

1 The aversive brain system of teleosts: Implications for neuroscience and biological psychiatry

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10

11 Abstract

12 Defensive behavior is a function of specific survival circuits, the “aversive brain system”, that are
13 thought to be conserved across vertebrates, and involve threat detection and the organization of
14 defensive responses to reduce or eliminate threat. In mammals, these circuits involve amygdalar and
15 hypothalamic subnuclei and midbrain circuits. The increased interest in teleost fishes as model
16 organisms in neuroscience created a demand to understand which brain circuits are involved in
17 defensive behavior. Telencephalic and habenular circuits represent a “forebrain circuit” for threat
18 processing and organization of responses, being important to mounting appropriate coping
19 responses. Specific hypothalamic circuits organize neuroendocrine and neurovegetative outputs, but
20 are the less well-studied in fish. A “midbrain circuit” is represented by projections to interneurons in
21 the optic tectum which mediate fast escape responses via projections to the central gray and/or the
22 brainstem escape network. Threatening stimuli (especially visual stimuli) can bypass the “high
23 road” and directly activate this system, initiating escape responses. Increased attention to these
24 circuits in an evolutionary framework is still needed.

25 *Keywords:* Fear; Anxiety; Aversive brain system; Comparative neuroanatomy; Teleost fish

26

27 **1. Introduction**

28 Defensive behavior is a function of specific survival circuits that are highly conserved
29 across vertebrates (LeDoux, 2012), and involve threat detection, as well as the organization of
30 species-specific defensive responses to reduce or eliminate threat (LeDoux, 2012). In mammals,
31 these circuits involve sensory systems that detect conditioned and unconditioned threatening
32 stimuli, amygdalar circuits that select appropriate behavioral and neuroendocrine responses and
33 promote aversive learning, hypothalamic circuits for both defensive behavior and neuroendocrine
34 responses, and midbrain circuits that control the output (LeDoux, 2012). This survival circuit for
35 defensive behavior links this appraisal phase, in which the global organismic state is determined
36 based on threat detection, to a response phase, in which “the instrumental physiological and
37 behavioural responses are determined and executed” (Andersen et al., 2016, p. 5). Different threat
38 levels restrict attention “in the current global organismic state [to make] the organism focus on a
39 short-term motive to [...] survive a threat” (Andersen et al., 2016, p. 6).

40 In fish, defensive circuits have not been thoroughly described, but there is and increasing
41 awareness that defensive behavior is a fundamental function in these animals (Andersen et al.,
42 2016; Kalueff et al., 2012; Kittilsen, 2013). As a result, fish are increasingly being used in
43 neurobehavioral research (Gerlai, 2014; Hall et al., 2014; Kalueff et al., 2014b; Stewart et al.,
44 2015). A framework for establishing homologies for defensive circuits in fish, therefore, is useful
45 for this enterprise. Classical neuroscientific approaches to describing behavioral circuits involve the
46 demonstration that a given structure is activated (or inhibited) by specific stimuli or behavioral
47 paradigms, as well as the effects of lesioning (or silencing) and stimulating the structure. Even
48 though there are many technical difficulties in performing functional neuroanatomy in the
49 diminutive zebrafish, novel genetic techniques (Friedrich et al., 2010) have allowed important
50 discoveries regarding the limbic system.

51 The increase in the use of zebrafish (Kalueff et al., 2014a) and other species (Hall et al.,
52 2014) as organisms in behavioral models for anxiety- and fear-like behavior also prompted the need
53 to describe the neural structures that are involved in these behaviors (Guo et al., 2012). While the
54 unspoken assumption in the field appears to be one of “multiple realizability” (Bickle, 2010) –
55 including the idea that it is the behavioral function that needs to be similar, and not the brain circuits
56 underlying it, in order to use these assays as models in psychopathology (Wright, 2002) –, at least
57 from an evolutionary point of view it is interesting to assess whether the putative conservation of
58 behavioral responses to aversive stimuli is accompanied by conservation of the neural substratum
59 (Striedter et al., 2014). The present article reviews the circuitry that is involved in detecting,
60 identifying, processing, and responding to threat in fish, with a special focus on zebrafish models.

61

62 **2. Discovering neural circuits for aversive behavior in fish**

63 An important starting point in determining the aversive behavior network in zebrafish is
64 activation studies. Typically, this is done by examining immediate early gene products, such as the
65 *cfos* gene and its protein, c-Fos. A network comprised of the medial nucleus of the dorsal
66 telencephalon (Dm), ventral nucleus of the ventral telencephalon (Vv), supracommissural nucleus
67 of the ventral telencephalon (Vs), and preoptic region (POA) shows upregulated *cfos* expression
68 after exposure to conspecific alarm substance [CAS] (Faustino et al., 2017). The expression of c-
69 Fos during CAS exposure appears to be specific, as no increases were observed in the lateral
70 nucleus of the dorsal telencephalon (Dl), dorsal nucleus of the ventral telencephalon (Vd), habenula,
71 or thalamus (Ruhl et al., 2017). Dm is also activated after exposure to the light/dark test (Lau et al.,
72 2011; von Trotha et al., 2014), an anxiogenic stimulus for zebrafish (Maximino et al., 2010a). Thus,
73 it appears that a network involving Dm, Vv, Vs, and POA is involved in detecting and/or processing
74 aversive stimuli and/or selecting behavioral strategies in zebrafish. A network for the retrieval of

75 fear memories include Dm, Dl, and thalamus, as these structures showed c-Fos expression during
76 presentation of a red light that has been previously associated with CAS (Ruhl et al., 2017).

77 Other aversive stimuli have also been used to investigate threat detection and processing in
78 zebrafish larvae. A looming stimulus, which simulates predatory attack, activates neurons in the
79 optic tectum (Dunn et al., 2016; Temizer et al., 2015). Tyrosine hydroxylase-positive neurons in the
80 caudal hypothalamus (Hc) respond to aversive stimuli, including pH change, high ammonia, and
81 handling stress (Semenova et al., 2014). Using immunohistochemical detection of phosphorylated
82 extracellular signal-regulated kinase (pERK), Randlett et al. (2015) mapped the activity of nuclei in
83 the larval brain in response to different aversive stimuli (exposure to mustard oil, dish taps, heated
84 water, electric shocks). The authors found that while each stimulus promoted pERK activity in
85 specific regions (due to the diversity of sensory channels tapped by each stimulus), all four
86 activated the locus coeruleus, caudal hindbrain, Hc, POA, and subpallium. It is not clear whether
87 the subpallial regions observed in this experiment overlap with those found in CAS-exposed adults
88 (i.e., Vv and Vs); nonetheless, these results suggest a wider network that is specialized in the
89 detection of aversive stimuli which includes pallial and subpallial components of the amygdala
90 homologues, the neuroendocrine hypothalamus, optic tectum, locus coeruleus, and the brainstem
91 escape network.

92 The focus of this review will now turn to each of these foci. We will quickly review some of
93 the evidence for proposing these areas as full or partial homologues of specific mammalian nuclei
94 which have been implicated in defensive behavior. Importantly, we will review evidence for the
95 participation of these regions in aversive behavior in fish, and delineate, when possible, the circuitry
96 that is involved in these roles.

97

98 **3. Pallial and subpallial components of the teleostean amygdala**

99 The fact that the teleostean telencephalon is everted precluded, for a long time, the easy
100 establishment of homologies for telencephalic structures (Nieuwenhuys, 2011). Different eversion
101 models lead to different proposed final topologies of pallial structures in adult fish (Braford, 2009;
102 Butler, 2000; Mueller et al., 2011; Nieuwenhuys, 2011; Northcutt, 2008; Wullimann and Mueller,
103 2004; Yamamoto et al., 2007); as a result, different propositions have been made regarding
104 amygdala homologues in zebrafish. Based on topology and topography, cytoarchitectonics,
105 neurochemistry, expression of developmental regulatory genes, and behavioral data, we will follow
106 our previous proposal that the medial nucleus of the dorsal telencephalon (Dm) is homologous to
107 the “limbic associative” amygdaloid system (lateral and basolateral amygdala of mammals), while
108 the subcommissural and postcommissural nuclei of the ventral telencephalon (Vs and Vp,
109 respectively) are homologous to the “autonomic” amygdaloid system (central extended amygdala
110 [CEXA]) (Maximino et al., 2013a)(Figure 1A).

111 Lesion studies implicated the Dm in associative learning of aversive memories in goldfish.
112 Ablation of the Dm, but not of the Dl, impair the acquisition and maintenance of two-way (active)
113 avoidance acquisition in this species (Portavella et al., 2004b, 2004a, 2002; Portavella and Vargas,
114 2005). c-Fos studies with zebrafish have also suggested a role for the Dm in aversive conditioning:
115 the association between a red light and CAS produces conditioned responses to the light that are
116 very similar to the innate repertoire of zebrafish (Hall and Suboski, 1995a, 1995b; Ruhl et al.,
117 2017). Interestingly, during the acquisition phase (when the red light and the CAS are presented
118 together), the number of c-Fos-positive cells is higher in the Dm, but not in the lateral nucleus of the
119 dorsal telencephalon (Dl, the hippocampal homologue; Goodson & Kingsbury, 2013) or in the
120 dorsal nucleus of the ventral telencephalon (Vd, the striatum homologue; Ganz et al., 2011). 24 h
121 later, during the retrieval phase (when only the red light is presented), c-Fos-positive cells are
122 increased in the Dm, Dl, and thalamus (Ruhl et al., 2017). In any case, it appears that the Dm is
123 involved in both the acquisition and the maintenance of aversive memories in cyprinids.

124 The majority of neurons in pallial regions are glutamatergic (von Trotha et al., 2014), with
125 some GABAergic interneurons. Lal et al. (2018) produced *Gal4FF* zebrafish lineages driving the
126 expression of the botulinum toxin B light chain (BoTxBLC) gene that showed deficits in active
127 avoidance conditioning. Among these lineages, two showed expression limited to the Dm, and in
128 both the trapped gene was *emx3*. In addition to showing deficits in conditioned aversive behavior,
129 these animals also showed changes in their response to CAS, with decreased freezing but increased
130 erratic swimming. Only 16% of the cells in the Dm express the Gal4FF construct, and the majority
131 are neurons; 94% of these neurons are glutamatergic. These neurons project heavily to the
132 hypothalamus, including anterior tuberal nucleus (ATN), lateral hypothalamic nucleus (LH), and
133 dorsal zone of the periventricular hypothalamus, as well as to telencephalic regions (entopeduncular
134 nucleus, preoptic area, Vd, and Vs)(Figure 1B), suggesting neural networks involved in both
135 conditioned and unconditioned aversive behavior (Lal et al., 2018).

136 The role of the Dm in unconditioned behavior has also been assessed. In zebrafish, exposure
137 to the light/dark test increases c-Fos expression in the Dm (Lau et al., 2011; von Trotha et al., 2014).
138 Moreover, CAS exposure also increases *cfos* in the Dm of zebrafish (Faustino et al., 2017),
139 suggesting that ethologically-relevant aversive stimuli recruit these cells even when conditioning is
140 not involved. As already discussed, expression of BoTxBLC in a subset of Dm neurons in zebrafish
141 decreases freezing responses and increases erratic swimming after CAS exposure (Lal et al., 2018).
142 A participation in restraint stress-induced behavioral responses has also been suggested. In Nile
143 tilapia (*Oreochromis niloticus*), restraint stress increases 5-HT metabolism in the Dm (Silva et al.,
144 2014). Likewise, acute stress (lowering water levels) increases *htr1aa* and *htr1ab* expression in the
145 Dm of Rainbow trout (Vindas et al., 2017); interestingly, 5-HIAA levels were increased in the Dm
146 only in those animals which were classified as “proactive” in relation to a behavioral response to
147 hypoxia (Vindas et al., 2017), suggesting that 5-HT acts in this nucleus to mount active responses to
148 remove stressors and restore homeostasis. Moreover, destroying 5-HT fibers in the telencephalon

149 with the serotonergic toxin 5-7-DHT decreases the ratio of correct avoidance responses in an active
150 avoidance paradigm in zebrafish (Amo et al., 2014). Thus, feedback mechanisms from
151 neuromodulators appear to also be important in these responses (Table 1). Consistent with the
152 hypothesis of a role of the Dm in active responses to aversive stimuli, inhibition of the Dm in
153 *Leporinus macrocephalus* by injecting midazolam abolished stress-induced analgesia (Wolkers et
154 al., 2015).

155 In rodents, different components of the limbic associative amygdalar system mediate
156 responses innate stimuli (auditory and visual components of predator cues) vs. conditioned stimuli
157 (i.e., stimuli which were previously paired with noxious stimuli)(Gross and Canteras, 2012). In
158 mammals, olfactory components of predator cues bypass the cortical components of the amygdala,
159 instead projecting directly to subcortical components (posteroventral part of the medial amygdala;
160 Gross and Canteras, 2012). These different components process distinct afferent inputs and channel
161 them to parallel downstream efferent pathways, responsible for output circuits for fear of predators
162 and fear of pain (Gross and Canteras, 2012). In zebrafish, sensory input to the posterior
163 telencephalon (including Dm) suggest massive olfactory information (e.g., Miyasaka et al., 2014),
164 while visual and auditory stimuli are processed initially in the optic tectum (Northcutt, 1983; Meek,
165 1990); however, the fact that an olfactory stimulus can be associated with a visual stimulus in the
166 Dm (Ruhl et al., 2017; Lal et al., 2018) suggests the participation of a projection from the optic
167 tectum, possibly via the lateral preglomerular nucleus (Carr, 2015). Thus, in zebrafish the Dm also
168 appears to be important for learned fear, but the participation in unconditioned behavior suggests
169 that either subpopulations of the Dm process different stimuli, or that the evolutionary shift towards
170 visual and auditory stimuli in mammals displaced the massive olfactory projections that was found
171 in fish.

172 If there is considerable evidence for the homology between Dm and the limbic associative
173 amygdala of mammals, this is not the case for the homology between Vs/Vp and the autonomic

174 amygdala; in fact, it is more probable that these regions are only *partially* homologous to the CEXA
175 (Goodson and Kingsbury, 2013; Maximino et al., 2013a). In mammals, the CEXA has been
176 proposed to mediate the selection of vegetative and behavioral responses to threatening stimuli
177 (Gozzi et al., 2010; LeDoux and Pine, 2016), as well as conditioned fear (Ciocchi et al., 2010;
178 Haubensak et al., 2010; LeDoux, 1998). The circuit acts by tonically inhibiting downstream
179 responses mediated by the periaqueductal gray area (PAG) and hypothalamus (freezing, flight), or
180 by the basal forebrain (risk assessment); inhibition of these circuits (by, e.g., signals from the
181 “limbic associative” amygdala) release behavior in one of these two streams, producing appropriate
182 active or passive responses to the threatening stimulus (Maximino, 2012). In zebrafish, these
183 neurons are also GABAergic (Mueller and Guo, 2009), suggesting a similar circuit.

184 Almost no behavioral evidence exists for the role of Vs/Vp in defensive behavior. One of the
185 possible reasons is that the Vs and Vp are continuous with the Vd in a rostrocaudal axis, leading
186 authors to report effects (or lack thereof) of interventions in wrongly identified structures. One
187 important exception is the observation that CAS increases *cfos* expression in the Vs (Faustino et al.,
188 2017). Interestingly, as reported above, this work also found increased *cfos* in the Dm, Vv, and
189 POA. Importantly, control animals showed functional connectivity between these regions, while
190 exposure to CAS decreased these correlations; in fact, animals exposed to CAS showed only co-
191 activation of Dm-Vs and Vv-Vs (Faustino et al., 2017). The precise mechanism for this decreased
192 coherence is yet to be described; however, considering the equivalent mammalian circuit, it is
193 possible that activation of the Dm activates inhibitory neurons from the Vs, which could be
194 responsible for decreased connectivity with Vv and POA, while at the same time disinhibiting
195 downstream (hypothalamic and mesencephalic) mechanisms for response emission (Figure 1B).

196 In summary, the Dm appears to be important for both conditioned and unconditioned
197 aversive responses, as well as to mounting appropriate (active vs. passive) responses. The Vs also

198 appears to participate in processing unconditioned threat, but its role in response selection and
199 learning is still unknown.

200

201 **3. Does the habenula participate in defense?**

202 The habenulae are paired structures located in the roof of the rostral diencephalon of fishes,
203 divided classically into a dorsal, asymmetric portion (dHb) and a ventral, symmetric portion (vHb).
204 The dHb can be further subdivided into medial and lateral subnuclei (Aizawa et al., 2011; Okamoto
205 et al., 2011, 2008), although eleven subdivisions (Hb01 - Hb10) are suggested by single-cell RNA-
206 Seq registered to anatomy (Pandey et al., 2018). Four clusters were identified in the zebrafish vHb
207 (Hb11, Hb12, Hb13, and Hb15), based on the expression of genetic markers and signaling
208 molecules (Pandey et al., 2018). The vHb receives (putatively glutamatergic) projections from the
209 ventral portion of the entopeduncular nucleus (Okamoto et al., 2011; Amo et al., 2014), which by its
210 turn receive excitatory projections from the Dm (Lal et al., 2018); as a result, a excitatory
211 feedforward circuit exists from Dm to vEN to vHb.

212 The difference in size between left and right habenulae is due mainly to the enlargement of
213 the lateral subnucleus (dHbL) in the left habenula in relation to the right, which shows an enlarged
214 medial subnucleus (dHbM) (Amo et al., 2010; Okamoto et al., 2011). This difference correlates with
215 parapineal asymmetry (Gamse et al., 2003) and is concordant with the lateralization of viscera
216 (Barth et al., 2005; Domenichini et al., 2011) and subsets of behavioral functions (Barth et al., 2005;
217 Dadda et al., 2010; Facchin et al., 2009). In the left dHb, high levels of the potassium channel
218 tetramerization domain-containing protein 12.1 (*kctd12.1/leftover/lov*) are expressed, while low
219 levels are observed in the right dHb. Conversely, two other members of the KTCD family, *kctd12.2*
220 (*right on/ron*) and *kctd8 (dexter/dex)*, are expressed exclusively in the right dHb (Beretta et al.,
221 2012; Gamse et al., 2005; Y. Kuan et al., 2007; Roussigné et al., 2011). High levels of *murb* and
222 the adrenoceptor beta 2 *adrb2a* are found in clusters enriched in the right dHb, while clusters

223 enriched in the left dHb show high levels of the protochaderin *pcdh7b*, the Wnt family member
224 *wnt7aa*, adenylate cyclase-activating polypeptide *adcyap1a*, and the protein phosphatase regulatory
225 inhibitor *ppplr1c* (Pandey et al., 2018). These genes have been used as markers for the
226 neuroanatomical divisions of the dHb. However, caution should be exercised in transposing these
227 neuroanatomical subdivisions to functional ones: DeCarvalho et al. (2014) showed that cholinergic
228 neurons in the left dHb are a distinct population from cells expressing *kctd12.1*, and part of the
229 developmental expansion of the lateral dHb from larvae to adults is accompanied by the invasion of
230 the left dHb with cholinergic neurons (in larvae, most neurons in the left dHb are glutamatergic
231 only) and an expansion of substance P and somatostatin-expressing neurons.

232 While the dHb is asymmetric, the ventral habenula (vHb) is symmetric (Amo et al., 2010;
233 Okamoto et al., 2011). The vHb is characterized by the expression of *diamine oxidase (dao)*, *lov* and
234 *protocadherin 10a (pcdh10a)* mRNA, in a complementary fashion with the expression of *POU*
235 *domain, class 4, transcription factor 1 (brn3a)* which is expressed exclusively in the medial
236 subnucleus (Amo et al., 2010). Cluster Hb11 is marked by high expression of the transcription
237 factor *sox1a*, cluster Hb12 by high expression of the 5-HT_{1A} receptor *htr1aa*, cluster Hb13 by high
238 expression of *beta-tubulin 5 (tubb5)*, and cluster Hb15 by the high expression of the kisspeptin gene
239 *kiss1* (Pandey et al., 2018). Again, these genes have been used as markers for the neuroanatomical
240 subdivision of the vHb.

241 The neuroanatomical asymmetry of the dHb has produced an interesting literature on its
242 behavioral correlates, especially in zebrafish. The neuroanatomical asymmetry correlates with
243 behavioral asymmetries in different assays. Animals with left parapineal position (L-PPO) tend to
244 use their right eye when viewing a mirror and swim preferentially in a clockwise direction, while
245 animals with right parapineal (R-PPO) do not show this preference; conversely, R-PPO zebrafish
246 use the right eye to inspect a live predator, while L-PPO animals show no preference (Taylor et al.,
247 2011). Importantly, while these animals show preference for eye use for a stimulus which elicits

248 aggression (mirror image) or defensive behavior (predator), no eye preference whatsoever is
249 observed towards “neutral” stimuli (Y.-S. Kuan et al., 2007) In addition to the lateralization of
250 behavior, some important differences are observed in adult behavior in other domains. In the first
251 minutes of an open tank test, R-PPO fish show less thigmotaxis than L-PPO fish; likewise, R-PPO
252 fish spend more time near a predator towards the end of the task (Aizawa et al., 2011; Beretta et al.,
253 2012; Concha et al., 2009; Okamoto et al., 2011). In the *frequent-situs-inversus* lineage, larvae with
254 a left-lateralized habenula (LH) begin to view their mirror image with the right eye, but then change
255 to the left eye by the end of a five-minute period; in larvae with a right-lateralized habenula (RH),
256 this pattern is reversed (Dadda et al., 2010; Domenichini et al., 2011). When adults are confronted
257 with a two-choice bead test, LH *fsi* fish bite the right target, while RHT *fsi* fish bite the left target. In
258 an emergence test, when LH *fsi* fish are confronted with a black stripe they show progressively
259 increasing latencies to change compartment; while this effect is observed in RH *fsi* fish, it is much
260 smaller (Barth et al., 2005). These results seem to suggest that a right-lateralized habenula decreases
261 responsiveness to novel and threatening stimuli.

262 Contrary to this hypothesis, however, zebrafish expressing the tetanus toxin light chain in
263 the dHbL [Tg(*narp:GAL4^{VP16}*; UAS:TeTxLC) lineage] show more bottom-dwelling in a novel tank,
264 as well as increased freezing responses to an alarm substance combined with the presentation of a
265 moving shadow above the tank (Mathuru and Jesuthasan, 2013); nonetheless, *c-fos* or *egr1*
266 expression is not significantly changed in the dHb after exposure to an alarm substance in zebrafish
267 (DeCarvalho et al., 2013). On the other hand, in larvae, expression of botulin toxin light chain
268 [Tg(*gng8: Gal4*; UAS:BoTxBLC-GFP)] in the dHbL (but not in the dHbR) impairs light preference
269 (Zhang et al., 2017), consistent with decreased anxiety (Steenbergen et al., 2011). Adult animals
270 with silenced dHbL show increased dark preference (Zhang et al., 2017), consistent with *increased*
271 anxiety (Maximino et al., 2010a). Moreover, exposure to the light/dark assay, a model of anxiety-
272 like behavior, increased *cfos* mRNA expression in the dHb of zebrafish, but only if they are handled

273 before the experiment (Lau et al., 2011). A subset of neurons in the dHbL, expressing the myosin
274 phosphatase Rho-interacting protein *mprip*, shows increased *cfos* mRNA expression after an electric
275 shock (Pandey et al., 2018). These results suggest that the zebrafish dHbL is an important center in
276 defensive behavior, being necessary to mount an adaptive response to innate aversive stimuli; in
277 adults, dHbL appears to inhibit anxiety- and fear-like behavior, while in larvae it appears to increase
278 it.

279 Other important pharmacogenetic work attempted to establish the role of the habenula in
280 stimulus appraisal and behavioral control in zebrafish. In the first work, Agetsuma and colleagues
281 (2010) expressed the tetanus toxin light chain [Tg(*narp:GAL4^{VP16}*; *UAS:TeTxLC*)] or a
282 nitroreductase-mCherry fusion protein in dHbL neurons, and were thus able to block synaptic
283 transmission in that area. When they reached maturity, animals were trained in a cued fear
284 conditioning task; while controls showed increased flight behaviors to the cue after conditioning,
285 dHbL-silenced fish instead showed persistent freezing to the cue. In another experiment, Lee et al.
286 (2010) expressed the photo-sensitizer KillerRed in a ventral telencephalon-habenula projection, and
287 photobleaching on the left habenula led to deficits in the acquisition of two-way avoidance and
288 hyperarousal after light onset before conditioning. They also expressed TeTxLC in the dHbL,
289 obtaining the same result regarding avoidance conditioning deficit.

290 Interestingly, the dHbL has also been implicated in social conflict resolution. Chou et al.
291 (2016) showed that, after the establishment of social dominance, loser (socially submissive) fish
292 show intense activity in the ventral IPN and median raphe and a very reduced responsiveness of the
293 dorsal IPN after acute electrical stimulation of the Hb. These results suggested a participation of
294 different dHb-IPN pathways in the behavioral plasticity that is associated with the losing
295 experience. Indeed, zebrafish expressing the tetanus neurotoxin in the dHbM
296 [Tg(*gpr151:GAL4VP16*; *brn3a-hsp70:GFP-Cre*; *UAS:loxP-DsRed-loxP-GFP-TeNT*)] show a
297 consistent trend to win fights, while dHbL-silenced [Tg(*narp:GALVP16*; *UAS:TeNT*)] fish showed a

298 consistent trend to lose fights (Chou et al., 2016). The authors proposed that the reduction in the
299 dHbL-iIPN-GC pathway could switch behavior from offensive behavior to defensive behavior (i.e.,
300 from attacking to fleeing), resulting in losing the fight. Conversely, reduction in the dHbM-vIPN-
301 MRN pathway results in winning the fight by disinhibiting the MRN, which would tend to increase
302 resilience to aversive stimuli.

303 The role of the vHb has been less extensively studied. An interesting exception is the work
304 of Amo et al. (2014), which demonstrated two types of responses in vHb neurons after fear
305 conditioning. One type responds phasically to an unconditioned stimulus (US) before conditioning,
306 a response which is gradually substituted by sustained firing for all duration of the presentation of
307 the conditioned stimulus (CS). A second type responds phasically to the US and gradually substitute
308 this response to phasic firing to the CS. Amo et al. (2014) suggested that these responses code for
309 aversive expectation value and prediction error, respectively. Multi-unit activity of the vHb
310 increases in the early stages of active avoidance conditioning and later returns to normal, suggesting
311 that the vHb encodes the negative reward expectation value in active avoidance learning (Amo et
312 al., 2014). Expression of TeTxLC in the vHb abolishes active avoidance learning without alterations
313 in basal anxiety-like behavior or fear conditioning, strengthening the hypothesis of a specific role in
314 active avoidance conditioning; finally, pairing the optogenetic activation of the vHb with a specific
315 tank compartment elicits avoidance of that compartment only when the stimulation is tonic, but not
316 when it is phasic. Overall, these results suggest that tonic responses in the zebrafish vHb represent
317 an aversive expectation value, participating in a larger vEP-vHb-MRN circuit (Amo et al., 2014).

318 Another important evidence for the role of the vHb comes from work with kisspeptin, a
319 system of peptides associated with reproductive behavior. In zebrafish, the neuropeptide Kiss1 is
320 expressed only in the vHb, while Kiss2 is expressed in the hypothalamus (Servili et al., 2011).
321 These Kiss1-positive co-express glutamate, and project to glutamatergic interneurons in the IPN
322 and MRN (Nathan et al., 2015a). Interestingly, kissr1 receptors are not found in the raphe, but can

323 be found in the vHb and the IPN (Ogawa et al., 2012). Intracranial administration of Kiss1 increases
324 c-Fos expression in the vHb and MRN, and increases the expression of serotonergic system-related
325 genes (*pet1* and *slc6a4a*) (Ogawa et al., 2014, 2012). It is not known whether CAS induces c-Fos
326 expression in the vHb, since DeCarvalho et al. (2013) only reported results from the dHb. Kiss1
327 also induces a dose-dependent effect on vHb neurons, with low concentrations leading to
328 depolarization and high concentrations leading to hyperpolarization (Lupton et al., 2017).
329 Importantly, Kiss1 blocks responses to CAS (Ogawa et al., 2014), an effect which is mediated by 5-
330 HT_{1A} and 5-HT₂ receptors (Nathan et al., 2015b). Paradoxically, Kiss1 mutant zebrafish with a stop
331 codon upstream of the active peptide show impaired performance in an active avoidance task
332 (Lupton et al., 2017). Interestingly, Kiss1 mutants also show blunted calcium responses in the
333 superior raphe after an electric shock (Lupton et al., 2017). These results underline a mechanism by
334 which the vHb-MRN circuit works: aversive stimuli activate Kiss1/glutamatergic neurons in the
335 vHb, which project to excitatory interneurons in the IPN and raphe. This feed-forward mechanism
336 induces the activation of the raphe, encoding expectations of dangerous outcomes. These
337 expectations can be compared to real outcomes by the activation of 5-HT_{1A} and 5-HT₂ receptors,
338 which have been shown to be important in controlling aversive behavior in zebrafish (Maximino et
339 al., 2014, 2013b; Nowicki et al., 2014). Lupton et al. (2017) suggested that this excitatory stage,
340 associated with increased aversive expectation, favors CS-US association, while the next stage in
341 avoidance learning is mediated by kisspeptin-evoked inhibition of the habenula and consequent
342 reduced aversive expectations once the strategy to avoid the US has been learned.

344 4. Hypothalamic circuits for defense

345 The hypothalamus of teleosts fish presents pair of ventrolaterally extending hypothalamic
346 lobes. These lobes ranges from moderately elevated lobes up to hemisphere-like corpora reaching
347 almost the size of the optic tectum (Senn, 1981). The use of diverse model systems to study

348 hypothalamus development has provided evidence that the molecular pathways regulating
349 hypothalamic induction and patterning are generally conserved from fish to mammals. In addition,
350 the basic hypothalamic cell types and the codes of gene expression that specify them are also highly
351 homologous throughout vertebrate species (Xie and Dorsky, 2017). The fish hypothalamus contains
352 equivalents to most if not all of the mammalian hypothalamic cell types. The hypothalamic neurons
353 are all located in stereotypical clusters within the ventral diencephalon hypothalamic and neuronal
354 populations that control the pituitary in fish have been conclusively shown to be functionally
355 analogous to their mammalian counterparts (Machluf et al., 2011).

356 In teleost fish, partial homologies for hypothalamic nuclei that are relevant for defensive
357 behavior have been proposed: the preoptic area (POA), for example, is composed of cells of
358 different sizes that form clusters considered homologous to mammalian hypothalamic nuclei
359 (Goodson and Kingsbury, 2013). Groups of magnocellular and gigantocellular cells are considered
360 homologous to the supraoptic nucleus, while the cluster of parvocellular cells is supposed to be
361 homologous to the paraventricular nucleus (Moore and Lowry, 1998). Between the POA and ventral
362 hypothalamic region there is a transition zone called ventral tuberal region (VTN) that is thought as
363 homologous to the mammalian anterior hypothalamus (AH), and in the ventrocaudal part of the
364 hypothalamus is located the anterior tuberal nucleus (ATN), a putative homolog of the mammalian
365 VMH (Forlano et al., 2005; Forlano and Bass, 2011; Goodson, 2005; O'Connell and Hofmann,
366 2011). The AH is a source of vasopressin in the mammalian brain, and activation of these
367 vasopressinergic neurons increases aggressive behavior in rodents (Gobrogge et al., 2007).

368 Hodology also supports the homology of these regions. Both the ATN Several
369 hypothalamic regions of teleosts, including ATN and VTN, are connected to the Vs (Folgueira et al.,
370 2004), which has been proposed as homologous to the medial amygdala (Biechl et al., 2017) or to
371 the extended central amygdala (Maximino et al., 2013a) – regions which, as discussed above,
372 participate in the processing of aversive stimuli and defensive responses of both mammals and

373 teleosts. Thus, it is possible that these connections of hypothalamic regions to Vs are related to
374 regulation of defensive behavior in fish.

375 Unfortunately, no functional studies have been made assessing the role of the ATN and
376 VTN in behavioral responses. In teleosts, arginine vasotocin (AVT, the homologue to vasopressin)
377 is also expressed in the VTN (Rodriguez-Santiago et al., 2017), and manipulating AVT levels
378 decreases antipredator behavior in zebrafish (Braidá et al., 2012). However, AVT is also expressed
379 in other brain regions, including the POA and pallial and subpallial amygdala (Rodriguez-Santiago
380 et al., 2017), precluding any speculation as to the role of the VTN. The ATN of teleosts also
381 expresses sex steroid hormone receptors, as is the case of the VTN (O'Connell and Hofmann,
382 2012). Stimulation of the ATN elicits reproduction-related vocalizations in male midshipman fish
383 *Porichthys notatus* (Goodson and Bass, 2000), suggesting a role in reproductive behavior; however,
384 a role in defensive behavior has not yet been determined.

385 Indirect evidence for a participation of the hypothalamus in defensive behavior is stronger
386 in relation to neuroendocrine endpoints, especially cortisol responses, given that these responses are
387 under descending hypothalamic control. Aversive stimuli which have been shown to elicit cortisol
388 responses in zebrafish include acute chasing stress (de Abreu et al., 2016; Idalencio et al., 2017;
389 Tran et al., 2014), acute restraint stress (Abreu et al., 2017a; Ghisleni et al., 2012), unpredictable
390 chronic stress (Piato et al., 2011), exposure to the novel tank or the light/dark test (Kysil et al.,
391 2017), and subordinate-dominant interactions (Pavlidis et al., 2011). CAS also elicits increases in
392 cortisol responses in Nile tilapia (Silva et al., 2015) and zebrafish (Abreu et al., 2017a; Schirmer et
393 al., 2013); interestingly, cortisol responses are also observed after visual contact with a predator in
394 zebrafish (Barcellos et al., 2010, 2007), and *D. rerio* also appears to be able to communicate
395 predation risk to conspecifics, since cortisol responses are observed after seeing a shoalmate
396 displaying antipredator behavior (Oliveira et al., 2017, 2013).

397 Presently little is known about either a causal relationship between cortisol and behavioral
398 responses to aversive stimuli in fish, or about which hypothalamic regions are involved in these
399 responses. A clue comes from Ziv et al. (2013) and Griffiths et al. (2012), which used a mutant
400 zebrafish lineage (*gr^{s357}*) with non-functional glucocorticoid receptors. Larvae from this line show
401 elevated whole-body cortisol levels, increased expression of pro-opiomelanocortin, and no
402 suppression of stress markers after dexamethasone treatment (Griffiths et al., 2012). These larvae
403 also show more auditory-evoked startle responses when compared to wild-type controls, an
404 phenotype that is rescued by treatment with fluoxetine (Griffiths et al., 2012). Adult *gr^{s357}* animals
405 show abnormal freezing behavior when introduced to a novel tank, reduced exploratory behavior,
406 and lack of habituation to environmental novelty (Ziv et al., 2013); chronic treatment with
407 fluoxetine rescue this phenotype and suppresses the stress-induced upregulation of the
408 mineralocorticoid receptor and the serotonin transporter *scl6a4a* (Ziv et al., 2013). These results
409 suggest a developmental role for the glucocorticoid receptor on shaping the serotonergic system
410 and, as a consequence, increasing anxiety-like behavior in both larvae and adults. The acute effect
411 of cortisol is unknown in this species; in goldfish, cortisol rapidly increases Mauthner cell
412 excitability (Bronson and Preuss, 2017), suggesting an acute, non-genomic mechanism to facilitate
413 the brainstem escape network (see below). Finally, in the crucian carp *Carassius carassius*, the
414 CRF₁ receptor antagonist antalarmin suppresses CAS-elicited responses (Lastein et al., 2008),
415 suggesting a role for CRF in these responses; however, it is not known whether this response is
416 mediated by the hypothalamus or by extra-hypothalamic sites, such as the Dm.

418 **5. Tectal circuits for detection of visual threatening stimuli**

419 The most prominent structures within the fish tectum are the optic tectum (TO) and torus
420 semicircularis (TS); they are homologous to the mammalian superior and inferior colliculi,
421 respectively (Nieuwenhuys et al., 1998). Zebrafish present at least six easily identifiable tectal

422 layers (from superficial to deeper: marginal [MS], optic [SO], superficial fibrous and gray [SFGS],
423 central gray [SGC], central white [SAC], and periventricular strata [SPV]). It has been observed
424 that, in all vertebrates, the upper layers of the tectum are retinorecipient, while the deeper layers
425 house the projection neurons (Butler and Hodos, 2005). In rodents, information from the upper
426 visual field is represented in the medial optic tectum, while information from the lower visual field
427 is represented in the lateral portion; likewise, stimulation of the lateral portion leads to approach-
428 like and appetitive movements, while stimulation of the medial portion leads to defensive-like
429 behavior (Brandão et al., 2003, 1999). In rodents, these medial regions receive exclusive projections
430 from multimodal and association sensory cortices, visual thalami, hypothalamic nuclei associated
431 with defensive behavior, and a few pretectal nuclei (Comoli et al., 2012). In goldfish, the medial
432 tectal zone seems to be related to orienting responses, the anteromedial zone to goal-directed
433 saccades, the extreme anteromedial zone to eye convergence, and the posterior zone to escape
434 responses (Herrero et al., 1998; Salas et al., 1997).

435 Many different interneurons have been described in the OT (Figure 2A). Superficial
436 interneurons (SINs) in the superficial layers (SO and SFGS) are GABAergic (Scott and Baier,
437 2009). About 75% of the cells from the periventricular stratum (SPV) are GABAergic, while 10%
438 are glutamatergic (Nevin et al., 2010; Robles et al., 2011). Glutamatergic cells from the tectum are
439 of the bistratified periventricular (bsPVIN) interneuron type, with somata which locate in the deeper
440 and intermediate regions of the SPV, a single apical process that spans the SGC and SFGS, and
441 glutamatergic axons which terminate in the layer between both of these strata (Nevin et al., 2010;
442 Robles et al., 2011). Another group of cells, the non-stratified periventricular (nsPVIN) type, is a
443 small population of GABAergic interneurons located deep in the SPV, arbor in the deeper regions of
444 the SGC and SFGS; they lack the stratification and laminar specificity of bsPVIN cells, with their
445 axons terminating mostly in the SGC and in between this stratum and the overlying SFGS and the
446 underlying SAC (Nevin et al., 2010; Robles et al., 2011; Scott and Baier, 2009). Finally,

447 periventricular projection neurons (PVPNs) are GABAergic cells with a dendritic arbor spanning
448 the regions between SFGS and SGC, the SGC itself, and the region between SGC and SAC, and an
449 axon that forms a sparse arbor of collaterals in the vicinity of the lateral longitudinal fascicle and
450 the hindbrain escape network (Nevin et al., 2010; Robles et al., 2011; Scott and Baier, 2009). Some
451 periventricular projection cells course medially and terminate in the superior raphe (Nevin et al.,
452 2010), but it is not known whether these cells are GABAergic or not; however, lipophilic dye
453 injection into the tectal neuropil marks a much larger population of axons than the GABAergic
454 neurons, suggesting that this is a small and very specialized projection (Nevin et al., 2010; Scott and
455 Baier, 2009).

456 bsPVINs, nsPVINs and PVPNs have been thought of as representing different parts of a
457 circuit for detecting visual threats and selecting appropriate responses (Fig. 2B). The distribution of
458 the neuropil, as well as the glutamatergic nature of the bsPVIN neuron type suggest that these cells
459 perform superficial to deep information transfer (Nevin et al., 2010; Robles et al., 2011). In contrast,
460 the inhibitory nsPVIN is a GABAergic interneuron, producing feed-forward inhibition to filter the
461 visual information transmitted onto projection neurons (Robles et al., 2011). Moreover, the location
462 of their dendrites primarily in the SGC layer, which receives non-visual afferents from the
463 telencephalon and thalamus, suggest that they might integrate inputs from visual and non-visual
464 areas (Nevin et al., 2010). Since projection cells from the tectum have dendrites in the deeper and
465 intermediate layers of neuropil (but not the superficial layers), this suggests that they are not
466 directly retinorecipient (Butler and Hodos, 2005), and therefore must receive information from
467 bsPVIN and nsPVIN cells.

468 In zebrafish, the role of these circuits in escape responses has been described in a series of
469 elegant experiments in larvae. Zebrafish larvae respond to a moving dot stimulus in a size-
470 dependent way: if the stimulus is relatively small (e.g. potential prey), the animal approaches it,
471 while larger stimuli (e.g., potential predators) are avoided (Barker and Baier, 2015). Silencing tectal

472 neurons completely abolishes both approach and avoidance responses, irrespective of stimulus size.
473 When SINs are ablated, however, large object avoidance is impaired (Barker and Baier, 2015).
474 Interestingly, ablating tectal cells marked in the *Gal4mpn354* line – most of which are glutamatergic
475 nsPVINs – shift behavior from approach to avoidance, with ablated larvae avoiding small stimuli;
476 optogenetic stimulation of these neurons produce the opposite effect (Barker and Baier, 2015).
477 Noteworthy, larvae acutely treated with fluoxetine, therefore increasing serotonergic tone, decrease
478 the probability of avoiding small and medium stimuli, but do not affect responses to larger stimuli
479 (Filosa et al., 2016), suggesting that this neurotransmitter does not participate in the control of
480 escape responses by the OT, but instead participates in shifting from fleeing to foraging.

481 Using a different stimulus – a “looming” dot that increases in size, simulating a predator
482 strike – it was demonstrated that zebrafish larvae attempt to escape this stimulus (Dunn et al., 2016;
483 Temizer et al., 2015), and that this stimulus activates three specific targets of retinal ganglion cells:
484 arborization fields AF6 and AF8, and the OT (Dunn et al., 2016; Temizer et al., 2015). In the OT,
485 responses were observed in the SFGS and SGS (Temizer et al., 2015); a specific response is
486 observed in SINs (Dunn et al., 2016). It appears that these SINs modulate the inputs from retinal
487 ganglion cells to periventricular projection neurons, “fine-tuning” the motor command produced by
488 these later cells to critical angle (Dunn et al., 2016). The higher density of looming-selective
489 responses in the OT in relation to extra-tectal retinorecipient areas suggest that the OT is involved
490 in processing looming stimuli, while AF6 and AF8 process other visual cues such as whole-field
491 motion and luminance changes. Indeed, ablation of retinotectal projections – leaving intact
492 projections to other AFs – impaired the ability of larvae to escape the looming stimulus (Temizer et
493 al., 2015). Ablation of the brainstem escape network – one important motor output from OT –
494 resulted in a specific bend deficit in response to looming stimuli, suggesting a participation of these
495 neurons in the escape response elicited by looming stimuli (Dunn et al., 2016).

496

497 **6. The brainstem escape network**

498 The concept of a “brainstem escape network” was introduced to describe gigantocellular
499 neurons in the brainstem of fish (the Mauthner [M-]cells, its two segmental homologs MiD2cm and
500 MiD3cm, and other identified neurons in the reticulospinal segments adjacent to the Mauthner cells)
501 that activate fast-start responses that are used by fish to escape predatory attacks (Eaton et al.,
502 2001). These reticulospinal system receives massive primary acoustic input as well as sensory
503 inputs from the optic tectum, and synapses on motoneurons that innervate trunk muscle on the
504 contralateral side (Kinkhabwala et al., 2010). The activation of M-cells produces a robust turn of
505 about 45°, leading to the initiation of a very fast response called C-start (Furukawa and Furshpan,
506 1963; Eaton et al., 1977) that is fine-tuned to the angle of stimulation; the participation of the other
507 components of the brainstem escape network code other kinematic features that result in propelling
508 the fish away from the stimulus (Eaton et al., 2001). These responses are very fast; in zebrafish, the
509 latency for a C-start after acoustic stimulation was recorded as about 5 ms (Eaton et al., 1977), and
510 the latency for looming visual stimuli varied from 10-20 ms (Temizer et al., 2015) to a few
511 hundreds of milliseconds (Dunn et al., 2016).

512 The brainstem escape network receives inputs from many different regions of the teleostean
513 brain. As described, escape responses to visual stimuli are mediated by the optic tectum (Dunn et
514 al., 2016; Temizer et al., 2015), which projects to this system. While telencephalic projections have
515 not yet been described, telencephalic ablation decreases startle probability in goldfish (Collins and
516 Waldeck, 2006), suggesting a facilitatory role; whether this is due to ablation of pallial or subpallial
517 amygdalar components is unknown.

518 Monoaminergic inputs are also important in the modulation of C-starts. In zebrafish larvae,
519 tyrosine hydroxylase and 5-HT immunoreactivity was observed closely apposed to the ventral
520 dendrites of the M-cell, MiD2cm, and MiD3cm, and tyrosine hydroxylase immunoreactivity was
521 observed near the lateral dendrite (McLean and Fetcho, 2004). 5-HT increases inhibitory currents

522 produced by activation of presynaptic pathways, while dopamine increases the amplitudes of
523 electrical and glutamatergic components of auditorily evoked responses (Korn and Faber, 2005).

524 While this apparently simple reflex has been considered to be a “fixed action pattern”, with
525 little modulation by upstream structures and therefore little relevance for fear- and anxiety-like
526 states, there is interesting evidence for considerable plasticity of this system (Medan and Preuss,
527 2014). Larval zebrafish present prepulse inhibition (PPI), in which the probability of a C-start is
528 reduced when it is preceded by a non-startling sound (Burgess and Granato, 2007). This prepulse
529 sound attenuates the synaptic response of M-cells to a subsequent auditory stimulus (Curtin et al.,
530 2013; Medan and Preuss, 2014, 2011). In zebrafish, the PPI is disrupted by apomorphine, a
531 dopaminergic agonist (Burgess and Granato, 2007); in goldfish, apomorphine blocks the prepulse
532 sound-evoked reduction in M-cell membrane resistance (Medan and Preuss, 2011; Neumeister et
533 al., 2008).

534 5-HT has been implicated in social modulation of startle responses in the African cichlid
535 *Astatotilapia burtoni* (Whitaker et al., 2011). In this species, dominant males show increased startle
536 probability and lower escape thresholds when compared to subordinate males (Neumeister et al.,
537 2010), perhaps as a compensation of the increased conspicuity caused by brighter body coloration
538 and higher activity (Medan and Preuss, 2014; Neumeister et al., 2010). The behavioral increases in
539 startle are accompanied by increased excitability of M-cells to auditory stimuli, as well as a
540 reduction on the inhibitory drive (Neumeister et al., 2010). The 5-HT₂ receptor antagonist
541 ketanserin decreases feedback inhibition in subordinate, but not dominant, African cichlids
542 (Whitaker et al., 2011). These represent presynaptic mechanisms, since only 5-HT_{5A} and 5-HT₆
543 receptors are expressed in Mauthner cells (Whitaker et al., 2011).

544 545 **7. Conclusions**

546 Different regions from the rostrocaudal axis appear to be involved in detecting, identifying,
547 processing, and responding to aversive stimuli in fish. In general, threats are detected and processed
548 at the level of the Dm, which may also be responsible for response selection at certain situations.
549 This region is homologous to the mammalian frontotemporal amygdalar cluster (Maximino et al.,
550 2013a). The “classical model” of the role of the amygdala in fear involves aversive learning in the
551 frontotemporal amygdala, while the behavioral output would be mediated by the autonomic/limbic
552 amygdala (Vargas et al., 2012). Recent evidence, however, suggests that the frontotemporal
553 amygdala is involved in encoding emotional events (including aversive stimuli and contexts) with
554 reference to particular sensory features, while the autonomic amygdala encodes the motivational or
555 affective significance (Balleine and Killcross, 2006). This is consistent with our hypothesis that the
556 Dm also is responsible for mounting appropriate coping (active vs. passive) responses.

557 A parallel circuit for negative incentive and coping has been described in the habenula as
558 well. The vHb appears to represent aversive expectation values and modulate aversive behavior via
559 the median raphe nucleus (Amo et al., 2014). Similarly, a dHBM-vIPN-MRN appears to be involved
560 in resilience to aversive stimuli and/or active coping, while the dHBL-iIPN-GC appears to be
561 involved in selecting appropriate responses (Okamoto et al., 2011). So far, it is not known if this
562 circuit is parallel to the (more classical) amygdalar/Dm one, or whether there are neuroanatomical
563 connections between Hb nuclei and Dm and/or Vs, but an indirect projection to the vHb via the
564 ventral entopeduncular nucleus has been described (Lal et al., 2018). Differently from the
565 disinhibitory Dm-Vs circuit, however, the Dm-vEN-vHb circuit is excitatory. It is possible that
566 both circuits run in parallel and compete (or collaborate) to decision-making in the GC.

567 The participation of these pathways needs to be interpreted in the context of a wider
568 modulation of overall brain state, given the relationships between these circuits and downstream
569 neuromodulators, such as dopamine and serotonin. In the case of the habenular circuit, this is more
570 clear, since both the dHb and the vHb appear to produce their effects by modulating activity in the

571 raphe; as a result, modifying serotonergic phasic and tonic responses in projections from the raphe.
572 However, extensive serotonergic projections have been described to the telencephalon of zebrafish,
573 with important projections to the Dm and Vs/Vp (Lillesaar et al., 2009). Therefore, it is highly
574 likely that not only do neuromodulators such as monoamines act as mediators in the responses, but
575 these transmitters also provide important feedback to the more rostral regions involved in threat
576 detection and response selection (Figure 1B)

577 In addition to this “slow” pathways (the “forebrain circuit”), a “quick-and-dirty” pathway
578 for visual aversive stimuli is organized in the optic tectum (the “midbrain circuit”)(Carr, 2015). This
579 midbrain circuit is sensitive to stimulus size and critical angle, and switches from approach (small
580 stimuli probably mean “food”) to avoidance (large stimuli arriving at a specific angle probably
581 mean “predator strike”). This circuit projects to motor regions of the medulla and spinal cord, as
582 well as to the GC, initiating and/or modulating escape responses that are highly stereotypical.
583 Moreover, a putative indirect OT-Dm projection, via the lateral preglomerular nucleus, has been
584 proposed in fish as an analogue of the superior colliculus-pulvinar-amygdala pathway of mammals
585 (Carr, 2015). It is expected that visually threatening stimuli activate the first circuit and bypass the
586 Hb-IPN/raphe-GC or the Dm-Vs-GC circuits. Very little is known about the GC of zebrafish –
587 contrary to the great importance that is given to the periaqueductal gray of mammals as a hub for
588 defensive behavior (see Motta et al., 2017, for a recent review) –, an interesting research
589 opportunity to clarify these circuits.

590 The forebrain circuit is functionally similar to LeDoux’s (1998, 2000) concept of a “high
591 road”, providing slower but highly processed sensory information with affective tones, while the
592 midbrain circuit would be functionally similar to the “low road”. In both the proposed forebrain
593 circuit and the “high road”, threatening stimuli pass through increased processing of the nature of
594 the sensory stimuli to reach a more precise identification; however, in LeDoux’s original
595 formulation the high road refers to cortical processing of sensory information before reaching the

596 amygdala, while the low road refers to information that is sent directly from the thalamus to the
597 amygdala. Not only is the isocortex not the main target of sensory thalamic projections in zebrafish,
598 the participation of putative homologues in aversive behavior has not been established (but see Aoki
599 et al., 2013). Indeed, Carr (2015) suggested that a “high road” is absent in fish, which would only
600 possess a “low road”. The present review, however, suggest that some granularity in this
601 interpretation is necessary, given that a differentiation between responses mediated by the forebrain
602 and midbrain circuits is possible.

603 The participation of the habenular pathways in neurovegetative responses is also less clear,
604 but a amygdalar-hypothalamic circuit has been suggested on the basis of c-Fos and p-ERK activity
605 (Faustino et al., 2017; Randlett et al., 2015). Aversive stimuli, including CAS and physical stressors,
606 induce cortisol (Abreu et al., 2017b; Schirmer et al., 2013) and norepinephrine and epinephrine
607 release (Maximino et al., 2014). This is consistent with a classical model of amygdalar control of
608 fear in rodents, in which CEXA-hypothalamic projections regulate the neurovegetative responses to
609 threats (Misslin, 2003). Based on coherence analysis of c-Fos expression to CAS (Faustino et al.,
610 2017), it is suggested that the Dm activates inhibitory Vs neurons, disinhibiting downstream
611 hypothalamic mechanisms.

612 What is the “switch” that regulates active vs. passive coping? What conditions favor
613 switching from the forebrain to the midbrain circuits? While the neural bases of these changes are
614 presently unknown, environmental characteristics which lead to decision-making have been
615 described thoroughly. For example, threat probability and distance vary in a continuum that restricts
616 attention (Andersen et al., 2016), and therefore whether fast or slower responses are needed (Brown
617 et al., 1999; Fanselow and Lester, 1988; Kavaliers and Choleris, 2001; Laundré et al., 2010;
618 McNaughton and Corr, 2004; Perusini and Fanselow, 2015). Thus, decision-making is biased
619 towards escape (flight) or fight responses when the threat is proximal, while avoidance and freezing
620 are elicited when threat is distal (Fanselow and Lester, 1988; McNaughton and Corr, 2004; Perusini

621 and Fanselow, 2015). Similarly, the decision to freeze or flee is dependent on environmental
622 affordances, such as the availability of escape routes (Blanchard and Blanchard, 1988). It has been
623 suggested that habenular circuits act as comparators for these stimuli, biasing the organism towards
624 careful approach or escape (Okamoto et al., 2011). An exciting possibility is that habenular circuits
625 produce this effect by modulating the activity of serotonergic projections to the Dm and/or Vs and
626 to the OT. This hypothesis is consistent with the Deakin/Graeff hypothesis for the role of serotonin
627 in switching responses from escape towards risk assessment (Deakin and Graeff, 1991; Graeff et al.,
628 1997; Maximino, 2012).

629 Much work is still needed to identify the regions that are responsible for aversive behavior
630 in fish. The present review suggests some of the regions involved and presents a roadmap and
631 framework for future research. The evolutionary relevance is clear, but it is also important to
632 consider the implications of these findings for work in the field of behavioral models and
633 experimental psychopathology: establishing homologies between regions involved in similar
634 behavior strengthens the hypotheses that these behaviors are indeed conserved, an assumption of
635 most models that is rarely tested (Maximino et al., 2010b) and of which depends the construct
636 validity of these models. This implication opens up novel avenues for future research which ought
637 to be prolific.

638

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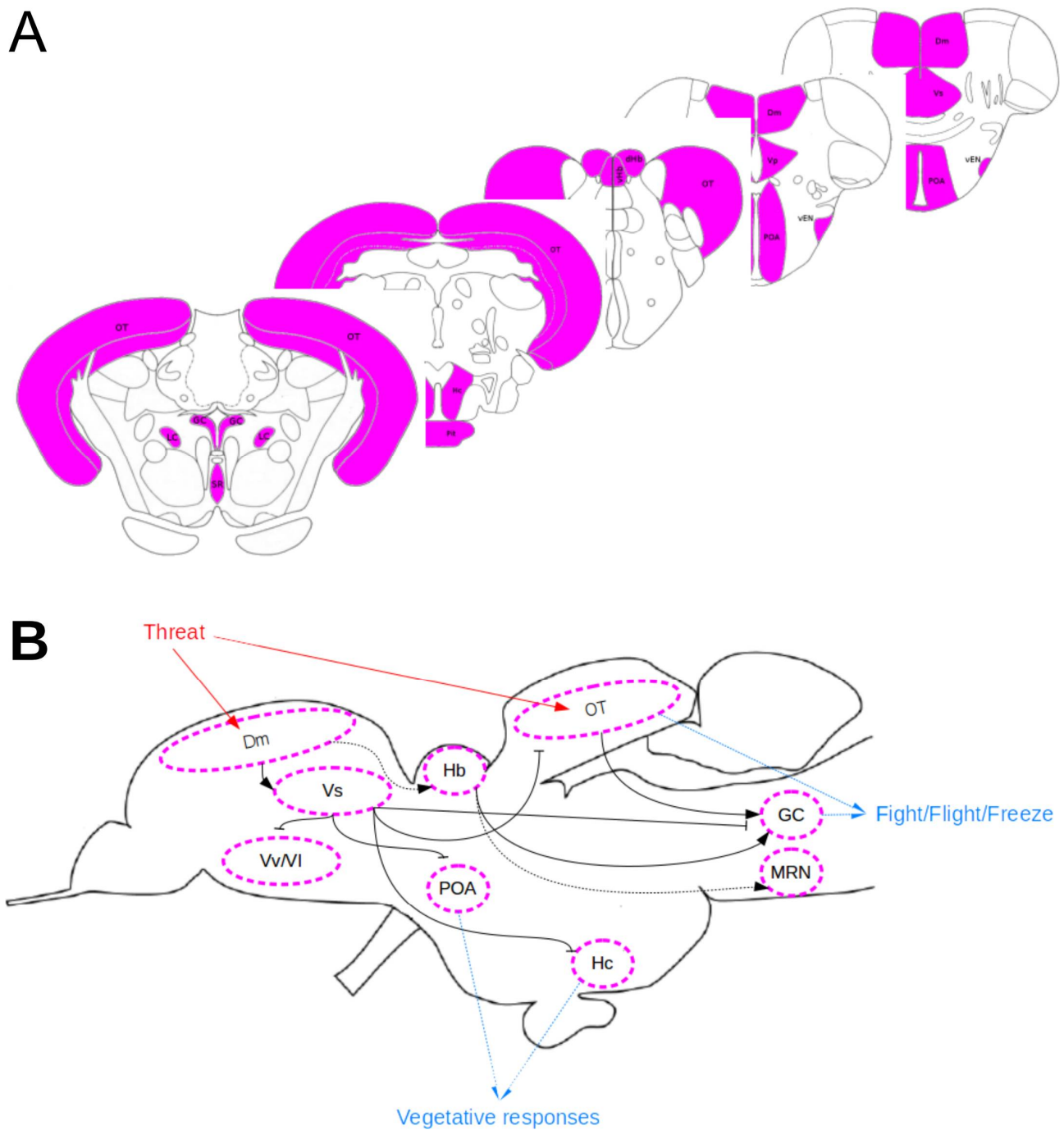
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1127 **Figure 1** – (A) Frontal view of the zebrafish brain circuits for threat detection, identification, and
1128 processing, and response selection (in purple), described in Fig. 1B. The right-to-left plans represent
1129 the rostrocaudal axis. Schemes adapted from Wulliman et al., 1996. (B) Circuits for threat detection,
1130 identification, and processing and response selection in the teleost brain. The dorsomedial
1131 telencephalon (Dm) is homologous to the frontotemporal amygdala system, and is the entry for the
1132 “forebrain circuit” that terminates in the mesencephalic central gray (GC) and in the hypothalamic
1133 circuits for neurovegetative responses (POA, Hc). An indirect projection is also depicted for the
1134 ventral habenula (Hb); this projection involves a feedforward glutamatergic projection from the Dm
1135 to the entopeduncular nucleus, and from there to the ventral habenula. This can represent part of a
1136 habenular circuit in the forebrain circuit which projects indirectly to the raphe (MRN) serotonergic
1137 neurons via glutamatergic interneurons in the interpeduncular nucleus and raphe (not shown). The
1138 “midbrain circuit” is represented by projections to interneurons in the optic tectum (OT) which
1139 mediate fast escape responses via projections to the GC and/or the brainstem escape network (not
1140 shown). Threatening stimuli (especially visual stimuli) can bypass the forebrain circuit and directly
1141 activate this system, initiating “quick-and-dirty” escape responses.

1142 Abbreviations: Dm: medial nucleus of the dorsal telencephalon; Vs: supracommissural nucleus of
1143 the ventral telencephalon; Vv: ventral nucleus of the ventral telencephalon; Vl: lateral nucleus of
1144 the ventral telencephalon; Hb: habenula; POA: preoptic area; Hc: caudal hypothalamus; OT: optic
1145 tectum; GC: central gray; MRN: median raphe nucleus.

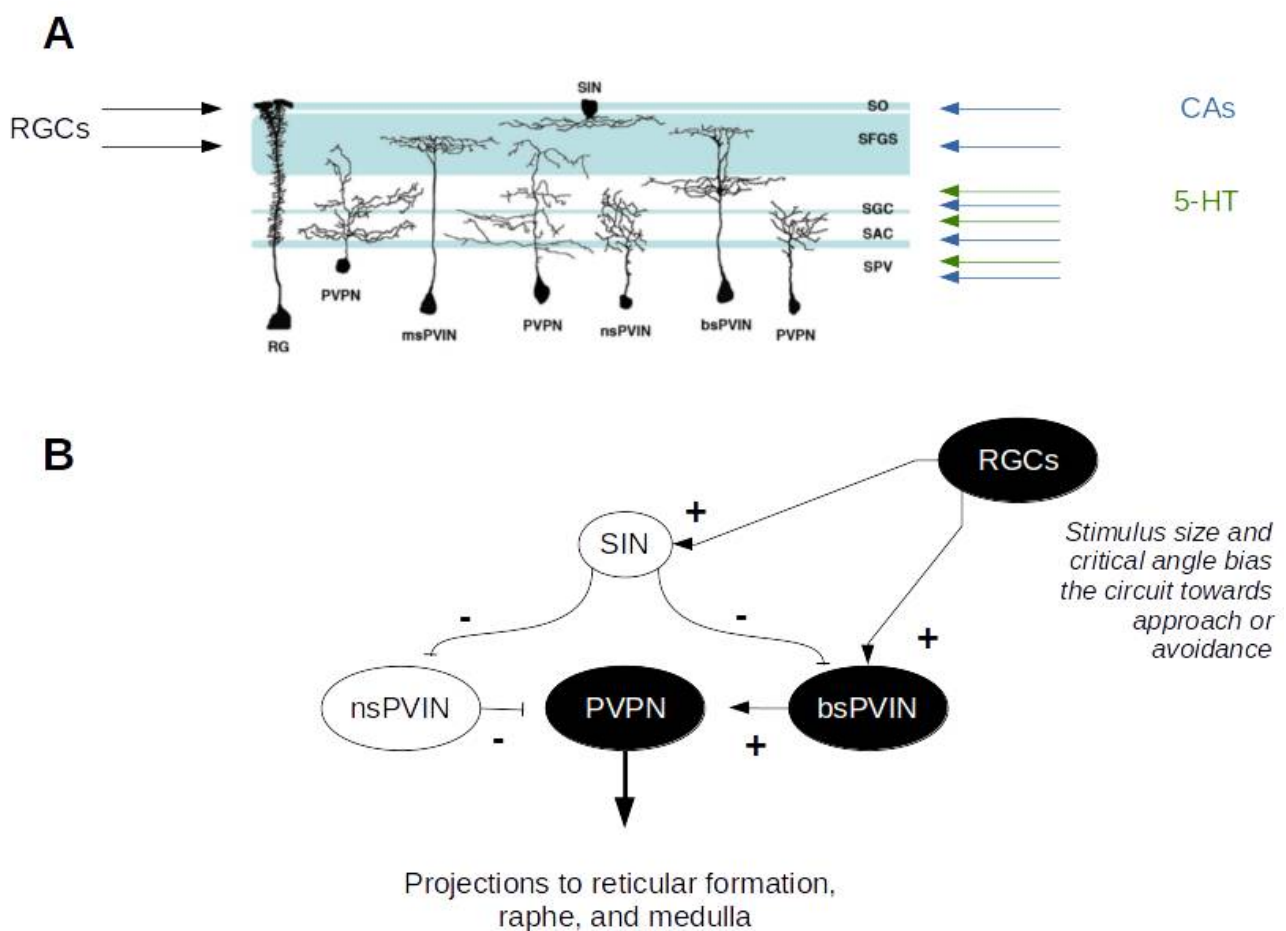
1146 Full black arrows: direct projections; dashed black arrows: indirect projections. Arrows terminating
1147 in dimension lines (⊥) represent inhibitory projections. Blue arrows represent outputs, while red
1148 arrows represent inputs.



1149

1150 **Figure 2** – (A) Depiction of the tectal interneurons that participate in the escape circuit. The upper
 1151 layers of the tectum (stratum opticum and stratum fibrosum et griseum superficiale) are
 1152 retinorecipient, and also receive catecholaminergic (CAs) projections. Interneurons in the deeper
 1153 layers (stratum griseum centrale, stratum album centrale, stratum periventriculare) participate in
 1154 decision-making, and receive serotonergic (5-HT) and catecholaminergic projections. Identified
 1155 interneuron types include the superficial inhibitory interneuron (SIN), as well as a variety of

1156 periventricular interneurons (non-stratified [nsPVIN], bistratified [bsPVIN], and mono-stratified
 1157 [msPVIN] interneurons). Morphologically distinct periventricular projection neurons (PVPNs) have
 1158 also been identified. (B) Stimulus properties, including size, are coded by retinal ganglion cells
 1159 (RGCs) bias the circuit towards approach (smaller size) or avoidance (large size). (B) These stimuli
 1160 activate inhibitory SINs and excitatory bsPVINs. SINs also fine-tune responses of bsPVINs, and
 1161 disinhibit PVPNs by inhibiting nsPVINs. These projection neurons, in its turn, modulate the activity
 1162 of the reticular formation, raphe, and medulla.



1163

1164

1165 **Table 1** – Neuromodulatory (monoaminergic) innervation of nuclei in the aversive brain system of
 1166 teleosts. Given that monoaminergic regions such as the raphe are important targets of forebrain and
 1167 midbrain circuits, it is likely that modulating the activity of these neuromodulators represent ways

1168 to “fine-tune” defensive responses, including by feedback. The number of ‘+’ symbols represent the
 1169 relative amount of innervation in these regions.

1170 Abbreviations: 5-HT: serotonin; DA: Dopamine; NE: noradrenaline; Dm: medial nucleus of the
 1171 dorsal telencephalon; Vs: supracommissural nucleus of the ventral telencephalon; Vv: ventral
 1172 nucleus of the ventral telencephalon; Vl: lateral nucleus of the ventral telencephalon; vEN: ventral
 1173 entopeduncular nucleus; Hb: habenula; POA: preoptic area; Hc: caudal hypothalamus; OT: optic
 1174 tectum; GC: central gray.

	5-HT innervation	DA innervation	NE innervation
Dm	+	0	
Vs	+	+	++
Vv	+	++	++
Vl	+	+	++
vEN	++	0	?
Hb	++	0	0
POA	+++	++	+
Hc	+++	+++	+
OT	++	++	+
GC	+++	?	?