tine et al., 2014). It remains to be
determined if both *prlr* paralogs display the same expression pattern according to salinity and if mRNA levels
differ between FW-tolerance phenotypes in European sea bass.

53 In fish, blood pH levels are salinity-dependent with lower blood pH at low salinity than in SW as it was 54 demonstrated in European sea bass by Shrivastava et al. (2019). In 8 month-old European sea bass, FW 55 intolerance was characterised by a lower blood osmolality associated with an elevated Na⁺/Cl⁻ ratio, indicating a 56 metabolic alkalosis (L'Honoré et al., 2019). To regulate a high blood pH, fish have to excrete bases as HCO₃, 57 mainly associated with Cl⁻ uptake (Maetz and García Romeu, 1964; Goss and Wood, 1990, 1991; Tresguerres et 58 al., 2006). The apical anion transporters SLC26A6, involved in Cl⁻/HCO₃⁻ and Cl⁻/oxalate exchanges, are known 59 to be widespread among species in osmoregulatory organs including the posterior kidney (Mount and Romero, 60 2004; Sardella and Brauner, 2007; Xie et al., 2013; Knauf et al., 2018) and gills (Perry et al., 2009; Boyle et al., 61 2015) and could be potential entry routes for HCO_3^- in intolerant European sea bass to FW. In the gulf toadfish 62 for example, *slc26a6* was highly expressed in kidney (Grosell et al., 2009) to reabsorb Cl⁻ from the lumen to the 63 blood. Among three different SLC26 anion transporters in zebrafish (SLC26A6, SLC26A3 and SLC26A4, 64 called za6, za3 and za4), za6 seemed to be the most expressed in gills and was overexpressed when fish were 65 transferred to water with low Cl⁻ or NaHCO₃ (Perry et al., 2009). Moreover, gene knockdown of *slc26a6c* 66 resulted in a reduction in Cl⁻ uptake in zebrafish larvae confirming the major role of this transporter in Cl⁻ 67 uptake (Bayaa et al., 2009; Perry et al., 2009). Guh et al. (2015) localised SLC26 genes apically in gill 68 ionocytes, called SLC26 cells, but to our knowledge nothing is known about SLC26 localisation in the fish 69 kidney. In D. labrax, the only record about SLC26A6 (most probably SLC26A6c) is very recent and shows a 70 high mRNA expression in the anterior intestine to promote Cl⁻ transpithelial transport to the blood and HCO₃⁻ 71 excretion (Alves et al., 2019).

The basolateral Na⁺/K⁺-ATPase (NKA) (Hwang et al., 2011; Kumai and Perry, 2012) is a key player in Na⁺ and Cl⁻ uptake and its importance in FW acclimation has been demonstrated in many fish species (Hiroi et al., 2008; Inokuchi et al., 2008; Watanabe et al., 2008; Hsu et al., 2014; Bollinger et al., 2016). *Nka α1a* transcriptional expression in the posterior kidney was previously shown to be related to FW tolerance in European sea bass (L'Honoré et al., 2019). One other key cotransporter known for Na⁺ and Cl⁻ uptake in fish is the NCC-2A or NCC-like (SLC12A10 or SLC12A10.2), that plays a crucial role in FW acclimation in many fish species including European sea bass (Inokuchi et al., 2008; Wang et al., 2009; Hwang et al., 2011; Blondeau-Bidet et 79 al., 2019). SLC12A10 is localised apically in FW-type branchial ionocytes (i.e. NCC-type cells) in several 80 species (Hiroi et al., 2008; Inokuchi et al., 2008; Guh et al., 2015; Blondeau-Bidet et al., 2019). It is highly 81 expressed in FW compared to SW fish gills (Hiroi et al., 2008; Inokuchi et al., 2008; Wang et al., 2009; 82 Blondeau-Bidet et al., 2019). In posterior kidney, *slc12a10.1* paralog has been shown to be highly expressed in 83 zebrafish and in Mozambique tilapia gills compared to other organs (Hiroi et al., 2008; Wang et al., 2009). The 84 Na⁺/H⁺ exchanger-3 (NHE3) is also localised in the apical membrane of another ionocyte subtype at the gill 85 level (Inokuchi et al., 2008; Watanabe et al., 2008; Hwang et al., 2011; Blondeau-Bidet et al., 2019) and 86 functionally coupled to several other ion transporters facilitating Na⁺ uptake (Dymowska et al., 2015). Anion 87 exchanger 1 (AE1) is mainly known to play a role in bicarbonate transport to the blood and Cl⁻ secretion at the 88 gill and kidney (mammal medullary collecting duct cells) levels (Barone et al., 2004, Lee et al. 2011). AE1 is 89 thought to be functionally linked to apical V-H⁺-ATPase (VHA) in order to complete acid secretion. At the gill 90 level, AE1 is localised basolaterally in zebrafish HR cells (Lee et al., 2011). Its role was investigated in medaka 91 Oryzias latipes and in zebrafish gills (Lee et al., 2011; Hsu et al., 2014; Liu et al., 2016). In both species, the 92 two analysed paralogs (aela and aelb) were expressed in gills with aelb being over-expressed in a low-Na⁺ 93 environment. It has also been suggested that AE1 is involved in Cl⁻ absorption and HCO_3^- secretion (Evans et 94 al., 2005; Hwang and Lee, 2007; Hwang and Perry, 2010) but its localisation in basolateral membranes of 95 ionocytes in pufferfish Tetraodon nigroviridis and in milkfish Chanos chanos seems not in favour with this 96 hypothesis (Tang and Lee, 2007; Tang et al., 2011). In European sea bass, no data is available on VHA and AE1 97 localization. VHA mRNA expression and protein activity measurements have shown the presence of VHA in D. 98 labrax gills in FW media (Sinha et al., 2015; Blondeau-Bidet et al., 2019).

99 Regarding Na⁺/K⁺/2Cl⁻ cotransporters (NKCC), three paralogs have been described in teleosts: basolateral 100 NKCC1a mainly expressed in the gills and NKCC1b, both dedicated to NaCl secretion, and the apical NKCC2 101 mainly expressed in the kidney and intestine and attributed to NaCl reabsorption (Haas and Forbush III, 2000; 102 Teranishi et al., 2013). NKCC1 paralogs are expressed in SW-type ionocytes in numerous teleost species 103 including the European sea bass (Lorin-Nebel et al., 2006; Inokuchi et al., 2008; Buhariwalla et al., 2012; 104 Breves et al., 2014). NKCC2 was detected in tilapia and in *D. labrax* intestine (Hiroi et al., 2008; Alves et al., 105 2019), but there is no data available on its localisation in the gills and in the posterior kidney of *D. labrax*.

European sea bass exhibiting a FW intolerant phenotype face a severe hydromineral imbalance and we will analyse if this failure originates from a failure in ion uptake at the kidney and/or gill levels. The involvement of the gill in differential ion uptake capacity of *D. labrax* will be analysed by comparing the expression profile of the main ion transporters in tolerant and intolerant fish to FW. Little information is available about ion
transporter expression at the kidney level. This is a first tentative to identify key renal ion transporters in FW
kidney and to detect intraspecific differences in the transcriptional profile between tolerant and intolerant fish to
FW.

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114 Materials and methods

115 1. Tissue sampling

European sea bass juveniles (N=350) were reared at Ifremer Station at Palavas-les-flots (Hérault, France) in recirculating SW (osmolality: 1208 mOsm kg⁻¹, Na⁺: 515 mmol L⁻¹, Cl⁻: 737 mmol L⁻¹) under a 12/12 hours light/dark photoperiod at 20°C. At the age of 8 months (13.59 \pm 0.12 cm, 32.19 \pm 2.62 g), 300 fish were then transferred to brackish water (BW; osmolality: 475 mOsm kg⁻¹) for 24h and then transferred to dechlorinated tap FW (osmolality: 8 mOsm kg⁻¹, Na⁺: 2 mmol L⁻¹, Cl⁻: 3.5 mmol L⁻¹) for two weeks. The 50 remaining fish were transferred from SW to SW as controls. Pellet food (Le Gouessant, France) was proposed to fish daily, but the fish did not feed during the two weeks of FW challenges.

123 Fish were sampled two weeks following FW challenge. FW intolerant fish were identified according to their 124 behavior e.g. an atypical swimming (lower velocity sometimes associated with a darker color, isolation from the 125 shoal) as described previously (Nebel et al., 2005; L'Honoré et al., 2019). Fish that did not show any of these 126 traits were considered as FW tolerant fish. In this study we compared three groups: FW tolerant fish (FW_t, 127 N=15), FW intolerant fish (FW_i, N=15), and SW control fish (SW, N=13). After the 2-week FW challenge, fish 128 were killed by a lethal dose of benzocaine (at 100 ppm) and the first left gill arch and the posterior kidney (last 129 posterior third of the excretory kidney) were excised and immerged into RNAlater (Qiagen, Mississauga, ON, 130 Canada) for 24h at 4°C and then stored at -80°C for further analyses.

The experiments were conducted according to the guidelines of the European Union (directive 86/609) and of
the French law (decree 87/848) regulating animal experimentation. The experimental design has been approved
by the French legal requirement concerning welfare of experimental animals (APAFIS permit no. 9045201701068219555).

135 2. Gene expression

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2.1 RNA extraction and complementary DNA (cDNA) synthesis

Tissues were thawed on ice in lysis buffer using the total RNA extraction kit (Nucleospin® RNA, MachereyNagel, Germany) before performing the extraction. Quantity and purity (A260/280 ratio) of extracted RNA were
verified using a spectrophotometer (NanoDropTM One/OneC Spectrophotometer, Thermo Scientific, Waltham,
MA, USA). One microgram of RNA was used to generate the complementary DNA (cDNA) using the
qScriptTM cDNA SuperMix (Quanta BiosciencesTM) providing all necessary components for first-strand
synthesis: buffer, oligo(dT) primers, random primers and qScript reverse transcriptase.

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2.2 Quantification of gene expression levels

146 384-wells plates were filled with an Echo®525 liquid handling system (Labcyte Inc., San Jose, CA, USA). Each 147 well contained a mix composed by 0.75 µL of LightCycler-FastStart DNA Master SYBR-Green I[™] Mix 148 (Roche, Manheim, Germany), 0.037 µL of each primer (forward and reverse primers at 0.2 µM final 149 concentration), 0.21 µL of ultrapure water and 0.5 µL of cDNA. The dilution of the samples was determined 150 according to the standard curves generated for each primer pair. Quantitative real-time PCR (qRT-PCR) was 151 performed using a LightCycler®480 Real-Time PCR System (Roche, Mannheim, Baden-Württemberg, 152 Germany), with the conditions described in Blondeau-Bidet et al., 2019. All the primers used are listed in Table 153 1. For each organ and gene, efficiency was determined (ranged from 1.8 to 2.1) and was used for gene 154 expression quantification (see below) (Table 1). PCR products were sequenced by Sanger sequencing in order to 155 validate the identity of the amplified sequences in comparison with the sea bass genome. Elongation factor 1 α 156 $(efl\alpha)$ was chosen as a reference gene according to previous studies performed on European sea bass challenged 157 to FW (Nebel et al. 2005; Mitter et al. 2009; Blondeau-Bidet et al. 2016, L'Honoré et al., 2019). Relative 158 expression of the target gene was performed using the delta delta Ct method (Pfaffl, 2001) with the SW 159 condition as the reference condition. Ultra-pure water was used as a negative control template, and each sample 160 was analysed in duplicate.

161 3. Phylogenetic analysis and protein comparisons

Phylogenetic analyses of *slc26a6* and *nkcc (slc12a1 and slc12a2)* have been performed (Figs 1, 2). Nucleotide sequences from different species were obtained at Ensembl or NCBI (Tables 2, 3). *D. labrax* sequences were obtained from the European sea bass genome project (Tine et al., 2014). Multiple nucleotide alignments were performed with MUSCLE V3.8.31 (Edgar, 2004) and ambiguous regions were removed with Gblocks V0.91b

166 (Talavera and Castresana, 2007). The phylogenetic trees were inferred using the maximum likelihood (ML) 167 method implemented in the PhyML program (v3.1/3.0 aLRT) (Guindon and Gascuel, 2003). Best model of 168 evolution was selected using Modelgenerator V.85 (Keane et al., 2006) following the corrected Akaike 169 Information Criterion (with four discrete gamma categories) and used to construct a phylogenetic tree. Bayesian 170 posterior probabilities were computed with MrBayes 3.2.1 (Huelsenbeck and Ronquist, 2001; Ronquist and 171 Huelsenbeck, 2003). Two different runs with four incrementally heated simultaneous Monte Carlo Markov 172 chains were conducted over one million generations, applying respective substitution models determined for 173 each partition in PartitionFinder2 (Lanfear et al., 2017). Trees were sampled every 100 generations to produce 174 10,000 trees. In order to estimate posterior probabilities, 25% of the trees were discarded as a burn-in stage, 175 observing when average standard deviation of split frequency (ASDSF) values dropped below 0.01. Tree were 176 generated and robustness of the phylogeny assumption was evaluated by bootstrapping procedure from 1000 177 data set replicates and with posterior probabilities (PP). The phylogenetic tree of *slc26a6* was rooted with the 178 slc26a6 sequences of two reptilians, the Chinese soft-shelled turtle (Pelodiscus sinensis) and the painted turtle 179 (Chrysemys picta bellii). The phylogenetic tree of nkcc (slc12a1 and slc12a2) was rooted with the ncc3 180 sequences of the tropical frog (Xenopus tropicalis) and the lizard (Anolis carolinensis) (Hartmann et al., 2013).

Amino acid identities (in %) were performed with MUSCLE V3.8.31 (Edgar, 2004) implemented in Geneious®
9.1.8 software (Biomatters).

183 4. Statistical analysis

Analyses were performed using GraphPad Prism (version 6, GraphPad Software Incorporated, La Jolla, CA 268, USA). When parametric conditions were verified, one-way analyses of variances were performed followed by a Tukey's post-hoc test. In the case of non-parametric conditions, a Kruskal-Wallis test was performed followed by a Dunn's post-hoc test. Data are represented as box and whisker plots (from first quartile to third quartile) showing median, minimum and maximum values. Statistical differences were accepted from P < 0.05.

189 Results

- Among the 300 fish challenged in FW, 28% of them were detected and characterised as FW intolerant fish and
 70% as FW tolerant fish.
- 192 1. Phylogenetic analysis of European sea bass *slc26a6* and *nkcc* paralogous genes

193 Three paralogs of *slc26a6* have been identified in the European sea bass genome (Tine et al., 2014). The three 194 paralogs are called in this study *slc26a6 a*, *slc26a6 b* and *slc26a6 c* according to their phylogenetic position 195 (Fig. 1) computed by maximum likehood analysis against sequences of selected teleost fish (Table 2). 196 Comparatively, three paralogs were identified in O. niloticus, O. latipes, G. aculeatus, F. heteroclitus and in D. 197 rerio whereas two paralogs were identified in T. nigroviridis (a and c clades). The European sea bass slc26a6 198 paralogous genes obtained in this study strongly grouped with other teleosts slc26a6 paralogs. D. labrax 199 slc26a6c is grouped with slc26a6c of O. niloticus, O. latipes, T. nigroviridis and shared 86%, 80% and 56% 200 identity in amino acid sequences respectively. European sea bass SLC26A6c only shares 49% and 54% amino 201 acid identity with its SLC26A6b and SLC26A6a. Slc26a6b of D. labrax is grouped with O. niloticus slc26a6b, 202 sharing 80% identity in amino acid sequences but only 53% with European sea bass SLC26A6a. European sea 203 bass slc26a6b is strongly grouped with O. latipes and T. nigroviridis, sharing 71% and 67% identity in amino 204 acid sequences respectively.

205 Two *slc12a1* (*slc12a1a* and *slc12a1b*) and one *slc12a2* were identified in the European sea bass genome that 206 correspond to NKCC1a, NKCC1b and NKCC2 according to their phylogenetical position (Fig. 2), computed by 207 maximum likehood analysis against sequences of selected teleost fish (Table 3). Three nkcc genes were also 208 found in O. mossambicus, O. latipes, T. rubripes, G. aculeatus and G. morhua while two were found in T. 209 nigroviridis (1b and 2 clades). European sea bass nkcc paralogs strongly grouped with teleost nkcc sequences. 210 European sea bass *nkcc1a* and *nkcc1b* are grouped with *O. latipes nkcc1a* and *nkcc1b* and share 90.2% and 211 91.9% identity in amino acid sequences respectively. D. labrax NKCC1a and NKCC1b only share 77.8% 212 identity between each other. European sea bass nkcc2 is strongly grouped with T. nigroviridis and shares 82.5% 213 amino acid identity whereas it only shares 57.1 and 55.6% identity with NKCC1a and NKCC1b of European sea 214 bass amino acid sequences.

215

2. Relative gene expression in osmoregulatory tissues

Expression of selected genes was quantified in the gills and in the posterior kidney (Figs 3, 4, 5). Analysed
genes were: *ae1a* (*slc4a1-1*), *ae1b* (*slc4a1-2*), *nhe3* (*slc9a3*), *nkcc1a* (*slc12a2*), *nkcc1b* (*slc12a2-like*), *nkcc2*(*slc12a1*), *vha-b* (*atp6v1b*), *ncc1* (*slc12a3*), *ncc-2a* (*slc12a3-like*), *slc26a6a*, *slc26a6b*, *slc26a6c*, *prlra* and *prlrb*. Among them, *ae1b*, *slc26a6a*, *slc26a6b* and *nkcc2* expression levels have not been quantified in the gills. *slc12a2-like* (*nkcc1b*) expression in the gill was below the quantification limit and is therefore not shown in the

results. *ae1a* and *nhe3* expression levels have not been quantified in the posterior kidney because their
expression was below the quantification limit.

223 2.1 Gill gene expression

224 *ae1a* was not differentially expressed between SW and both FW groups (FW_t and FW_i, Kruskal-Wallis test, P =225 0.13, Fig. 3A). *nkcc1a* expression was significantly lower in both FW phenotypes (FW_t and FW_i) compared to 226 SW (Dunn's test, P < 0.0001 and P = 0.0142 respectively, Fig. 3B), but no significant differences were 227 measured between FW phenotypes.

228 *nhe3* relative expression showed no significant differences between SW, FW_t and FW_i. (Kruskal-Wallis test, P =229 0.1081, Fig. 3C) but a higher variability was observed in FW conditions. vha-b expression levels were 230 significantly higher in SW than in both FW phenotypes (FW_t and FW_i), but no differences were measured 231 between FW_t and FW_i (Dunn's test, P < 0.0001, P = 0.0007 and P = 0.9215 respectively, Fig. 3D). No 232 significant differences have been measured for ncc1 expression levels between the three groups (Kruskal-Wallis 233 test, P = 0.0626, Fig. 3E), but there is more variability in FW_i compared to SW and FW_t, with an almost 234 significant difference between FW_t and FW_i (Dunn's test, P = 0.0676). ncc-2a expression was significantly 235 higher in FW_t than in SW (Dunn's test, P = 0.0170, Fig. 3F), but no differences were measured between SW and 236 FW_i as between FW_t and FW_i (Dunn's test, P = 0.0562 and P > 0.9999 respectively). Among *slc26a6* paralogs, 237 only *slc26a6c* expression has been detected in the gills. FW_t exhibited a significantly lower expression of *slc26a6c* compared to both SW and FW_i (Dunn's test, P = 0.0309 and P = 0.0367, Fig. 3G) whereas no 238 239 difference was observed between SW and FW_i (Dunn's test, P > 0.9999).

Prolactin receptor *prlra* relative expression was significantly higher in FW_i compared to SW fish (Dunn's test *P* = 0.0001, Fig. 5A). No difference in *prlra* expression was recorded between SW and FW_t or between FW_i and FW_t (Dunn's test, *P* = 0.1324 and *P* = 0.0652 respectively) but a high variability is observed in FW_i. *Prlrb* expression was lower in FW groups compared to SW, but only significant between FW_t and SW fish (Dunn's test, *P* = 0.0045, Fig. 5B). No difference was observed between FW_i and FW_t.

245 2.2 Posterior kidney gene expression

No significant differences were measured in *ae1b* expression levels between SW and FW_t. However a significantly lower expression was measured in FW_i compared to SW but not to FW_t (Dunn's test, P = 0.0261and P = 0.8787, Fig. 4A). 249 No significant differences were measured in *ncc1* and *nkcc1a* expression levels between the three groups 250 (Kruskal-Wallis test, P = 0.1016 and P = 0.2475 respectively, Fig. 4B, C). Concerning *nkcc1b*, no differences 251 were measured between SW and FW_t (Dunn's test, P = 0.5448, Fig. 4D) but FW_i expression levels were 252 significantly lower than SW and almost significantly lower than FW_t (Dunn's test, P = 0.0043 and P = 0.0529253 respectively). nkcc2 expression was not significantly different between SW and FWt or between FWt and FWi, 254 but it was significantly lower in FW_i than in SW (Dunn's test, P = 0.2269, P = 0.8044 and P = 0.0096255 respectively, Fig. 4E). In FW₁, among *slc12a* cotransporters analysed, *ncc1* is the most expressed paralog in the 256 posterior kidney followed by the 5-fold less expressed *nkcc2* and the 25- and 50-fold less expressed *nkcc1a* and 257 *nkcc1b* (not shown).

Concerning *slc26a6* paralogs, *slc26a6a* exhibited a significant lower relative expression in FW_t compared to SW (Dunn's test, P = 0.0044, Fig. 4F). In FW_i, *slc26a6a* expression was not significantly different from FW_t, whereas for *slc26a6b* and *slc26a6c*, FW_t were at significant higher levels than FW_i (Dunn's test, P = 0.0141 and Tukey's test, P = 0.0018, Fig. 4F, G, H) but not significantly different from SW. Overall, in FW_t the three paralogs were similarly expressed (not shown).

prlra expression was significantly higher in FW_t than in SW (Tukey test, P < 0.0001, Fig. 5C) whereas *prlrb* expression levels did not differ between SW and FW_t (Tukey test, P = 0.8217, Fig. 5D). A significantly lower expression of both *prlr* paralogs (*prlra*, *prlrb*) was measured in FW_i compared to FW_t (Tukey test, P = 0.0139and P = 0.0472 respectively, Fig. 5C, D).

267 Discussion

268 The comparative analysis of key genes and proteins involved in the maintenance of hydromineral balance in fish 269 exhibiting different capacities to tolerate FW is a powerful tool to investigate intraspecific variation in FW 270 tolerance in euryhaline species. In FW, fish have to minimise ion loss and compensate through active ion uptake 271 occurring at interfaces with their surroundings. Most studies investigating FW osmoregulation focused at the gill 272 level in adults or at integument level in larvae (mostly zebrafish), showing that FW tolerance relies on the 273 expression of key genes involved in hyper-osmoregulatory mechanisms (Evans and Clairborne, 2009). Less data 274 are available on the involvement and importance of the kidney in ion re-uptake. In this study, we highlight the 275 importance of taking into account not only the gill but also the posterior kidney in order to fully understand the 276 mechanisms of ion and acid-base regulation.

277 1. Ion uptake capacities are not altered in gills of FW intolerant sea bass

278 Regarding gill ion uptake, slightly higher $nka \alpha la$ expression levels and NKA protein activity were previously 279 measured in FW_i compared to FW_t in European sea bass (Nebel et al., 2005; L'Honoré et al., 2019). This could 280 indicate a compensatory response to the blood ion imbalance. In fact, several studies on F. heteroclitus or D. 281 *labrax* suggested that a higher Na⁺/Cl⁻ ratio may indicate a more alkaline state (Scott, 2004; Scott et al., 2004a; 282 L'Honoré et al., 2019). In D. labrax, the FW_i phenotype was in this physiological condition (L'Honoré et al., 283 2019) due to a high Na⁺/Cl⁻ ratio. Regarding diverse transporters, we measured no difference in *ae1a* (*slc4a1a*) 284 relative expression between SW and FW phenotypes in gills but a higher variability in FW_i (Fig. 3A). AE1b has 285 been demonstrated as the dominant paralog in medaka gills and it was more expressed in FW media than in 286 saline waters from 10 to 30 % (Liu et al., 2016). This seems not to be the case in European sea bass, as we 287 could only quantify *ae1a*, which was not differently expressed between salinities. In medaka gills, an opposite 288 response was measured regarding the two AE1 paralogs, AE1a and AE1b, when comparing SW and FW-289 acclimated fish suggesting different functions according to the salinity. Using inhibitors and the scanning ion-290 selective electrode technique (SIET), the role of AE1 seemed to be attributed to acid and Cl⁻ secretion in 291 seawater-acclimated medaka larvae (Liu et al., 2016). In European sea bass, the physiological role of AE1 292 paralogs still needs to be investigated. Moreover, a different expression pattern is observed between *ae1a* and 293 vha-b, which is not consistent with a functional link between apical VHA and basolateral AE1a as shown in 294 medaka. The slightly higher aela expression in FW_i compared to FW_t is therefore probably not linked to an 295 increased acid secretion by VHA. A 3-fold lower expression of nkcc1a was measured in FW compared to SW 296 acclimated fish gills (Fig. 3B), as it has previously been shown in this species (Lorin-Nebel et al. 2006; 297 Blondeau-Bidet et al., 2019) and many other teleosts (Hiroi et al., 2008; Inokuchi et al., 2008 Bollinger et al., 298 2016). This is consistent with the role played by basolateral NKCC1 in ion secretion. According to Inokuchi et 299 al. (2017) and Blondeau-Bidet et al. (2019), FW transfer in Japanese and European sea bass might be followed 300 by a migration towards gill lamellae of seawater-type ionocytes differentiating into FW-type ionocytes (NHE3-301 type cells) involved in Na⁺ uptake. The observed decrease of *nkcc1a* expression in FW_i suggests a successful 302 functional shift from SW to FW-type ionocytes. nhe3 seems also slightly more expressed in both FW 303 phenotypes compared to SW but unlike previous studies (Blondeau-Bidet et al., 2019), there is no significant 304 difference between salinities. Interestingly, the variability of expression levels for nhe3 is higher in FW_i 305 compared to the other conditions, which could indicate differential responses within this group.

307 We measured a higher expression of ncc-2a in FW acclimated European sea bass than in SW (Fig. 3F) as it was 308 already demonstrated in this species (Blondeau-Bidet et al., 2019). This reinforces the hypothesis of the 309 differentiation of *ncc-2a* expressing ionocytes in FW, as it was observed in many other fish species such as 310 medaka and tilapia, exhibiting an higher relative expression of ncc (homologous to European sea bass ncc-2a) in 311 FW than in SW conditions (Hiroi et al., 2008; Inokuchi et al., 2008 Bollinger et al., 2016). FW_i fish also 312 increase slightly but not significantly their ncc-2a expression suggesting that at least some individuals 313 successfully differentiate NCC-type cells in gill filaments as shown previously in this species (Blondeau-Bidet 314 et al., 2019). Interestingly, expression levels of ncc1 were slightly higher and extremely variable in FW_i 315 compared to the other groups (Fig. 3E), which was surprising as *ncc1* expression in the gills is generally low 316 (Inokuchi et al., 2017). A compensatory role of ncc1 in Na⁺, Cl⁻ uptake in intolerant fish gills could be 317 hypothesised as fish gills are known to have multiple ion uptake pathways, that compensate each other in case of 318 stressful and ion-poor environments (Hwang et al., 2011; Hwang and Lee, 2007).

319 *Slc26a6c* expression is significantly higher in SW and FW_i compared to FW_t (Fig. 3G). In SW, where fish do 320 not need to take up Cl⁻, the high *slc26a6c* expression is surprising and might be linked to other functions as an 321 increased need to excrete HCO3⁻. In fact, studies dealing with slc26a6c in fish mostly focus on FW 322 environments and/or FW species (Bayaa et al., 2009; Perry et al., 2009). In this study we show that this gene 323 might also be of importance in SW conditions. The high expression in FW_i indicates that these fish may 324 compensate their low blood Cl⁻ levels through an increased Cl⁻ absorption and also activate the secretion of 325 HCO_3^- to restore blood pH levels (L'Honoré et al., 2019). In zebrafish, a higher expression level of za6c was 326 observed in the gills in hypo-osmotic conditions which is consistent with an involvement of this ion exchanger 327 in chloride uptake in apical membranes (Bayaa et al., 2009; Perry et al., 2009). By displaying enhanced 328 expression levels of Cl⁻ uptake genes such as *ncc1* and *slc26a6c*, FW_i sea bass seemed to activate Cl⁻ absorption 329 at the branchial level to compensate low Cl⁻ circulating levels measured previously (L'Honoré et al., 2019).

330

2. FW intolerant sea bass differentially regulate chloride reabsorption by the kidney

In FW_i posterior kidney, lower expression of *nka* a1a were previously measured compared to FW_t, associated with lower Cl⁻ blood levels possibly due to a renal ion leakage (L'Honoré et al., 2019). In this study, we have measured a lower expression of two other transporters involved in acid-base and hydromineral regulations (*ae1b* and *slc26a6c*) in FW_i compared to FW_t suggesting that FW_i are actually less efficient to compensate ion loss at the kidney level. Contrary to zebrafish in which *ae1a* is the predominant paralog of AE1 in the kidney, in *D*. 336 labrax only aelb could be detected (Lee et al., 2011). Expression of aelb was low in FWi compared to the other 337 conditions. Its basolateral localisation has been demonstrated in goldfish (Fehsenfeld and Wood, 2018), 338 suggesting a role in acid-base regulation. FW_i may decrease the expression of this transporter in order to limit 339 excessive HCO₃⁻ levels in the blood as these fish already experience blood alkalosis (L'Honoré et al., 2019). In 340 D. labrax kidney, expression levels of ncc1 (slc12a3) are higher than other slc12a cotransporters, which suggest 341 that NCC1 is predominant in D. labrax kidney for ion uptake. It is slightly more expressed in FW_t compared to 342 SW fish and is highly variable in FW_i. ncc1 (slc12a3) has been demonstrated to be also highly expressed in FW 343 in Japanese sea bass kidney compared to SW (Inokuchi et al., 2017). In mefugu Takifugu obscurus, ncc 344 (homologous to European sea bass *ncc1*) is localised apically and highly expressed in collecting duct of FW-345 acclimated fish, with a putative role in Na⁺, Cl⁻ reabsorption (Kato et al., 2010). In killifish F. heteroclitus, 346 rainbow trout Oncorhynchus mykiss and torafugu Takifugu rubripes, NKCC2 was localised apically and a role 347 in Na⁺, Cl⁻ reabsorption was suggested (Katoh et al., 2008; Kato et al., 2010). Compared to ncc1, nkcc2 was 348 about 6.5 times less expressed in European sea bass kidney in FWt and it rather seems downregulated in FW 349 media vs SW. Its role in ion uptake seems therefore secondary. FWi sea bass exhibited the lowest expression 350 levels of *nkcc2* and are thus certainly not using this ion uptake pathway to compensate ion loss. The basolateral 351 nkcc1a and nkcc1b are respectively 15.8 and 27 times less expressed than ncc1 in FWt which is consistent with 352 the role of NKCC1 paralogs in ion excretion rather than ion absorption. Whereas *nkcc1a* expression did not 353 change between the different treatments and phenotypes, nkcc1b showed significantly lower expression in FW_i 354 compared to the other conditions. The physiological significance of this transcriptional down-regulation needs 355 to be further explored.

356 Among the three SLC26A6 paralogous genes investigated in the posterior kidney of European sea bass, 357 *slc26a6a* exhibited significant lower expression in FWt compared to SW: *slc26a6b* and *slc26a6c* expressions 358 were not significantly modified between SW and FW_t , but they were lower in FW_i . A higher *slc26a6c* 359 expression in low-Cl⁻ FW compared to normal FW has already been demonstrated in zebrafish gills and kidney 360 with a suggested role in HCO₃⁻ excretion associated with a Cl⁻ (re)absorption (Bayaa et al., 2009). SLC26A6 361 family can exchange various anions with Cl⁻ such as sulfate (SO_4^{2-}) , oxalate $([COO^-]_2)$ and bicarbonate (HCO_3^{-}) 362 (Mount and Romero, 2004). The role of SLC26A6a in SO₄²⁻/Cl⁻ exchange in mefugu proximal tubules has been 363 already highlighted. According to Kato et al. (2009), SLC26A6a is acting as the main apical SO_4^{2-}/Cl^{-} exchanger 364 and its expression is enhanced under SW conditions. Our results are concordant with this study since both FW 365 sea bass phenotypes (FW_t and FW_i) exhibited lower *slc26a6a* expression levels than in SW. Less is known

about the role of SLC26A6b in the kidney. SLC26A6a and SLC26A6b have been shown to be both implicated in HCO_3^- secretion by the intestine (Xie et al., 2002) but their role in SO_4^{2-} or/and HCO_3^- secretion needs to be further investigated in the kidney of European sea bass and other teleost species.

369 3. Prolactin receptor transcript levels differ according to FW tolerance capacities

370 Contrary to mammals where only one *prlr* gene exists, teleost can have two distinct paralogs due to genome 371 duplication events (Ocampo Daza and Larhammar, 2018). The relationship between PRLR and gene expression 372 patterns of osmoregulatory-related genes has not yet been described in European sea bass and needs more 373 attention. Gill *prlr* paralog expression levels seem to depend on circulating PRL levels as shown in Seale et al. 374 (2012) for prlra and on extracellular osmolality as shown for prlrb in Fiol et al. (2009). Pituitary PRL acts as 375 one of the major endocrine signals for FW acclimation in euryhaline teleosts (Manzon, 2002). Lower prlra 376 expression levels in SW compared to FW have been already shown in Mozambique tilapia (Breves et al., 2010; 377 Breves et al., 2011), whereas no differences where observed concerning branchial prlrb relative expression 378 between salinities. In our study, prlr paralogs expression patterns are not concordant with those described in tilapia gills. Even if *prlra* seems to be slightly more expressed in FW_t than in SW, *prlrb* is significantly 379 380 overexpressed in SW compared to FW_t. We have previously shown that FW_i exhibited decreased blood 381 osmolality levels (L'Honoré et al., 2019), which could be one of the trigger to activate branchial *prlr* expression. 382 In fact, we measured slightly but not significantly higher mRNA levels of both *prlr* paralogs in FW_i compared 383 to FW_t. In zebrafish and Nile tilapia, PRL and PRLR positively regulate the expression of *ncc* in gill filament 384 cultures in vitro (Breves et al., 2013, 2014), which suggests that NCC-type cells might express PRLR. In our 385 study, we also highlighted higher branchial ncc1 relative expression in FW_i compared to FW_i, suggesting a 386 similar regulation of *ncc1* expression levels by *prlr*. Moreover, slightly higher *nka* αla expression levels and 387 increased gill NKA activity were previously measured in FW_i compared to FW_t (Nebel et al., 2005; L'Honoré et 388 al., 2019), potentially linked to an increased number of ionocytes to compensate low blood osmolality and low 389 circulating chloride levels.

In the posterior kidney, *prlra* is the predominantly expressed paralog (10-fold higher expressed than *prlrb*) whereas in the gills, *prlrb* was 10-fold more expressed than *prlra*. No differences in *prlra* relative expression have been reported following salinity change in Seale et al. (2012) but a lower relative expression of *prlrb* was shown in FW compared to SW (Fiol et al., 2009; Breves et al., 2011). It seems different in European sea bass posterior kidney where significantly higher expression levels of *prlra* are measured in FW_t compared to SW. 395 Moreover, no differences were observed for *prlrb* between SW and FW_t. Interestingly, both *prlra* and *prlrb* 396 were less expressed in FW_i compared to FW_t, highlighting that FW_i and FW_t do not exhibit the same regulation 397 mechanisms in response to the FW challenge. We showed a differential expression of prolactin receptors in FW 398 between FW_t and FW_i, with a higher, though not significant, over-expression of branchial prolactin receptors in 399 FW_i and a significantly lower expression of both renal prolactin receptors. These results suggest that FW 400 intolerance in European sea bass might be due to an altered PRL-associated endocrine response in FW_i.

401 Conclusion

402 This study is the first to highlight that freshwater intolerance in European sea bass is linked with lower mRNA 403 expression of *slc26a6* genes involved in Cl⁻ uptake in the posterior kidney. Regarding prolactin receptors, we 404 showed a differential endocrine control in FW between the tolerant and intolerant European sea bass associated 405 to the incapacity to maintain blood hydromineral balance in FW_i sea bass. Facing strong rainfalls during their 406 migration to transitional habitats like lagoons, European sea bass may undergo strong salinity decreases 407 triggering mortality in the freshwater intolerant phenotype. Thus, freshwater intolerance could affect some 408 European sea bass in their migrations by preventing them to access transitional environments. It remains to be 409 characterised how freshwater intolerance in D. labrax affects seasonal migratory behaviour in European sea 410 bass.

411

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416 Conflict of interest

417 The authors declare that there is no conflict of interest regarding the publication of this manuscript.

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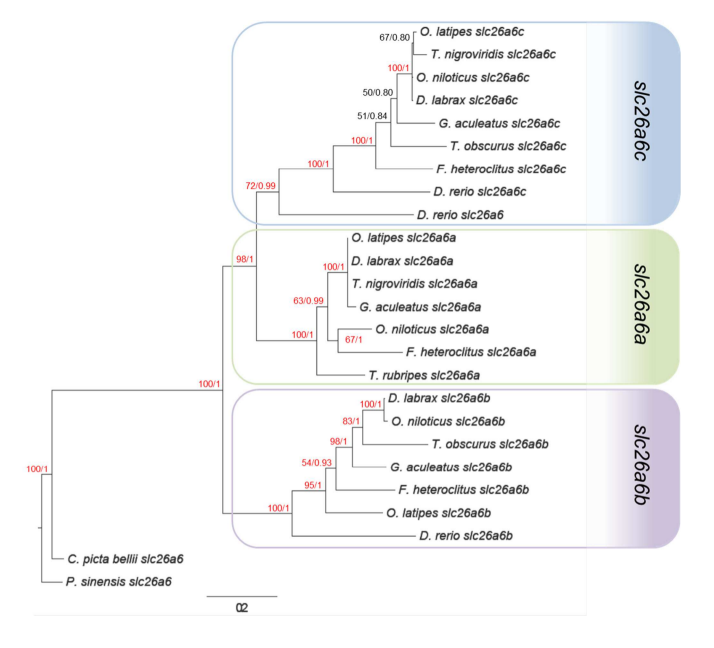


Fig. 1 Bayesian tree of *slc26a6*. A ML tree had identical topology for all nodes. Bootstrap values (in %) from the ML are shown before the Bayesian PP, confident nodes are indicated in red. Branch lengths represent the degree of divergence, with the scale bar indicating the distance representing 0.2 substitutions per position. The blue frame mainly comprises sequences of teleost *slc26a6a* and purple frame comprises sequences of teleost *slc26a6b*.

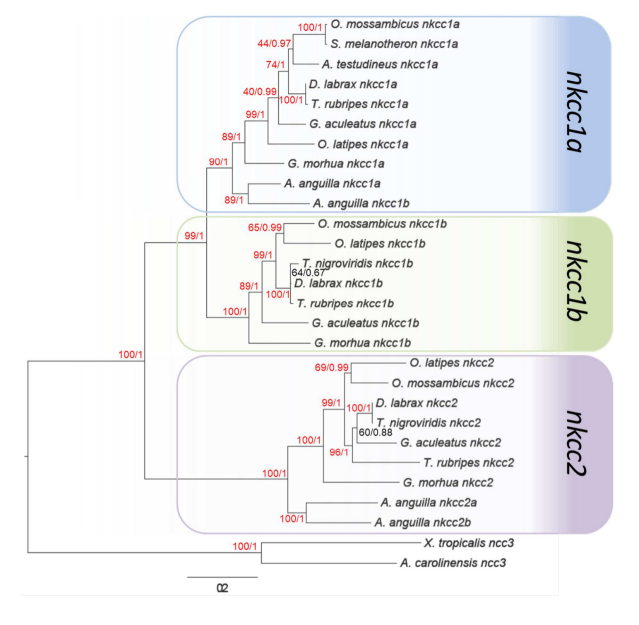


Fig. 2 Bayesian tree of *nkcc1a*, *nkcc1b* and *nkcc2* (*slc12a1a*, *slc12a1b* and *slc12a2*). A ML tree had identical topology for all nodes. Bootstrap values (in %) from the ML are shown before the Bayesian PP, confident nodes are indicated in red. Branch lengths represent the degree of divergence, with the scale bar indicating the distance representing 0.2 substitutions per position. The blue frame mainly comprises sequences of teleost *nkcc1a*. The green frame mainly comprises sequences of teleost *nkcc1b* and purple frame comprises sequences of teleost *nkcc2*.

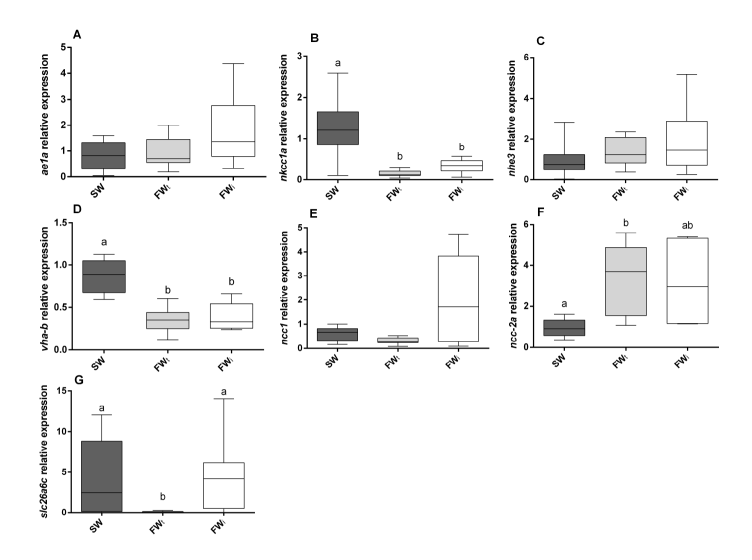


Fig. 3 Relative expression in gills of (A) ae1a / slc4a1-1, (B) nkcc1a / slc12a2-1, (C) nhe3, (D) vha-b, (E) ncc1, (F) ncc-2a and (G) slc26a6c in SW and after two weeks in FW. The expression has been normalised according to the expression of the elongation factor ef1a. Different letters denote significant differences between groups (one-way Anova followed by Tukey's test or Kruskal-Wallis followed by Dunn's test, P < 0.05, means \pm s.e.m, N=6-16). SW: control fish in SW, FW_i: intolerant fish to FW, FW_i: tolerant fish to FW.

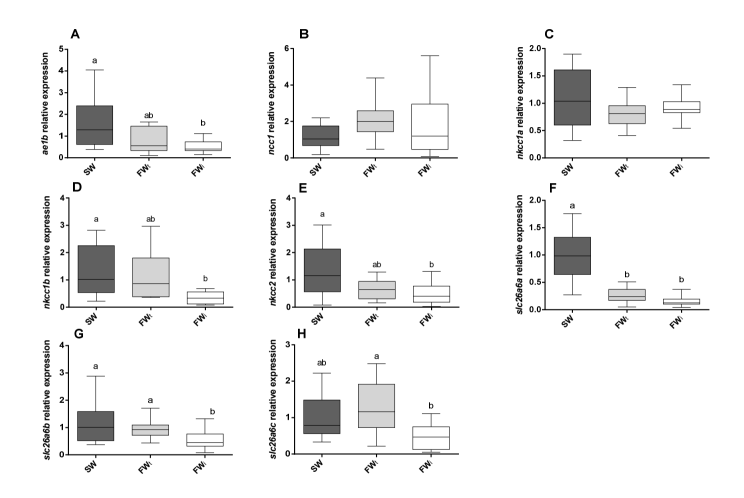


Fig. 4 Relative expression in posterior kidney of (A) *ael b / slc4a1-2*, (B) *ncc1 (slc12a3)*, (C) *nkcc1a / slc12a2*, (D) *nkcc1b / slc12a2-like*, (E) *nkcc2 (slc12a1)*, (F) *slc26a6a*, (G) *slc26a6b* and (H) *slc26a6c* in SW and after two weeks in FW. The expression has been normalised according to the expression of the elongation factor *ef1a*. Different letters denote significant differences between groups (one-way Anova followed by Tukey's test of Kruskal-Wallis followed by Dunn's test, P < 0.05, means ± s.e.m, N=6-16). SW: control fish in SW, FW_i: intolerant fish to FW, FW_i: tolerant fish to FW.

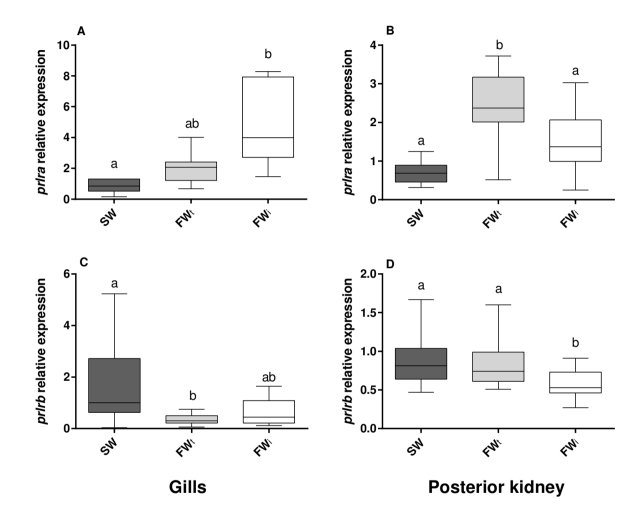


Fig. 5 Relative expression in the gills and in the posterior kidney of *prlr a* (A-C) and *prlr b* (B-D) in SW and after two weeks in FW. The expression has been normalised according to the expression of the elongation factor *ef1a*. Different letters denote significant differences between groups (one-way Anova followed by Tukey's test or Kruskal-Wallis test followed by Dunn's test, P < 0.05, means ± s.e.m, N=6-16). SW: control fish in SW, FW_i: intolerant fish to FW, FW_i: tolerant fish to FW.

Table 1 Primer sequences used for the gene expression analysis

Target gene	Primer name	Sequences ID	Sequence (from 5' to 3')	Efficiency	Reference
eflα	EF1a F	AJ866727.1	GGCTGGTATCTCTAAGAACG	1.9 (gills)	Nebel et al., 2005
	EF1a R		CCTCCAGCATGTTGTCTCC	1.9 (posterior kidney)	
nkcc1a	NKCC1A F	DLAgn_00123120	AGTTGGCAGTAAGGAGGTGG	2.1 (gills)	Blondeau-Bidet et al., 2019
	NKCC1A R		TCAGACTCAGAGGAGACTTGG	1.9 (posterior kidney)	
nkcc1b	NKCC1B F	DLAgn_00080120	TCAGCTCACAGTTCAAGGCC		This study
	NKCC1B R		TTGTGGAGTCCATAGCGGC	2.1 (posterior kidney)	
nkcc2	NKCC2 F	DLA_LG5_005810	GACACTGTGGAGGACGATGG		This study
	NKCC2 R		AGCATGCATCTCACCAGGAC	2.0 (posterior kidney)	
nhe3	NHE3 F	DLAgn_00204050	GGATACCTCGCCTACCTGAC	1.9 (gills)	Blondeau-Bidet et al., 2019
	NHE3 R		AAGAGGAGGGTGAGGAGGAT	1.9 (posterior kidney)	
ncc1	NCC1 F	DLAgn_00172790	TGACGTACTTGATCGCTGCC	1.9 (gills)	This study
	NCC1 R		AGTTGGTGATGGAGGCATGG	2,0 (posterior kidney)	
ncc2-a	NCC2-A F	DLAgn_00038210	ATGATGAGCCTCTTCGAGCC	2.1 (gills)	Blondeau-Bidet et al., 2019

	NCC2-A R		ACAGAAGGTGATGAGAGCAGC	2.0 (posterior kidney)	
slc26a6a	26A6-A F	DLAgn_00129890	TCATGTGTGTGTCTCCCAGC		This study
	26A6-A R		GAGAGTGCTACCAGGCTGAC	2.0 (posterior kidney)	
slc26a6b	26A6-B F	DLAgn_00133440	ATACCTGTGGAGCTGCTTGG		This study
	26A6-B R		TCAGGGCAAAGATTCGTCCC	1.9 (posterior kidney)	
slc26a6c	26A6-C F	DLAgn_00221070	GAAACGGACACAGAGGAGGG	1.9 (gills)	This study
	26A6-C R		GTCTGGTGTTCGAGGAGTGG	2.0 (posterior kidney)	
aela	AE1-A F	DLAgn_00101870	TCTGAAGGAATCGGTGGTGC	1.9 (gills)	This study
	AE1-A R		CTCGTTCTGGATCTCGGTGG		
aelb	AE1-B F	DLAgn_00193420	TGAACAAGGGTGAGATCCGC		This study
	AE1-B R		ACAAAGCGAATAGGGACGGG	2.0 (posterior kidney)	
prlr a	PRLR-A F	DLA_LG19_005350	GGGACAGAGGCAGAAGACAT	2.0 (gills)	This study
	PRLR-A R		GAGGTAGGAGGATGTGGAGC	2.0 (posterior kidney)	
prlr b	PRLR-B F	DLA_LG20_006210	GAATGACAATGGGCCTCTGC	2.0 (gills)	This study
	PRLR-B R		GTTCCGTTGCTTCTGGGTTT	2.0 (posterior kidney)	

Nucleotide sequence ID	Fish species	Gene
XM_024104973.1	Chrysemys picta bellii	slc26a6
DLAgn_00129890	Dicentrarchus labrax	slc26a6 a
DLAgn_00133440	Dicentrarchus labrax	slc26a6 b
DLAgn_00221070	Dicentrarchus labrax	slc26a6 c
XM_680900.6	Danio rerio	slc26a6
XM_001344207.7	Danio rerio	slc26a6 b
FJ170818.1	Danio rerio	slc26a6 c
XM_012876467.2	Fundulus heteroclitus	slc26a6 a
XM_021318616.1	Fundulus heteroclitus	slc26a6 b
XM_012872750.2	Fundulus heteroclitus	slc26a6 c
ENSGACP0000002987	Gasterosteus aculeatus	slc26a6
ENSGACT00000007648.1	Gasterosteus aculeatus	slc26a6 b
ENSGACT00000001978.1	Gasterosteus aculeatus	slc26a6 c

Table 2 Cl⁻/HCO₃⁻ exchanger (slc26a6) nucleotide sequences used in the phylogenetic analysis

ENSORLP00000011507	Oryzia latipes	slc26a6 a
XM_011476835.3	Oryzia latipes	slc26a6 b
ENSORLP0000002390	Oryzia latipes	slc26a6 c
ENSONIT00000015705.1	Oreochromis niloticus	slc26a6 a
ENSONIT0000002151	Oreochromis niloticus	slc26a6 b
ENSONIP00000018142	Oreochromis niloticus	slc26a6 c
XM_014577432.2	Pelodiscus sinensis	slc26a6
CAF95115.1	Tetraodon nigroviridis	slc26a6 a
CAG06912.1	Tetraodon nigroviridis	slc26a6 c
AB200328.1	Takifugu rubripes	slc26a6 a
AB200329.1	Takifugu rubripes	slc26a6 b
AB200330.1	Takifugu rubripes	slc26a6 c

Table 3 NKCC 1, 2 (*slc12a1/slc12a2*) nucleotide sequences used in the phylogenetic analysis

Nucleotide sequence ID	Fish species	Gene
JN180944.1	Anabas testudineus	nkccla
AJ486858.1	Anguilla anguilla	nkcc1a
AJ486859.1	Anguilla anguilla	nkcc1b
AJ564602.1	Anguilla anguilla	nkcc2a
AJ564603.1	Anguilla anguilla	nkcc2b
XM_003223867.1	Anolis carolinensis	ncc3
DLAgn_00080120	Dicentrarchus labrax	nkcc1a
DLAgn_00123120	Dicentrarchus labrax	nkcc1b
DLA_LG5_005810	Dicentrarchus labrax	nkcc2
ENSGMOT00000004750.1	Gadus morhua	nkcc1a
ENSGMOT0000001772.1	Gadus morhua	nkcc1b
ENSGMOT0000009346.1	Gadus morhua	nkcc2b
ENSGACT00000024304.1	Gasterosteus aculeatus	nkcc1a
ENSGACT00000019494.1	Gasterosteus aculeatus	nkcc1b

ENSGACT00000022179.1	Gasterosteus aculeatus	nkcc2
AY513737	Oreochromis mossambicus	nkcc1a
AY513738	Oreochromis mossambicus	nkcc1b
AY513739.1	Oreochromis mossambicus	nkcc2
ENSORLT00000024013.2	Oryzias latipes	nkcc1a
ENSORLT00000021359.2	Oryzias latipes	nkcc1b
ENSORLT00000038800.1	Oryzias latipes	nkcc2
GU066877.1	Sarotherodon melanotheron	nkcc1
ENSTRUP0000004453	Takifugu rubripes	nkcc1a
ENSTRUP00000027260	Takifugu rubripes	nkcc1b
ENSTRUT00000043584.2	Takifugu rubripes	nkcc2
CAG09669.1	Tetraodon nigroviridis	nkcc1b
CAF99849.1	Tetraodon nigroviridis	nkcc2
XM_002934280.4	Xenopus tropicalis	ncc3