

1 Mini-Review

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3 **Endocrine and osmoregulatory responses to tidally-changing salinities in fishes**

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32 **Abstract**

33 Salinity is one of the main physical properties that govern the distribution of fishes across  
34 aquatic habitats. In order to maintain their body fluids near osmotic set points in the face of  
35 salinity changes, euryhaline fishes rely upon tissue-level osmotically-induced responses and  
36 systemic endocrine signaling to direct adaptive ion-transport processes in the gill and other  
37 critical osmoregulatory organs. Some euryhaline teleosts inhabit tidally influenced waters such  
38 as estuaries where salinity can vary between fresh water (FW) and seawater (SW). The  
39 physiological adaptations that underlie euryhalinity in teleosts have been traditionally identified  
40 in fish held under steady-state conditions or following unidirectional transfers between FW and  
41 SW. Far fewer studies have employed salinity regimes that simulate the tidal cycles that some  
42 euryhaline fishes may experience in their native habitats. With an emphasis on prolactin (Prl)  
43 signaling and branchial ionocytes, this mini-review contrasts the physiological responses  
44 between euryhaline fish responding to tidal versus unidirectional changes in salinity. Three  
45 patterns that emerged from studying Mozambique tilapia (*Oreochromis mossambicus*) subjected  
46 to tidally-changing salinities include, 1) fish can compensate for continuous and marked changes  
47 in external salinity to maintain osmoregulatory parameters within narrow ranges, 2) tilapia  
48 maintain branchial ionocyte populations in a fashion similar to SW-acclimated fish, and 3) there  
49 is a shift from systemic to local modulation of Prl signaling.

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## 63 **1. Introduction**

64 Fishes are found across a wide range of aquatic environments where physical and  
65 chemical conditions shape their distribution and life history strategies. For instance, fish are  
66 distributed across environments with salinities ranging from freshwater (FW) (e.g., rivers, lakes,  
67 and marshes) to seawater (SW) (e.g., coastal waterways, bays, and the pelagic ocean). Further,  
68 fishes may inhabit extreme salinity environments such as hypersaline ponds and estuaries that  
69 undergo large tidal variations (Brauner et al., 2013). Canonically, euryhaline fish inhabit  
70 environments subject to wide salinity changes, though the extent of their tolerance to these  
71 changes is variable and largely dependent on the native environment and life history of the  
72 animal. Physiologically, euryhaline fishes (<10% of teleosts) can tolerate salinities ranging from  
73 FW ( $\leq 0.5$  ppt) to SW (30-40 ppt) through their capacity to maintain salt and water balance (Brill  
74 et al., 2001; Evans and Claiborne, 2008; Schultz and McCormick, 2013). When residing in FW,  
75 fish produce large volumes of dilute urine while actively absorbing environmental ions to  
76 counterbalance their loss by diffusion. Alternatively, marine/SW-acclimated fish drink to replace  
77 (via solute-linked water absorption in the gut) the water that is lost by osmosis and actively  
78 extrude excess ions (Evans et al., 2005). The maintenance of hydromineral balance is largely  
79 achieved through the major osmoregulatory organs, namely the gill, intestine, and kidney, which  
80 respond to osmosensory and endocrine stimuli. The myriad adaptive responses that fishes from  
81 various clades exhibit in response to changes in environmental salinity have been thoroughly  
82 reviewed (Fiol and Kültz, 2007; Gonzalez, 2012; Marshall, 2012; McCormick and Bradshaw,  
83 2006; Sakamoto and McCormick, 2006; Schulte, 2014). Studies which utilized euryhaline  
84 models to improve the collective understanding of how fish acclimate to environmental salinity  
85 have traditionally described the physiological states of animals acclimated to two or more  
86 steady-state salinities, or subjected to one way-transfers. There is no question these general  
87 paradigms will continue to provide the framework for understanding the physiological  
88 mechanisms that underlie environmental adaptation. Nonetheless, a need for employing  
89 experimental designs that address the highly-dynamic nature of particular aquatic environments  
90 has emerged in order to better understand the impacts of climate change and urbanization  
91 (Blewett et al., 2022). In turn, this mini-review focuses on findings that resulted from the use of  
92 experimental paradigms where physiological responses were described in fish subjected to

93 tidally-changing salinity regimes that simulate the periodicity of salinity fluctuations in their  
94 native environments.

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## 96 **2. The influence of salinity acclimation history**

97       Very few euryhaline fishes have been studied under tidally-changing salinity regimes. To  
98 our knowledge, studies investigating the physiological responses of euryhaline fish under a  
99 simulated tidal regime have been limited to the mummichog (*Fundulus heteroclitus*) and the  
100 Mozambique tilapia (*Oreochromis mossambicus*), which are known to withstand marked and  
101 frequent changes in salinity. Mummichogs are native to shallow coastal waters including  
102 estuaries and salt marshes of the western Atlantic Ocean between the Gulf of St. Lawrence and  
103 northeastern Florida (Robins et al., 1986). The Mozambique tilapia is native to estuaries and the  
104 lower reaches (within a mile from the tidal ebb and flow) of rivers from the Zambezi River to the  
105 southeast coast of South Africa (Trewavas, 1983). Thus, in certain locations, it is probable that  
106 both species are continuously exposed to changing salinities. In part because many acclimation  
107 studies have focused on the physiological responses following one-way transfers between  
108 salinities, a lingering question remains: to what extent do fish actually undergo physiological  
109 transformations/reversals when exposed to salinity challenges in periodic fashions? One  
110 approach to address this question is to consider whether a fish that experiences a ‘one-time’  
111 change in salinity from FW to SW, or vice-versa, responds in a similar manner as an individual  
112 previously, and periodically, exposed to both salinities. Survival, the ultimate indication of  
113 acclimation success, provides a clue to understanding the effects that acclimation history imparts  
114 on future salinity challenges.

115       In euryhaline Japanese medaka (*Oryzias latipes*), pre-adaptation to elevated salinity  
116 facilitated subsequent acclimation to SW (Miyaniishi et al., 2016). FW-acclimated fish were first  
117 pre-acclimated to 50% SW and then returned to FW. Twenty-five percent of the fish that had  
118 been pre-acclimated to 50% SW survived a direct transfer from FW to SW, while fish that were  
119 not previously exposed to 50% SW did not survive transfer to SW. Likewise, Mozambique  
120 tilapia, which can eventually be acclimated to salinities far exceeding those of SW (Uchida et al.,  
121 2000), fail to survive a direct transfer from FW to SW unless they are transitioned over an  
122 extended period, pre-acclimated to an intermediate salinity, or pre-exposed to a regime of  
123 changing salinities (Inokuchi et al., 2021; Moorman et al., 2015; Seale et al., 2002). The

124 congeneric Nile tilapia (*O. niloticus*) is less tolerant of elevations in salinity, as evidenced by its  
125 inability to survive direct transfer from FW to brackish water (BW; 20 ppt) for more than 24 h  
126 (Yamaguchi et al., 2018). The more narrow salinity tolerance of Nile tilapia is reflected by a  
127 sharp and uncontrolled rise in plasma osmolality following transfer from FW to BW (Yamaguchi  
128 et al., 2018). Plasma osmolality, a measure of the amount of solutes per kg of solution, largely  
129 reflects the concentrations of Na<sup>+</sup> and Cl<sup>-</sup>, which are also the principal ionic constituents of  
130 environmental water and are actively regulated through epithelial cells for maintaining  
131 organismal salt and water balance. It stands unresolved whether the ability of *O. niloticus* to  
132 tolerate BW/SW conditions can be improved with pre-exposure to sublethal periodic changes in  
133 salinity.

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### 135 **3. Osmotic homeostasis in tidally-changing salinities**

136         Studies with the mummichog on the effects of acute salinity transfers on transepithelial  
137 potential (TEP) indicated the prevalence of Na<sup>+</sup> diffusion in SW-acclimated fish and a limitation  
138 of Na<sup>+</sup> loss in FW-acclimated fish (Wood and Grosell, 2008). Interestingly, when fish were  
139 tested under conditions simulating a tidal cycle their TEP was distinct from fish acclimated to  
140 FW and more closely resembled fish acclimated to SW (Wood and Grosell, 2009). Nonetheless,  
141 when mummichogs transiently move into lower salinities they can do so without large  
142 disruptions in osmotic homeostasis (Wood and Grosell, 2009). By tracking ion fluxes in response  
143 to salinity changes, these studies also provided insight into how transient shifts in salinity can be  
144 tolerated with the least metabolic cost and with minimal impact on the configuration of  
145 physiological processes required for osmotic homeostasis.

146         As observed in mummichogs, Mozambique tilapia can also maintain osmotic  
147 homeostasis under tidally-changing salinities. Despite the wide range of salinities in which the  
148 Mozambique tilapia is found, it regulates plasma osmolality within a narrow range, typically  
149 around 320 mOsm/kg in FW-acclimated fish and 335 mOsm/kg in SW-acclimated fish (Seale et  
150 al., 2006a; Seale et al., 2002; Yada et al., 1994). However, following a one-way transfer from  
151 SW to FW, plasma osmolality may briefly reach as low as 280 mOsm/kg, while after a change  
152 from FW to 80% SW, it may climb to over 400 mOsm/kg for a short period (Breves et al., 2011;  
153 Seale et al., 2006a; Seale et al., 2012a; Seale et al., 2002; Yada et al., 1994). Most previous work  
154 investigating the links between salinity tolerance and key physiological processes such as

155 osmoregulation and growth in Mozambique tilapia has focused on fish reared under steady-state  
156 salinity conditions (FW or SW) or transferred from FW to SW and vice-versa (Borski et al.,  
157 1994; Breves et al., 2010a; Riley et al., 2003; Seale et al., 2006a; Seale et al., 2002). In  
158 Mozambique tilapia, a tidal regime (TR) paradigm was first tested in fish that were continuously  
159 reared in salinities alternating between FW and SW every 6 h from the early fry stages until 4  
160 months of age (Fig. 1A) (Moorman et al., 2014). A tidally-changing environment was simulated  
161 by maintaining fish in aquaria supplied with water from reservoirs containing FW or SW  
162 equipped with submersible pumps hooked to automatic timers offset by 6 h (Fig. 1B). Studies  
163 employing the TR paradigm allowed for a comparison between the FW- and SW-phases of the  
164 tidal cycle (TF and TS, respectively) relative to comparisons between FW and SW steady-state  
165 regimes (Fig. 1A). In contrast to the wide excursions in plasma osmolality typically observed by  
166 6 h after a ‘one-time’ transfer from FW to SW, tilapia reared under a TR showed a minor  
167 elevation of plasma osmolality in TS (Fig. 2) (Moorman et al., 2014; Moorman et al., 2015;  
168 Pavlosky et al., 2019; Seale et al., 2019). In fact, changes in osmolality were within the range  
169 found in fish maintained in steady-state FW or SW (Fig. 2). The capacity to prevent wide shifts  
170 in plasma osmolality under a TR suggests that tidal fish are particularly effective in maintaining  
171 osmoregulatory homeostasis.

172         The pathways that connect changes in environmental salinity with adaptive physiological  
173 responses involve signaling mechanisms that couple the perception of osmotic stimuli with the  
174 molecular and cellular processes central to the maintenance of osmotic balance. These signaling  
175 mechanisms, which involve direct osmosensing, osmoreceptors, and the endocrine control of ion  
176 and water transport within osmoregulatory epithelia, have been thoroughly reviewed (Breves et  
177 al., 2014; Fiol and Kültz, 2007; Guh et al., 2015; Marshall, 2012; McCormick and Bradshaw,  
178 2006; Seale et al., 2005, 2006b; Seale et al., 2020a; Seale et al., 2012b) and synthesized into a  
179 framework that integrates multiple sensory and effector elements (Evans and Kültz, 2020; Kültz,  
180 2012). Here, these signaling mechanisms are discussed within the context of animals reared  
181 under tidal paradigms.

182

#### 183 **4. Hormonal responses to tidally-changing salinities**

184         In response to an osmotic challenge, euryhaline species integrate osmosensing, systemic  
185 and local endocrine signaling, and effector mechanisms. Aspects of these distinct, yet interacting,

186 processes are summarized in Fig. 3 and compared between Mozambique tilapia acclimated to  
187 steady-state salinities (FW and SW) and TR (TF and TS). At the organismal level, tidal changes  
188 in environmental salinity (Fig. 3A) elicited narrow oscillations in plasma osmolality, between  
189 about 320 and 345 mOsm/kg (Fig. 3B). At the level of the pituitary, systemic osmoreception in  
190 Mozambique tilapia occurs at least partially through prolactin (Prl), a pleiotropic hormone that is  
191 both highly osmosensitive and effective at directing hyperosmoregulatory processes (Grau and  
192 Helms, 1989; Seale et al., 2006b; Seale et al., 2012b). In Mozambique tilapia, the release of two  
193 Prl isoforms, Prl<sub>177</sub> and Prl<sub>188</sub>, increased in response to falls in extracellular osmolality *in vitro* or  
194 following an *in vivo* transfer from SW to FW (Grau et al., 1981; Nicoll et al., 1981; Seale et al.,  
195 2012a; Seale et al., 2002; Yada et al., 1994). One notable exception to the general pattern of an  
196 inverse relationship between plasma Prl and extracellular osmolality was observed in fish reared  
197 in a TR. The gene expression and release of both Prls became decoupled from extracellular  
198 osmolality; circulating Prl levels resembled those found in SW-acclimated fish (Fig. 3C and D)  
199 (Moorman et al., 2014; Moorman et al., 2015; Seale et al., 2019). These findings suggest that Prl  
200 release is less sensitive to ambient osmolality under a TR. However, if extracellular osmolality  
201 abruptly breaks below 320 mOsm/kg, plasma Prl levels will rise (Seale et al., 2019).  
202 Accordingly, Prl increased only after fish previously maintained under a TR had been transferred  
203 to FW for 24 h. This pattern of circulating Prl paralleled *prl* gene expression in the pituitary  
204 (Moorman et al., 2015). Collectively, these observations indicate that a point exists between 6  
205 and 24 h after FW-exposure when elevated plasma Prl becomes necessary to maintain osmotic  
206 balance.

207         Teleost Prls exert their actions by interacting with transmembrane receptors, two of  
208 which, Prl receptor 1 (Prlr1) and -2 (Prlr2), have been identified in tilapia (Fiol et al., 2009).  
209 Branchial gene expression of *prlr1* and -2 responds to both osmotic stimuli and circulating Prl.  
210 While branchial *prlr1* was stimulated by Prl<sub>177</sub> and Prl<sub>188</sub> yet unresponsive to extracellular  
211 osmolality, *prlr2* was unresponsive to both Prls and upregulated following an increase in  
212 osmolality (Inokuchi et al., 2015). In pituitary Prl cells, the expression patterns of both *prlrs* in  
213 relation to extracellular osmolality were similar to those observed in the gill, indicating that in  
214 addition to mediating Prl signaling in target tissues (Seale et al., 2012a), Prlrs also mediate  
215 osmotically-modulated autocrine responses in the pituitary (Yamaguchi et al., 2016). In contrast  
216 to the modest changes in the levels of *prl* transcripts and plasma Prls seen under a TR, the gene

217 expression of branchial *prlrs* changed during each phase of the tidal cycle. For instance, *prlr1*  
218 increased in TF and decreased in TS (Fig. 3E) (Moorman et al., 2014; Moorman et al., 2015;  
219 Seale et al., 2019). By contrast, *prlr2* responded in the opposite direction by increasing in TS and  
220 decreasing in TF. These observations led us to propose that under tidal conditions the endocrine  
221 control of osmoregulatory processes shifts from the systemic regulation of hormone secretion to  
222 the local modulation of hormonal action via changes in receptor expression (Seale et al., 2019).

223 Changes in salinity affect the release of other pleiotropic pituitary hormones such as  
224 growth hormone (Gh). Unlike Prl, however, Gh increased in response to elevations in salinity, *in*  
225 *vivo* (Helms et al., 1987; McCormick, 2001; Pierce et al., 2007; Sakamoto and Hirano, 1993;  
226 Sakamoto et al., 1997), and extracellular osmolality, *in vitro* (Borski et al., 1994; Seale et al.,  
227 2006a; Seale et al., 2002). These responses are consistent with the higher growth rates observed  
228 in tilapia reared in SW versus lower salinities (Kuwaye et al., 1993; Morgan and Iwama, 1991;  
229 Riley et al., 2002; Ron et al., 1995; Shepherd et al., 1997; Sparks et al., 2003). Moreover, Gh is  
230 also tied to branchial processes that support SW-acclimation (McCormick and Bradshaw, 2006).  
231 The actions of Gh on osmoregulatory systems is mediated, at least partially, by increased plasma  
232 insulin-like growth factor 1 (Igf1) levels (Mancera and McCormick, 1998; Sakamoto and  
233 McCormick, 2006; Seidelin and Madsen, 1999; Tipsmark et al., 2007) and is associated with the  
234 enhanced expression of branchial ion pumps and transporters (Pelis and McCormick, 2001;  
235 Tipsmark and Madsen, 2009).

236 Rearing Mozambique tilapia under a TR increased pituitary *gh* expression and plasma Gh  
237 to levels above those of both steady-state SW- and FW-acclimated fish; however, there were no  
238 differences detected between the two phases of the TR (Fig. 3C and D) (Moorman et al., 2016).  
239 The elevations in pituitary *gh* and plasma Gh observed in tilapia reared in a TR are consistent  
240 with the sexually dimorphic regulation of growth and the higher growth rates, in both sexes, of  
241 fish reared in TR versus steady-state FW or SW (Moorman et al., 2016; Seale et al., 2020b). The  
242 transcriptional activation of downstream components of the Gh/Igf system in fish reared in TR,  
243 such as the growth hormone receptor (*ghr*), *igf1*, and *igf2* in liver and muscle, followed patterns  
244 resembling the *prlrs* where transcript levels changed between TF and TS (Fig. 3F and G)  
245 (Moorman et al., 2016). Specifically, the expression of *ghr* in liver and muscle, and *igfs* in liver,  
246 was higher in TF than in TS. These patterns suggest that Gh-sensitivity in muscle and liver is  
247 impacted by environmental salinity in an oscillatory fashion that may underlie the increased



248 growth observed in fish reared under TR. Moreover, because the changes in *ghr* and *igfs*  
249 resembled the dynamic regulation of *prlrs* observed in the gill under an identical regime, they  
250 contribute to a broader pattern of endocrine regulation where relative systemic stability is  
251 coupled with organ-level changes at the level of receptors and downstream effectors in response  
252 to tidally-changing salinities.

253

## 254 **5. Effectors of ion transport in tidally-changing salinities**

255 Whether regulated by endocrine factors, or in direct response to changes in salinity,  
256 effectors of hydromineral balance respond uniquely to tidally-changing salinities. A closer look  
257 at the branchial ‘ionocytes’ (also termed ‘mitochondria-rich cells’) of tilapia illustrates this  
258 notion. In Mozambique tilapia, four ionocyte sub-types have been identified based upon the  
259 expression of ion-transporting proteins: Type-I cells express an apical renal outer medullary K<sup>+</sup>  
260 channel (Romka) and basolateral Na<sup>+</sup>/K<sup>+</sup>-ATPase (Nka; Atp1a); Type-II cells express an apical  
261 Na<sup>+</sup>/Cl<sup>-</sup> cotransporter 2 (Ncc2; Slc12a10) and basolateral Nka; Type-III cells are characterized by  
262 an apical Na<sup>+</sup>/H<sup>+</sup> exchanger 3 (Nhe3; Slc9a3) in addition to basolateral Nka; Type-IV cells  
263 express an apical cystic fibrosis transmembrane conductance regulator (Cftr) and basolateral  
264 Na<sup>+</sup>/K<sup>+</sup>/2Cl<sup>-</sup> cotransporter 1a (Nkcc1a; Slc12a2a) and Nka (Hiroi et al., 2005; Kaneko et al.,  
265 2008; Furukawa et al., 2014).

266 The functional and morphological changes that occur among branchial ionocytes have  
267 been traditionally identified through one-way salinity transfer paradigms. By examining the  
268 ionocytes of tilapia maintained under a TR, we observed that ionocytes do not necessarily  
269 change in response to the FW- and SW-phases of a tidal cycle as would be expected if such  
270 changes mimicked those seen during one-way salinity transfers. For example, tilapia raised under  
271 a TR exhibited Cftr- and Nkcc-immunopositive cells that did not change between the FW- and  
272 SW-phases of the tidal cycle; immunoreactivity more closely approximated the patterns observed  
273 in fish continuously maintained in SW (Moorman et al., 2014). In addition, fish acclimated to the  
274 TR maintained the gene expression of pumps and transporters involved in ion uptake, such as  
275 *ncc2*, *nkaα1a*, and *nhe3* (Hiroi et al., 2008; Inokuchi et al., 2008; Tipsmark et al., 2011) at higher  
276 levels than fish acclimated to steady-state SW, but much lower than fish acclimated to FW. In  
277 some cases, such as with *ncc2*, there were differences between TF and TS, but they typically  
278 occurred within a much narrower range than observed between steady-state FW and SW fish

279 (Fig. 3E) (Moorman et al., 2014). By contrast, the expression of genes encoding effectors of ion  
280 secretion, such as *nkcc1a* and *cftr*, fluctuated between TF and TS with the same magnitude  
281 observed between steady-state FW and SW acclimated fish (Fig. 3E). These patterns of ionocyte  
282 regulation in tilapia are consistent with the similarities observed in TEP measurements between  
283 mummichogs reared in SW versus tidally-changing environments (Wood and Grosell, 2009),  
284 though some important differences in the regulation of molecular transducers of ion and water  
285 transport are worth noting. While tilapia in TF and TS maintained ionocyte numbers similar to  
286 fish in SW, *Cftr* immunoreactivity was dramatically decreased in TF fish (Moorman et al., 2014).  
287 The decrease in *Cftr* abundance that was evident in TF is indicative of salinity-dependent  
288 trafficking of *Cftr* to the apical membrane (Marshall and Singer, 2002). This apparent regulation  
289 of *Cftr* expression is important because otherwise the extrusion of Cl<sup>-</sup> by Type-IV ionocytes  
290 would be highly deleterious to fish in FW. Hence, the rapid decrease in *Cftr* contributes to how  
291 fish exposed to tidal conditions can survive in FW for 6 h without recruiting ion-absorptive  
292 ionocytes.

293 In addition to the changes in *Cftr*, branchial *aquaporin 3* (*aqp3*) expression differed  
294 between the two phases of the tidal cycle. This suggests that modulating *Aqp3* function (possibly  
295 water/urea/glycerol permeability) is important even in fish that experience a different salinity for  
296 just 6 h (Moorman et al., 2014; Moorman et al., 2015). Although responding in opposing  
297 fashions, *cftr* and *aqp3* were highly sensitive to continually changing salinities, and especially in  
298 response to transfers from TR to steady-state salinities (Moorman et al., 2015). The high  
299 sensitivity of these transcripts to tidally-changing salinities is also consistent with the previously  
300 proposed roles of their encoded proteins in branchial mechanosensory signal transduction (Cutler  
301 and Cramb, 2002; Madsen et al., 2014; Marshall, 2003; Marshall, 2011; Marshall and Singer,  
302 2002; Watanabe et al., 2005) and PrI cell osmoreception through osmotically-driven changes in  
303 cell volume (Seale et al., 2003; Seale et al., 2012b; Watanabe et al., 2009; Weber et al., 2004).  
304 Therefore, the dynamic nature of *Cftr* and *Aqp3* expression in TR further strengthens the notion  
305 that each one plays indispensable roles in maintaining hydromineral balance, where an  
306 abundance of the former in FW would underlie Cl<sup>-</sup> loss and an abundance of the latter in SW  
307 could lead to excessive osmotic water loss. In addition to the acute responses of *cftr* and *aqp3* to  
308 extracellular osmolality, evidence from studies employing the TR paradigm indicates that these  
309 and other osmoregulation-related transcripts may be tied to hormone receptor dynamics in TR

310 (Seale et al., 2019). The expression of *ncc2* and *aqp3*, for example, is upregulated during FW  
311 acclimation and is stimulated by Prl (Breves et al., 2016; Breves et al., 2010b), whose actions in  
312 tidal conditions may be mainly modulated by receptor expression rather than circulating levels of  
313 hormone. Thus, the observed changes in *prlr* expression may represent a strategy to fine-tune the  
314 actions of Prl in circulation within time-frames that may be too short for pituitary-based  
315 processes to respond.

316 The functional consequences of tilapia experiencing tidal conditions has also been  
317 experimentally addressed through transfers from TR to steady-state environments (Moorman et  
318 al., 2015) and vice versa (Pavlosky et al., 2019). Notably, when compared with fish reared in  
319 FW, fish reared in a TR had a significantly improved ability to survive direct transfer to SW  
320 (Moorman et al., 2015). When adult fish (>2 years in age) were transferred from either steady-  
321 state FW or SW to TR, however, they were able to survive the abrupt and cyclical salinity  
322 changes imposed by the TR, and within seven days, exhibited ion-transporter/channel  
323 transcriptional patterns similar to those observed in fish reared in TR from the fry stage  
324 (Pavlosky et al., 2019). Thus, acclimation to a TR can occur within a range of life stages, where  
325 a combination of direct and endocrine-mediated responses to osmotic stimuli enable fish to cope  
326 with the vicissitudes of estuarine environments.

327

## 328 **6. Conclusion and future perspectives**

329 Studies employing simulations of tidally-changing salinities provide an important  
330 perspective on the adaptive processes that support salinity acclimation. Euryhaline teleosts, such  
331 as mummichogs and Mozambique tilapia, tolerant of environments characterized by frequent  
332 changes in salinity are ideally suited for such investigations. The rearing conditions/life history  
333 of a given individual is a key determinant of that individual's capacity to exhibit plastic  
334 osmoregulatory activities. In both species, extended pre-exposures to tidally-changing salinities  
335 enabled their performance in stable hypo- or hyperosmotic environments (Moorman et al., 2015;  
336 Wood and Grosell, 2009). Underlying this adaptive capacity are distinct patterns of endocrine  
337 control over downstream effectors of ion transport that enable plasma osmolality to remain  
338 within a narrow range. While more closely resembling fish acclimated to hyperosmotic  
339 environments, fish reared under a TR maintain a distinct osmoregulatory strategy that does not  
340 fully coincide with animals exposed to 'one-time' transfers between FW and SW.

341           There is no question that a broad collection of hormones beyond Prl and Gh play key  
342 roles in promoting hydromineral balance in teleosts (Takei et al., 2014). For instance, cortisol is  
343 deemed a ‘SW-adapting hormone’ because it directly stimulates the activities and/or expression  
344 of Na<sup>+</sup>/K<sup>+</sup>-ATPases and ion transporters tied to ion extrusion pathways in the gill; cortisol also  
345 acts indirectly by synergizing with Gh/Igf1 signaling (McCormick, 2001). While cortisol  
346 exhibits clear responses to ‘one-time’ changes in salinity (Kajimura et al., 2004), to our  
347 knowledge, there is no information on the patterns of its release during tidally-changing  
348 conditions. It will be interesting to learn whether plasma cortisol levels change in parallel with  
349 environmental salinity, or like circulating Prl levels, become uncoupled from changes in plasma  
350 osmolality during tidal cycles.

351

#### 352 **Declaration of interest**

353           The authors declare that there are no conflicts of interest that could be perceived as  
354 hindering the impartiality of the research reported.

355

#### 356 **Figure legends**

357 **Figure 1.** Experimental tidal regime (TR) paradigm with fish reared in alternating FW and SW  
358 phases (TF and TS, respectively); blue bands represent SW and light blue bands represent FW  
359 (A). Illustration of the tank setup employed for TR experimental paradigms (B). The setup is  
360 designed to supply FW or SW to experimental tanks every 6 h, thereby simulating a tidal cycle.  
361 Reservoir tanks containing either FW or SW supply water to experimental tanks via a  
362 submersible pump. The pump is plugged into a timer that is offset by 6 h between the FW and  
363 SW reservoirs.

364

365 **Figure 2.** Plasma osmolalities of Mozambique tilapia: 1) in steady-state seawater (SW) and fresh  
366 water (FW); 2) transferred one-way from FW to SW (F-S) and from SW to FW (S-F); and 3) in a  
367 tidal regime sampled in the SW phase (TS) and in the FW phase (TF). Transferred fish were  
368 sampled at 6 h. Bars represent mean ± S.E.M. Plasma osmolality of fish in SW (black) and FW  
369 (grey) (*n* = 10-20). A complete transition between salinities in the transfer and tidal regime  
370 groups occurred within 1 h. Data were analyzed by 2-way ANOVA with acclimation regime and  
371 salinity as main effects, followed by Bonferroni’s post-hoc test. \*\*\* indicate main and

372 interaction effects at  $P < 0.001$ ; †, ††† indicate significant differences between salinities at each  
373 acclimation regime at  $P < 0.05$  and  $P < 0.01$ , respectively. Letters not shared between bars  
374 indicate significant differences between acclimation regimes within salinities at  $P < 0.05$ . Figure  
375 adapted from Moorman et al. (2015).

376

377 **Figure 3.** Effects of salinity regime on endocrine regulators of growth and ionoregulation. Plus  
378 and minus signs next to each gene transcript indicate an increase or decrease, respectively,  
379 relative to each other; faded signs denote attenuated responses and equal signs denote similar  
380 values that are intermediate to the other values being compared. Environmental salinity ranges in  
381 steady-state FW and SW and tidally-changing salinities between FW (TF) and SW (TS) (A).  
382 Changes in salinity elicit changes in plasma osmolality within narrow ranges for fish in steady-  
383 state FW and SW, and TF and TS (B). *prl<sub>177</sub>* and *prl<sub>188</sub>* expression is higher in steady-state FW  
384 versus SW while no difference is observed between fish in TF and TS; the opposite pattern is  
385 seen with *prlr1*, which was unchanged between FW- and SW-acclimated fish and different  
386 between TF and TS; expression of *gh* is highest in TF and TS, followed by SW and FW (C).  
387 Plasma Prl<sub>188</sub> and Prl<sub>177</sub> levels follow a similar pattern as pituitary levels of their associated gene  
388 transcripts; Prl<sub>188</sub> and Prl<sub>177</sub> are elevated in steady-state FW relative to SW, but unchanged  
389 between TF and TS; plasma Gh levels are similar between all salinity regimes (D). Branchial  
390 *prlr1* was consistently elevated in FW and TF fish relative to SW and TS fish, respectively,  
391 while *prlr2*, *nkcc1a*, and *cfr* levels were elevated in fish in SW and TS relative to those in FW  
392 and TF; *ncc2*, *nka $\alpha$ 1b*, and *aqp3* showed the same direction of response in TF and TS fish  
393 compared to FW and SW, but the magnitude of expression was attenuated and generally lower  
394 than that of steady-state fish. Lastly, *nhe3* and *nka $\alpha$ 1a* were differentially expressed between FW  
395 and SW but not between TF and TS (E). In liver, *ghr* and *igf1* did not vary between steady-state  
396 salinities, but changed between TF and TS; *igf2* changed between steady-state salinities and  
397 between TF and TS (F). In muscle, *ghr* did not change between steady-state salinities but  
398 changed between TF and TS; transcript levels in FW and SW fish were low compared with those  
399 in TF. The expression of *igfs* was relatively unchanged across all salinity regimes, with the  
400 exception of the elevation of *igf1* in fish in SW relative to those in FW (G). This summary of  
401 plasma hormone and gene expression patterns in steady-state and tidal salinities is based on  
402 results from Moorman et al. (2014), Moorman et al. (2015), Moorman et al. (2016), Pavlosky et

403 al. (2019), Seale et al. (2020b), and Seale et al. (2019). Abbreviations: Osm: osmolality  
404 (mOsm/kg); FW: fresh water; SW: seawater; TF: FW phase of tidal cycle; TS: SW phase of tidal  
405 cycle; *prl<sub>188</sub>*: prolactin 188; *prl<sub>177</sub>*: prolactin 177; *gh*: growth hormone; *ghr*: growth hormone  
406 receptor; *igf1* and *-2*: insulin-like growth factor 1 and -2; *prlr1* and *-2*: prolactin receptor 1 and -  
407 2 ; *ncc2*: Na<sup>+</sup>/Cl<sup>-</sup> cotransporter 2; *nkcc1a*: Na<sup>+</sup>/K<sup>+</sup>/2Cl<sup>-</sup> cotransporter 1a; *nhe3*: Na<sup>+</sup>/H<sup>+</sup>  
408 exchanger 3; *nka $\alpha$ 1a* and *- $\alpha$ 1b*: Na<sup>+</sup>/K<sup>+</sup>-ATPase alpha 1a and -alpha 1b; *cftr*: cystic fibrosis  
409 transmembrane conductance regulator; *aqp3*: aquaporin 3. Abbreviations in italics denote gene  
410 transcripts.

411

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419

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424

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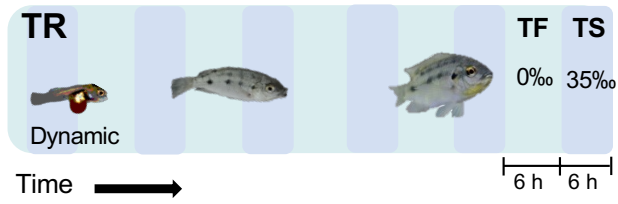
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Figure 1

A



B

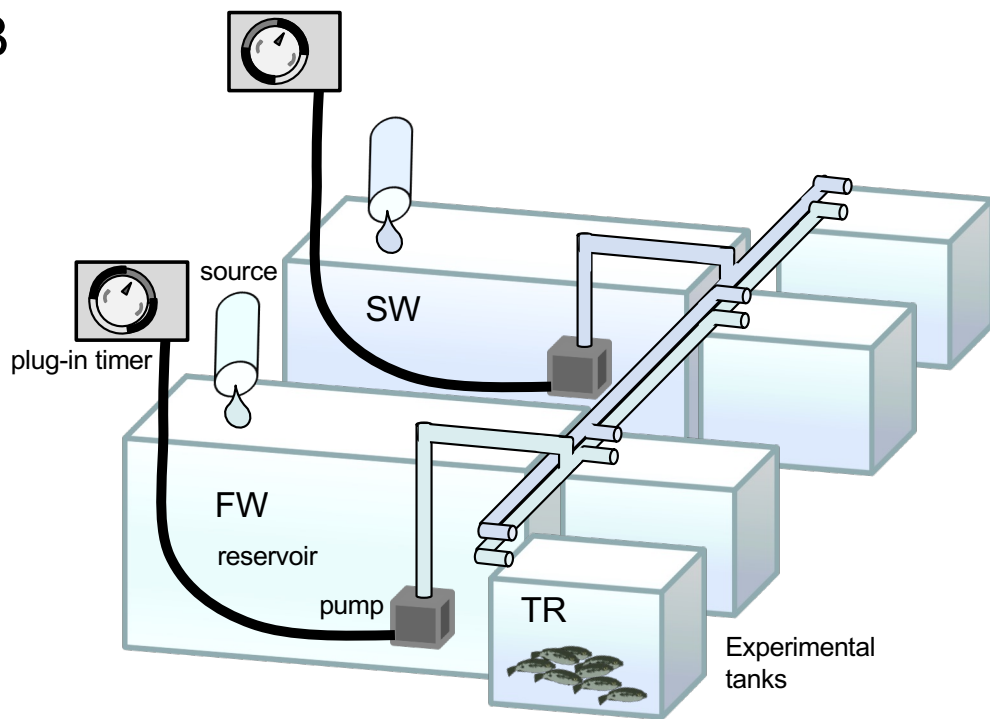


Figure 2

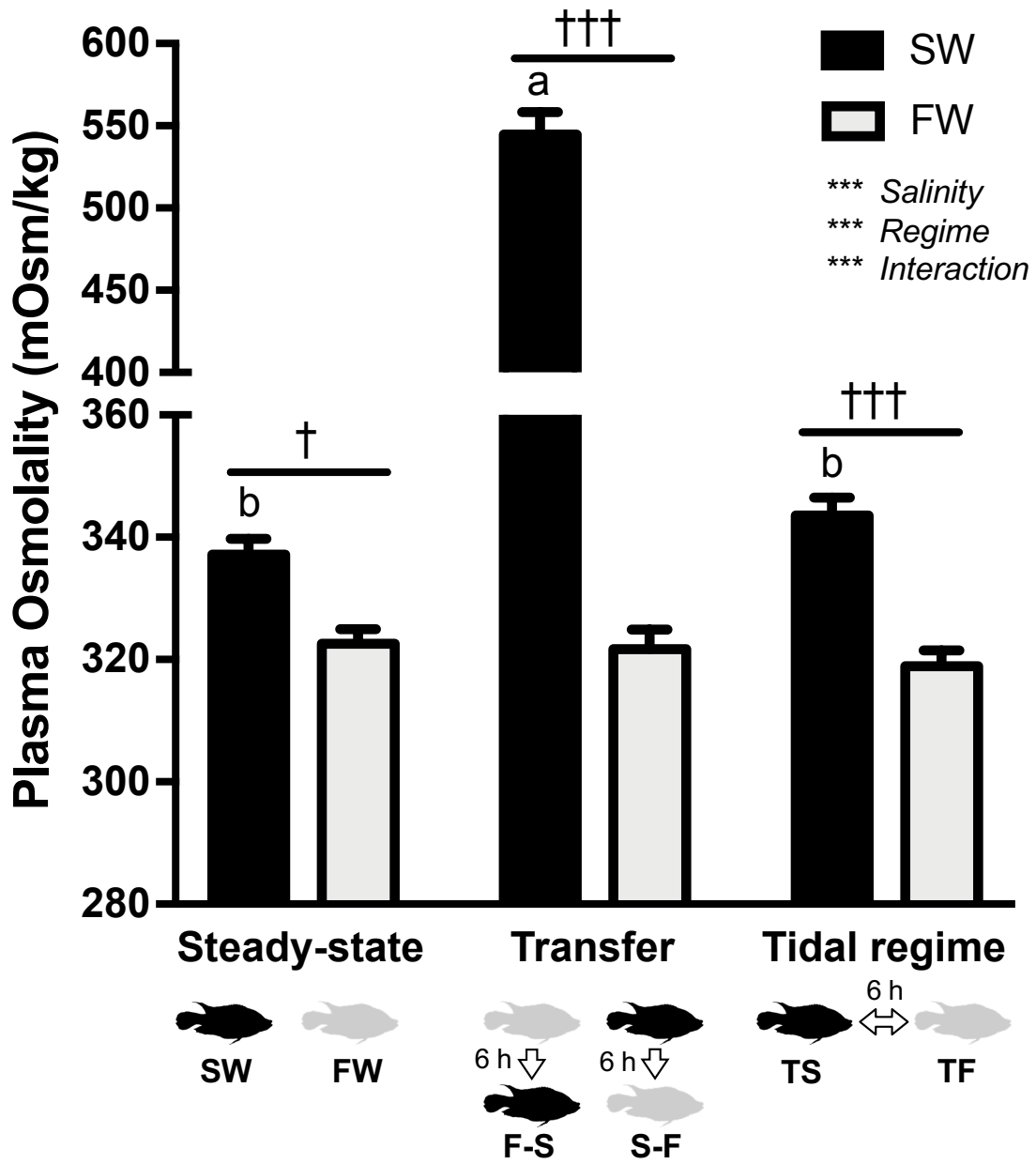


Figure 3

