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4	Morphogenetic and patterning defects explain the coloboma phenotype of
5	the eye in the Mexican cavefish
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24 Abstract

25 The morphogenesis of the vertebrate eye consists of a complex choreography of cell 26 movements, tightly coupled to axial regionalization and cell type specification processes. Any 27 disturbance in these events can lead to developmental defects and blindness. Here we have deciphered the sequence of defective events leading to coloboma phenotype in the 28 embryonic eye of the blind cavefish of the species Astyanax mexicanus. Using comparative 29 live imaging on targeted enhancer-trap Zic1:hsp70:GFP reporter lines of both the normal, 30 river-dwelling morph and the cave morph of the species, we identified major defects in initial 31 optic vesicle size and optic cup invagination in cavefish. Combining these results with gene 32 33 expression analyses, we also discovered defects in axial patterning affecting mainly the 34 temporal retina, in optic stalk tissue specification, and in the spreading processes involving the retinal pigmented epithelium cells. Based on these results, we propose a developmental 35 scenario to explain the cavefish phenotype and discuss developmental constraints to 36 morphological evolution. The cavefish eye appears as an outstanding natural mutant model 37 to study molecular and cellular processes involved in optic region morphogenesis. 38

40 Introduction

41

The morphogenesis of the vertebrate eye is a complex choreography of cell movements 42 starting from a flat neural plate and leading to the formation of a spherical multi-layered 43 structure. Owing to technological improvements, this process has been increasingly 44 investigated in the last decade, especially on teleost models which are amenable to live 45 imaging due to their external development and transparency (England et al., 2006; Ivanovitch 46 et al., 2013; Kwan et al., 2012; Martinez-Morales et al., 2009; Nicolas-Perez et al., 2016; Picker 47 48 et al., 2009; Rembold et al., 2006; Sidhaye and Norden, 2017). This focus on morphogenesis led to the description of cell and tissue movements during eye development in fish (reviewed 49 in Cavodeassi, 2018) (Fig. S1). 50

At the end of gastrulation, the eyefield is specified in the anterior neural plate, surrounded 51 anteriorly and laterally by the prospective telencephalon, and posteriorly by the future 52 hypothalamus and diencephalon (Varga et al., 1999; Woo and Fraser, 1995; Woo et al., 1995). 53 54 During neurulation, the first step of eye formation is the lateral evagination of the optic vesicles. Already at this step, cell behaviours are complex as some eye-fated cells behave like 55 the nearby telencephalic cells and converge toward the midline to form the neural keel, while 56 others lag behind and keep the eyefield wide (Ivanovitch et al., 2013; Rembold et al., 2006). 57 The eye vesicles then elongate due to a flow of cells entering the anterior/nasal optic vesicle, 58 in a process recently re-described as "extended evagination" (Kwan et al., 2012). 59 Simultaneously, the optic vesicles are separated from the neural keel by the anterior-wards 60 progression of a posterior furrow between them and the diencephalon, leaving a connection 61 with the neural tube at the optic stalk (England et al., 2006). Cells from the medial part (inner 62 leaflet) of the optic vesicle then migrate around the rim of the eye ventricle (the optic recess) 63 64 into the lens facing neuroepithelium through a process called rim movement (Heermann et al., 2015; Kwan et al., 2012). The cells in the outer layer of the optic cup, fated to the retinal 65 pigmented epithelium (RPE), expand and flatten to cover the back of the retina (Cechmanek 66 and McFarlane, 2017; Heermann et al., 2015). Together with the basal constriction of lens-67 facing epithelial cells (Martinez-Morales et al., 2009; Nicolas-Perez et al., 2016), these 68 movements lead to optic cup invagination. The invagination process leads to the formation of 69

70 the optic fissure at the level of the connection of the eye with the optic stalk. This fissure allows bloods vessels to invade the eye and leads the way of retino-fugal axons, but needs to 71 close to have a functional and round eye (Gestri et al., 2018). Finally, the entire eye, together 72 with the forebrain, rotates anteriorly, bringing the optic fissure in its final ventral position. 73 Hence, cells that are initially located in the dorsal or ventral part of the optic vesicles 74 contribute to the nasal or temporal quadrant of the retina, respectively (Picker et al., 2009). 75 Failure to correctly complete any of these steps can lead to vision defects; for example, failure 76 to properly close the optic fissure is termed coloboma and can lead to congenital blindness. 77

During the eye morphogenetic process, three types of tissues emerge: (1) the neural retina, facing the lens and composed of various neuronal types, (2) the RPE at the back of the neural retina, with multiple functions including nurturing of photoreceptors (Strauss, 2018), and (3) the optic stalk, transiently connecting the retina to the neural tube. This thin ventral structure is invaded by the ganglionic cells axons and guides them on their way to the tectum. Optic stalk cells then differentiate into reticular astrocytes surrounding the optic nerve (Macdonald et al., 1997). All these tissues derive from the optic vesicles and have a neural origin.

85 Concomitantly to morphogenesis, retinal cells acquire axial positional identity. Indeed, the 86 visual sense requires a topographic perception of the light stimuli and processing of the signal 87 to form images. In fishes, the neuronal map of the retina is replicated onto the contralateral tectum, in a symmetrical manner so that the nasal retina projects to the posterior optic 88 tectum, while the temporal retina projects to the anterior tectum. This retinotopy requires a 89 precise regional identity of both the retina and the tectum for a proper matching (Rétaux and 90 Harris, 1996; Sperry, 1963). In the retina, several transcription factors such as Vax2, FoxG1, 91 Tbx5a, or FoxD1 present a strong regional expression, already present at optic vesicle stage, 92 93 and define the future quadrants identity by controlling the expression of effector guidance 94 molecules (French et al., 2009; Picker et al., 2009; Sakuta et al., 2006; Schulte et al., 1999). The regionalized expression of the transcription factors themselves is mainly achieved through 95 Hedgehog and Fgf signalling from the embryonic midline, and reciprocal interactions between 96 them (Asai-Coakwell et al., 2007; Hernandez-Bejarano et al., 2015; Kruse-Bend et al., 2012; 97 Picker and Brand, 2005; Picker et al., 2009). Shh is involved in the regulation of the ventral and 98 temporal fates while Fgfs secreted by the anterior neural ridge and olfactory placode seem to 99 100 be involved in nasal specification. Extra-ocular Bmps direct the acquisition of dorsal fates

101 (Gosse and Baier, 2009; Hernandez-Bejarano et al., 2015; Picker and Brand, 2005; Picker et al.,
102 2009; Take-uchi et al., 2003).

Astyanax mexicanus is a teleost fish that arises in two morphs: the classical river-dwelling eyed 103 morph and the cave-dwelling blind morph. Although eyes are absent in adult cavefish, they 104 105 first develop in embryos before degenerating during larval stages. The early cavefish eyes 106 display several morphogenesis abnormalities: the optic vesicles are short (Alunni et al., 2007), the optic cup and lens are small (Hinaux et al., 2015; Hinaux et al., 2016; Yamamoto and 107 Jeffery, 2000) and the ventral part of the optic cup is severely reduced or lacking, leaving the 108 109 optic fissure wide open (coloboma phenotype) (Pottin et al., 2011; Yamamoto et al., 2004). Moreover, cavefish exhibit several modifications of morphogen expression including an 110 expanded Shh expression at the anterior midline, a heterochrony of Fqf8 onset in the anterior 111 neural ridge, and variations of *Bmp4* and *Dkk1b* expression in the prechordal plate (Hinaux et 112 al., 2016; Pottin et al., 2011; Torres-Paz et al., 2018; Yamamoto et al., 2004). These morphogen 113 alterations trigger modifications of the cavefish eyefield and subsequent eye, as evidenced by 114 modified expression of Lhx9/2 or Pax6 in the neural plate when compared to surface fish 115 116 embryos (Pottin et al., 2011; Yamamoto et al., 2004), and they have been linked to cavefish 117 eye development defects: overexpression of Shh in surface fish shortens its optic cups and triggers the apoptosis of the lens, while inhibition of Fgf signalling in a cavefish restores the 118 ventral retina (Hinaux et al., 2016; Pottin et al., 2011; Yamamoto et al., 2004). 119

Because of these various modifications, the cavefish is a remarkable natural mutant model to study eye development, beyond the mechanisms of eye degeneration and loss. We therefore undertook to study the morphogenesis and regionalization of the cavefish eye, in comparison with surface fish, with two aims: a better understanding of the defects of the cavefish embryonic eye as well as the mechanisms of eye morphogenesis in general.

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Using CRISPR/Cas9-mediated targeted enhancer trap cavefish and surface fish *Zic1:hsp70:GFP* lines, we performed comparative live imaging of eye morphogenesis on developing embryos. We propose that the reduction in cavefish optic vesicle size, associated with a proper extended evagination, leads to a bias towards nasal fate and an abnormal final ventral position of the lens. In addition, optic cup invagination is impaired and RPE spreading over the surface of the neural retina is delayed in cavefish. Through gene expression analyses, we show

- moreover that the cavefish eyes display modifications of their axial regionalization, with a
- 133 tendency to preserve or increase nasal and dorsal fates while strongly diminishing temporal
- 134 fate. Finally, the optic stalk fate is widely increased throughout the retina.

136 **Results**

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138 Screening candidate genes to establish Astyanax GFP reporter lines

139 To perform live imaging and label regions of interest, i.e. the entire optic region of cavefish (CF) and surface fish (SF) embryos, we sought to find a reporter gene that would be expressed 140 in the eyefield from the neural plate stage (10hpf) until at least 24 or 36hpf, when optic recess 141 region (ORR) and retina are clearly separated (Affaticati et al., 2015). An in situ hybridization 142 mini-screen for candidate genes selected from publications and the ZFIN expression database 143 144 was performed. Chosen candidates were Vax1, Vax2 (Take-uchi et al., 2003), Zic1 (Hinaux et al., 2016; Maurus and Harris, 2009; Rohr et al., 1999; Tropepe et al., 2006), Zic2a (Sanek et al., 145 2009), Rx3 (Deschet et al., 1999; Rembold et al., 2006; Stigloher et al., 2006), Lhx2 and Lhx9 146 (Pottin et al., 2011). We performed *in situ* hybridization on CF and SF embryos at 5 different 147 stages (10, 12, 14, 24 and 36hpf) (Fig. S2A). 148

Among the 7 genes, 5 were expressed in the anterior neural plate at 10hpf while 2 were not: 149 Vax1 and Vax2, whose expressions were detectable from 12hpf only (Fig. S2A). Five of them 150 were expressed at least partially in the optic vesicle per se (excluding ORR and optic stalk): 151 152 Vax2, Zic1, Rx3, Lhx9 and Zic2a (faintly). At 36hpf, only 4 of them were still expressed in the optic cup: Zic2a and Zic1 (around the lens), Lhx2 (faintly) and Vax2 (in the ventral retina). 153 154 Subtle differences between CF and SF expression patterns were observed (not shown), and only one candidate genes was consistently expressed in the eye from neural plate to 36hpf: 155 Zic1 (Figure 1A and Fig. S2B; see legend of Fig. S2B). Even though the Zic1 pattern was complex 156 and encompassed a region wider than the optic region of interest, it was chosen for 157 transgenesis due to its early and persistent expression throughout the eye and the ORR/optic 158 159 stalk regions.

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161 Comparative expression of *Zic1* in surface fish and cavefish embryos

A closer examination of *Zic1* pattern highlighted some patterning and morphological differences between SF and CF (**Figure 1A**). In the anterior neural plate at 10hpf, *Zic1* expression was wide in the bilateral eyefield in SF, with a medial indentation (**Figure 1A**,

asterisk); in CF, *Zic1* was expressed in narrower lateral bands, with a wider staining anteriorly.
At 12hpf, *Zic1* pattern confirmed that the CF optic vesicles were shorter than those of SF, but
they also looked more "plump". Indeed, from a dorsal view SF optic vesicles were elongated,
slender and pointier while CF optic vesicles looked rounder. On lateral views, the optic vesicles
showed a trapezium shape in SF and an oval shape in CF. At 36hpf the *Zic1*-expressing ORR
was wider in cavefish. These differences in expression patterns decisively convicted us to use *Zic1* as reporter gene.

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173 Establishing Zic1:hsp70:GFP surface fish and cavefish knock-in lines

174 We used a targeted enhancer-trap strategy into the *Zic1* locus to generate reporter lines. This option presented the advantage that the GFP reporter insertion site would be identical in CF 175 and SF lines and avoid positional effects, which is crucial for comparative purposes. The 176 genomic region around Zic1 was examined to find conserved elements that might point 177 toward putative regulatory elements (Fig. 1BC). The region was large (~500 kb in zebrafish) 178 179 and complex. In the zebrafish genome which has a better quality and annotation than the Astyanax genome (McGaugh et al., 2014), Zic1 and Zic4 were very close to each other in a 180 181 head to head configuration, and located in the middle of a gene desert (~200kb downstream 182 of Zic1 and ~220kb downstream of Zic4 in zebrafish, and ~275kb downstream of Zic1 and ~235kb downstream of Zic4 in Astyanax). This gene desert contained many elements 183 conserved among fishes, also partly conserved with tetrapods and mammals (Fig. 1BC). Such 184 a regulatory landscape suggested that the elements driving *Zic1* expression are probably 185 modular and difficult to identify, further strengthening the choice of a directed enhancer-trap 186 approach. 187

Cavefish and surface fish embryos were co-injected with sgRNA2 (targeting the region 188 189 between conserved non-coding elements 1 and 2), Cas9 protein and a linearized minimal promoter *hsp70:GFP* repair construct and were screened at 30hpf for fluorescence patterns 190 consistent with Zic1 endogenous pattern (Fig. 1CD). Excellent Zic1 pattern recapitulation in F0 191 was observed at low frequency (1-2% of injected embryos), while other more partial patterns 192 were also seen at higher frequencies. All potential founder embryos were sorted and raised 193 until males were sexually mature (5-6 months' post-fertilization) and could be screened by 194 individual in vitro fertilization. We detected 3 founders for SF (out of 15 F0 males screened) 195

196 and 5 founders for CF (out of 9 F0 males screened) with various transmission rates: 4%, 7% and 30% for SF founders and 4%, 45%, 48%, 50% and 54% for CF founders, respectively. The 197 fish were screened based on their GFP pattern, matching Zic1 (Fig. 1D). In both morphs some 198 variations in relative intensities of fluorescence were observed, with some lines exhibiting 199 more homogeneous levels of expression and others having an extremely strong GFP 200 fluorescence in the telencephalon and a dimer fluorescence in the eye. We focused on the 201 most homogeneous lines for imaging purposes. Importantly, in those lines, genomic analyses 202 confirmed the proper insertion of the transgene at the exact targeted site (Fig. S3), and double 203 fluorescent in situ hybridisation for Zic1 and GFP mRNAs demonstrated that the reporter fully 204 recapitulated the endogenous *Zic1* pattern at the stages of interest (Fig. 1E). 205

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207 Comparing eye morphogenesis in surface fish and cavefish through live imaging

Live imaging was performed on a light-sheet microscope on both CF and SF lines from ~10.5hpf to 24-30hpf, depending on the acquisitions (**Fig. 2** and **Movies 1 and 2**). The planes used for analysis were chosen to always cross the middle of the lens and the optic stalk (see drawings on **Fig. 2**), in order to follow the anterior rotation of the eye. Overall, eye morphogenesis in SF recapitulated step by step the events described in zebrafish, while in CF the morphogenetic movements were conserved but their relative timing and extent seemed different. The following analyses result from quantifications made on 4 eyes for each morph.

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216 **Evagination and elongation.** The CF optic vesicles were much shorter from the beginning of evagination onwards and they spanned about half the length of the SF optic vesicles at the 217 same stage (139µm compared to 216µm at 11.5hpf) (Fig. 2A,B and Fig. 3A,B). Elongation then 218 proceeded at approximately the same pace as in SF until 17.5hpf (Fig. 3B). However, while the 219 length of the eye primordium decreased between 17.5hpf and 25.5hpf in SF, due to the 220 221 invagination movement, elongation continued at slower pace until 25.5hpf in CF. Hence, the increase in length from 11.5hpf (beginning of the furrow progression to separate the optic 222 223 vesicles from the neural tube in both morphs) to 31.5hpf was more important in CF than in SF, because it remained flatter (Fig. 3B). Moreover, the final size of the SF optic cup was very 224 similar to that of its early evaginating eyefield (252µm at 31.5hpf compared to 240µm at 225

10.5hpf) while in CF a net increase in the optic primordium size was visible (186µm at 31.5 hpf compared to 146µm at 10.5hpf) (Fig. 3A). In addition, while in SF the optic vesicles stayed closely apposed to the neural tube, in CF they first started growing away from the neural tube before getting back closer between 18.5 and 21.5hpf (Fig. 2B). Finally, throughout development, the width of the optic stalk (defined in its wide meaning as the connection between the optic vesicle/cup and the neural tube) was similar in the two morphs despite an initially smaller size in the CF due to the smaller optic vesicles (Fig. S4).

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Optic cup invagination and lens formation. The posterior end of the optic vesicles started 234 235 curling back in both CF and SF around 15.5hpf, probably due to basal constriction. The lens 236 started being identifiable at 17.5 hpf in both morphs (Fig. 2B and movies 1 and 2). At this stage, the lens was in a central position with regard to the antero-posterior extension of the optic 237 238 vesicles, in both morphs (Fig. 2B and 3D). Then, in SF the invagination quickly brought closer the two edges of the optic cup, in contact with the lens (Fig. 2B and Fig. 3C). In contrast, 239 despite initially harbouring a curvature typical of an invaginating optic cup, the edges of the 240 CF optic cup did not undergo the same change of form and stayed "flat", with an apparent 241 impairment of the rim movement in the posterior optic cup (Fig. 2B, Fig. 3C and Movie 2). In 242 243 fact, the CF optic vesicles continued to elongate while the lens remained static, therefore 244 seemingly shifting the lens position anteriorly (Fig. 2B and Fig. 3D). During this period, the posterior part of the CF optic cup showed slow and reduced curling, which in some embryos 245 led to a separation of the posterior optic vesicle from the lens. Yet the posterior (prospective 246 dorsal) optic cup finally curved and contacted the lens (see progression, especially on the right 247 eye from 22.5hpf to 30.5hpf, Movie 2; Fig. 2B and Fig. 3D). However, the CF optic cup 248 remained shallower as its lens, albeit smaller, always bulged out while in SF the lens was 249 250 always contained inside of the optic cup curvature.

In sum, live-imaging experiments suggested that in CF (1) the optic vesicles were reduced in size after the initial evagination, (2) elongation of the optic vesicle occurred properly, including for the extended evagination, while (3) invagination was transiently compromised. We reasoned that these impaired morphogenetic movements should impact the proper patterning and regionalization of the CF retina.

257 Comparing eye regionalization in surface fish and cavefish

258 In order to label the different quadrants of the eye, four classical markers were chosen: FoxG1 for the nasal quadrant, Tbx5a for the dorsal quadrant, FoxD1 as a temporal marker and Vax2 259 as a ventral marker spanning both the nasal and temporal sides of the optic fissure (Fig. 4). 260 The difficulty of comparing these quadrants between CF and SF resided in the difference of 261 262 morphology and size of their eyes. We therefore decided to measure different parameters including angles of expression (taking as a reference the centre of the lens and the middle of 263 the optic fissure if needed) and gene expression areas (expressed either as absolute values or 264 265 as relative values normalized to eye size). Below the expression patterns of the four markers 266 are described in a clock-wise manner starting at the optic fissure.

267 The nasal marker *FoxG1* presented a larger angle of expression in CF (149°) than in SF (132°) due to a dorsal expansion of the expression domain (Fig. 4A-A"). In CF the dorsal marker Tbx5a 268 started at the same angle from the optic fissure but spanned an increased angle towards the 269 temporal retina (angle to mid-fissure: 106° for SF and 111° for CF; span: 108° for SF and 139° 270 for CF) (Fig. 4B-B"). Reciprocally to these two increases of expression domains in the clock-271 wise direction, the temporal marker FoxD1 span was reduced in its dorsal part in CF (179° in 272 SF, 141° in CF) (Fig. 4 C-C"). Finally, the ventral marker Vax2 had different features on the 273 274 nasal and temporal margins of the optic fissure. Vax2 span was increased in the ventro-nasal guadrant, indicating a dorsal-wards or clock-wise expansion (66° in SF, 94° in CF) while it was 275 unchanged in the temporal quadrant when compared to SF (Fig. 4 D-D"). 276

The smaller size of the CF eye was reflected by the absolute values of markers expression areas 277 (Fig. 4E). Indeed, all of them but one (*Tbx5a*) were decreased in size, including nasal *Vax2* and 278 *FoxG1* expression domains which were slightly but significantly reduced. *Tbx5a* was the only 279 gene showing the same area of expression in CF and SF eyes, suggesting that the dorsal 280 281 quadrant was proportionately increased in cavefish. The two temporal genes, FoxD1 and Vax2, exhibited a strong reduction of expression area, pointing to a temporal reduction in the 282 cavefish eye. This conclusion was further supported when the gene expression areas were 283 expressed in relative values normalized to eye size (Fig. 4F) or in percentages of SF labelling 284 (Fig. 4G). Indeed, the most strongly reduced quadrant was the temporal quadrant, labelled by 285 FoxD1 and the temporal aspect of the Vax2 domain. 286

Thus, in CF all markers examined presented modifications of expression in a fan-opening fashion, from nasal towards temporal, overall increasing the nasal and dorsal fates at the expense of the temporal retina. Contrarily to the usual description of the cavefish eye as ventrally-reduced, we unmasked here a temporal reduction.

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292 Comparing tissue and cell identity markers in surface fish and cavefish

To assess tissue identity in the developing retina, *Pax2a* was used as a marker for optic stalk and optic fissure margin identity (Macdonald et al., 1997) and *Bhlhe40* (Cechmanek and McFarlane, 2017) was used as a RPE identity marker.

Pax2a showed wider expression in cavefish, expanding beyond the optic fissure margins and occupying a large part of the ventral quadrants both nasally (angles, SF: 36°, CF: 67°) and temporally (angles, SF: 21°, CF: 55°) (**Fig. 5A-A''**). Surprisingly, in CF eyes *Pax2a* expression sometimes expanded throughout the retina with a lighter, although specific, expression intensity. Staining was even sometimes present in the dorsal quadrant, opposite to the optic stalk or optic fissure margins without obvious staining in between (see **Fig. 5A'**). These "dorsal" phenotypes were observed in 60% of CF embryos, but never in SF.

In 36hpf SF embryos the RPE marker Bhlhe40 was expressed all around the eye, often 303 contacting the lens (4.7µm away) (Fig. 5B, B', D), which was taken as an indicator of the correct 304 engulfment of the retina by the migrating RPE. The edges of the optic fissure margins 305 306 sometimes lacked staining but overall the expression spanned 326° around the eye (Fig. 5B"). 307 Conversely, in CF, Bhlhe40 expression area was reduced, with a wider ventral gap possibly resulting from the wider optic fissure opening, and a significantly diminished covering of the 308 retina by the RPE (289° around the eye). Bhlhe40-positive cells were also found further away 309 from the lens (15µm), reinforcing the idea of a reduced retina covering by the RPE at this 310 stage; at 48hpf however, the staining span was increased and no longer significantly different 311 from the 36hpf SF (CF 48hpf: 309°; SF 36hpf: 326°) (Fig. 5B, C, B"). These data suggest a slower 312 but still occurring engulfment of the back of the retina by the RPE. 313

The back of the retina was usually well covered with *Bhlhe40*-expressing RPE cells, but with a ventral gap which was well delimited by strongly contrasted, sharp edges (**Fig. S5**). We interpreted this gap as the "exit point" of the optic stalk and used it as a proxy for optic stalk

- width. The *Bhlhe40* gap was wider in CF than in SF (SF: 41µm, CF: 60µm) (Fig. 5E). This was
 consistent with the above-reported increase in *Pax2a* expression domain and points towards
 an increased optic stalk size in cavefish.
- 320 Altogether, these results suggested that both the RPE engulfment/spreading movement and
- 321 the optic fissure margins juxtaposition are slowed -but occur in a delayed manner- in cavefish.
- 322 Below these findings are discussed altogether with regards to earlier observations of eye
- 323 morphogenesis through live imaging.

324 **Discussion**

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326 Astyanax knock-in reporter lines obtained by CRISPR/Cas9

327 Several transgenesis techniques are available in zebrafish and have already been adapted to Astyanax: a classical approach consists in cloning a promoter or regulatory element driving 328 the expression of a reporter or effector gene under its control, flanked with either transposons 329 or meganuclease cutting sites (Kawakami, 2005; Thermes et al., 2002). After injection with the 330 331 appropriate enzyme, transposase or meganuclease, into the 1st cell of the embryo, the construct is randomly inserted one or several times in the genome. This method is technically 332 333 simple and works in Astyanax (Elipot et al., 2014; Hinaux et al., 2015; Stahl et al., 2019). It has however several drawbacks: the correct identification of most if not all the regulatory 334 sequences driving expression is necessary and often difficult; the insertion of the transgene is 335 random and its expression depends on the insertion site; the insertion of the construct can 336 disrupt coding or regulatory sequences. 337

The advent of the flexible CRISPR/Cas9 technique now allows for an "easy" RNA-mediated 338 targeting at a precise genomic location. However, if performing a targeted knock-out by 339 cutting DNA and relying on the imprecise non-homologous end joining (NHEJ) DNA repair 340 mechanism to generate indels and frameshifts has proven quite efficient, more precise repair 341 and insertions are still difficult to obtain, at least in zebrafish. The more precise methods 342 343 require homology-directed repair and therefore involve homology arms flanking a repair construct. Although different ways to increase homology-directed repair efficiency have been 344 tested (Albadri et al., 2017; Morita et al., 2017), it is still a very challenging method as NHEJ is 345 the preferred repair mechanism in fish embryos (Hagmann et al., 1998). 346

The difficulty of identifying *Zic1* regulatory elements with comparative genomics led us to adopt a targeted enhancer-trap strategy, using a NHEJ DNA repair mechanism-based approach to maximize integration efficiency. This option also had the advantage that the insertion site would be similar in CF and SF lines, which is crucial for comparative purposes. Enhancer traps were originally performed by random insertions in a two-step process, the first step being the selection of the expression pattern of interest, the second the identification of the region in 353 which the transgene was inserted. This allowed the generation of transgenic lines with various patterns of expression, reflecting the activity of one or more enhancers and regulatory 354 elements. Here, we "addressed" the enhancer-trap construct to Zic1 downstream genomic 355 region using CRISPR/Cas9 methodology, similarly to what was performed by Kimura and 356 colleagues (Kimura et al., 2014). This method yielded good results as its limited efficiency was 357 compensated by the possibility of using a pattern-based fluorescence screening in FO 358 embryos. Hence, we obtained an excellent ratio of founder fish in the pool of selected F0 359 embryos (more than 50% in cavefish). Finally, for both morphs the different Zic1:hsp70:GFP 360 lines, although recapitulating the Zic1 pattern, showed slight variations in the relative 361 intensities of reporter fluorescence in the telencephalon and in the eyes. The insertion 362 method being based upon non conservative NHEJ mechanism, those variations are likely due 363 to sequence differences from one line to the next (indels and duplications in the genomic DNA 364 or the transgene), which may affect the nearby regulatory sequences and slightly modify 365 transgene expression. However, those variations remain anecdotal compared to the 366 differences observed between lines generated by traditional transgenesis techniques (such as 367 368 Tol2 promoter transgenesis), validating this approach as a tool to follow gene expression in Astyanax morphotypes. 369

CRISPR/Cas9 has been previously reported in surface *Astyanax mexicanus* to generate an *Oca2*null mutant and confirm the role of *Oca2* in the control of pigmentation (Klaassen et al., 2018).
But this is to our knowledge the first report of the CRISPR/Cas9 technology used in this
emergent model species to generate identical reporter lines in the two morphotypes, and in
a targeted genome edition perspective.

375

376 Live imaging

The choice of live imaging microscopy technique was directed by several parameters, from the intensity of the labelling to temporal-spatial resolution trade-off. Light-sheet or SPIM (single plane illumination microscopy) microscopy offered several advantages such as wide dynamic range of the camera, allowing to detect both the strong labelling of the telencephalon and the fainter labelling of the eye. The next step of this study will be to track the cells participating in the formation the optic region, i.e. eye, optic stalk, ORR. Cells in these regions 383 undergo fast movements during early morphogenesis, so that a good time resolution is 384 necessary. They are also densely-packed and therefore require a good spatial resolution. Moreover, live embryos are fragile and laser power must be kept minimal to avoid bleaching 385 of the fluorescence and photo-damage. In contrast to the point acquisition of the confocal 386 microscope, the plane acquisition of the SPIM allowed fast imaging while retaining sufficient 387 spatial resolution. The orthogonal illumination induced minimal photo-damage and embryos 388 developing for more than 20 hours under the microscope recovered well and were alive with 389 a normal head shape at 48-60hpf -even though the tail was usually twisted due to the 390 391 mechanical constraint in the low-melting agarose.

392

393 Recapitulating cavefish eye development

Live imaging performed on the Zic1:hsp70:GFP transgenic lines combined to gene expression 394 studies allowed us to reveal striking differences in eye morphogenesis, morphology and 395 patterning between the two Astyanax morphs. Because the ventral quadrant of the eye was 396 originally described as reduced and because the expression of the ventralizing morphogen Shh 397 398 was known to be enlarged in cavefish (Yamamoto et al., 2004), we expected a global "ventralization" of the eye quadrants at the expense of the dorsal quadrant which we assumed 399 400 would be reduced. However, the data revealed a guite different story and we venture to propose a developmental scenario for cavefish morphogenesis and patterning (Fig. 6). 401

402 (1) First, the shorter size of the cavefish optic vesicles (Alunni et al., 2007; Strickler et al., 2001) seems principally due to the smaller eyefield, since elongation proceeds similarly in CF and SF 403 until 17.5hpf. Increased Hedgehog signalling in CF was shown to decrease the size of its optic 404 vesicles (Yamamoto et al., 2004) and probably the eyefield size, which could account for the 405 final smaller eye size. Moreover, albeit smaller, CF optic vesicles are "correctly" patterned in 406 407 their future naso-temporal axis, as shown by Hernández-Bejarano and colleagues at 10/12 somite stage (~13.5hpf) using FoxG1 and FoxD1 markers (Fig. 6A) (Hernandez-Bejarano et al., 408 409 2015).

(2) Second, after the initial evagination and patterning of a small optic vesicle, morphogenesis
proceeds with the "extended evagination", whereby cells from the neural tube continue
entering the optic vesicle to contribute exclusively to the ventro-nasal part of the eye (Kwan

et al., 2012). If this step proceeds normally in CF, which seems to be the case from live imaging
experiments, it could partially compensate the originally small size of the eyefield/optic
vesicle, but only in the nasal part, while the temporal part would remain fully affected (Fig.
6B). This would well explain the increased angles of nasal *FoxG1* and *Vax2* expression in
comparison to the reduced temporal *FoxD1* and *Vax2* territories we observed. It is also worth
noting that early Hedgehog signalling can increase *Vax2* expression (before early optic vesicle
stage) in *Xenopus* (Wang et al., 2015), which could further explain *Vax2* nasal expansion.

420 (3) Third, the lens forms correctly at the centre of the optic vesicle in both morphs, in a proper 421 place with regard to the initial invagination of the optic cup; it is only at later stages that the 422 lens appears more anterior (i.e., facing the presumptive ventral retina after final eye rotation) in cavefish. This apparent displacement of the lens relative to the retina is not due to a 423 movement of the lens -which remains fixed throughout eye morphogenesis, attached to the 424 overlying ectoderm from which it delaminates around 22hpf in Astyanax (Hinaux et al., 2017)-425 but rather to a persistent elongation of the optic vesicle. This suggests normal interactions 426 between the optic vesicle and the lens to adjust their relative position and initiate optic cup 427 428 invagination. Indeed, in chick, the pre-lens ectoderm is required for normal optic cup 429 invagination while the lens placode itself is dispensable (Hyer et al., 2003). In cavefish such mechanisms could exist and correctly initiate optic cup folding. Finally, the anterior-shifted 430 position of the lens, due to elongation of the cavefish optic vesicle without invagination, 431 explains why the lens is ventrally displaced in the mature eye: the final anterior rotation 432 movement brings it in a ventral position (Fig. 6C). 433

434 (4) Fourth, although the invagination of the CF optic vesicle seems to start normally between 15.5 and 19.5 hpf, it only progresses very little afterwards so that the optic cup remains shallow 435 436 and elongated. This timing is very reminiscent of the 2 described steps of optic cup 437 invagination: basal constriction initiating the primary folding between 18 and 20hpf in zebrafish (18ss to 22ss), followed by the rim movement which brings the presumptive retina 438 from the inner optic vesicle leaflet into the lens-facing epithelium by an active migration 439 around the rims of the optic recess between 20 and 24hpf (Heermann et al., 2015; Nicolas-440 Perez et al., 2016; Sidhaye and Norden, 2017). In Astyanax, 18 ss corresponds to ~16.5hpf 441 (Hinaux et al., 2011), suggesting that the initial basal constriction leading to the onset of optic 442 cup invagination is well conserved in cavefish -this will have to be confirmed. In contrast, the 443

444 prolonged extension of the optic vesicle and the fact that the curvature of the retina remains 445 shallow with the lens bulging out of the eye suggests that the rim movement must be impaired in cavefish. We propose that the continuous flow of cells entering the retina leads to its 446 elongation, in the absence of an efficient rim movement. This movement does seem weaker 447 448 but not absent in CF, as the posterior part of its optic cup still manages to contact the lens, but at later stages. This difference of rim movement might be due to various causes such as 449 defects in the basal membrane or failure to establish proper focal adhesion as seen in the 450 ojoplano medaka mutant (Martinez-Morales et al., 2009; Nicolas-Perez et al., 2016; Sidhaye 451 452 and Norden, 2017). Alternatively, the active migration described by Sidhaye and Norden could be altered by extrinsic signals, as in BMP overexpression experiments where the cell flow 453 toward the lens-facing epithelium is reduced (Heermann et al., 2015). The various morphogen 454 modifications known in cavefish, and the fact that the ventral eye can be restored by delaying 455 the onset of the Fgf signalling in CF to match the SF scenario (Pottin et al., 2011), support this 456 457 possibility.

The "continuous extension of the optic cup with only little rim movement" hypothesis explains cavefish phenotypes. It clarifies the ventral and "floating" position of the lens in the final eye, without much contact with the edges of the retina. It also explains how grooves/folds in the retina (typically between the dorsal and temporal quadrants) are sometimes formed (**Fig. S6**).

(5) Fifth, Bhlhe40 expression reveals that RPE identity is maintained in the CF eye. Yet, the 462 expansion and engulfment movement that this tissue is supposed to achieve to reach the rim 463 of the retina and cover the whole retina is delayed compared to SF. In fact, this spreading and 464 migration of RPE cells is concomitant with the rim movement and may contribute to it as a 465 driving force (Cechmanek and McFarlane, 2017; Moreno-Marmol et al., 2018). This 466 467 observation reinforces the idea that the rim movement is impaired in cavefish. Potentially, the 468 RPE movement may also be involved in optic fissure closure, as suggested by the presence of a coloboma upon impairment of the rim movement by BMP4 overexpression in the optic 469 vesicle (Heermann et al., 2015). Deficiency in this morphogenetic movement might participate 470 in the cavefish coloboma phenotype. Of note, the transplantation of a healthy SF lens into a 471 CF optic cup rescues the eye as a structure, i.e., prevents lens-induced degeneration, but does 472 473 not rescue the coloboma phenotype (Yamamoto and Jeffery, 2000). This is in agreement with 474 our findings showing that the improper closure of the optic fissure is autonomous to the CF
475 retinal tissues and results from defective morphogenetic movements.

476 (6) Sixth, in addition and concomitantly to impaired morphogenetic processes, morphogen signalling modifications previously reported in cavefish might contribute to the axial 477 478 patterning variations observed. On the naso-temporal axis, there seems to be a slight increase in *FoxG1* domain in the early optic vesicle (Hernandez-Bejarano et al., 2015), which may be 479 explained by the earlier onset of Fqf8 expression in the cavefish anterior neural ridge (Pottin 480 et al., 2011), but also by the increase in size of the cavefish olfactory placodes which also 481 482 secrete Fgf24 (Hernandez-Bejarano et al., 2015; Hinaux et al., 2016). These Fgfs, along with Fgf3 promote retinal nasal fate (Hernandez-Bejarano et al., 2015; Picker and Brand, 2005; 483 Picker et al., 2009), and may be responsible for the slight expansion of *FoxG1* expression in 484 cavefish, itself limited or counteracted by increased Shh signalling (Hernandez-Bejarano et al., 485 2015). This might contribute to the importance taken by nasal fates relative to temporal fates 486 in cavefish. It may also account for the Tbx5a angle enlargement towards the temporal 487 quadrant. In fact, Fgf signalling has been shown to promote dorsal identity temporally, as 488 489 demonstrated by the temporal reduction of Tbx5a span and its nasal shift in Fgf8 mutant 490 zebrafish (Ace) (Picker and Brand, 2005). Finally, besides Shh, Fgf and Bmp signalling, it is to note that the mouse Lrp6 insertional mutant displays microphthalmia and coloboma (Zhou et 491 al., 2008), and that zebrafish treated with the Wnt- β cat pathway activator LiCl have small eyes 492 (Shinya et al., 2000), therefore the modulations of canonical Wnt signalling observed in 493 cavefish (Torres-Paz et al., 2018) probably also impact the development of their eyes. 494

495 (7) Pax2a expression domain is increased at the expense of the Pax6-expressing part of the eyefield at neural plate stage in CF (Yamamoto et al., 2004). Here we expand these results at 496 497 later stage and show that Pax2a is no longer confined to the optic stalk and optic fissure 498 margins but invades the ventral retina and even often, the dorsal retina. Contrarily to Vax2, Pax2a/optic stalk identity is expanded in both temporal and ventral directions. This phenotype 499 is probably caused by the increase in Shh signalling in cavefish, as Shh injections in zebrafish 500 501 embryos or Ptch1 (a negative regulator of Hedgehog pathway) loss of function in blowout mutants cause very similar phenotypes with a larger optic stalk and an invasion of the ventral 502 retina by Pax2a expression (Lee et al., 2008). Nevertheless, the observation of cells with an 503 optic stalk identity in the cavefish dorsal retina was surprising, as in zebrafish, Shh signalling 504

505 maintains the optic stalk-retina interface through the regulation of *Vax* genes (Take-uchi et 506 al., 2003), and here we found that *Vax2* was mis-expressed in a much milder manner than 507 *Pax2a*. A possibility would be that these *Pax2a* "ectopic cells" stem from abnormal 508 morphogenetic movements that could not be traced in our study.

509

510 Evolutionary considerations

511 In all eyeless or eye-reduced cave vertebrates examined so far, initial eye development occurs, including the evagination of optic vesicles and further morphogenetic steps (e.g., Durand, 512 1976; Stemmer et al., 2015; Wilkens, 2001). This represents an energetically-costly process for 513 embryos, raising the question of why would all these different species first develop eyes which 514 515 are after all fated to degeneration, and suggesting that initial eye development cannot be circumvented (Rétaux and Casane, 2013). Our results help further refine the step(s) in eye 516 morphogenesis that are subjected to developmental constraint. Indeed, in cavefish, the 517 (small) eyefield is correctly specified and the evagination/elongation steps, which correspond 518 to cell movements leading to the sorting of retinal versus adjacent telencephalic, preoptic and 519 hypothalamic cells of the neural tube, occurs properly. It is only after the segregation between 520 these differently-fated cell populations that the cavefish eye morphogenesis starts going awry, 521 522 with a defective invagination process, that will soon be followed by lens apoptosis and the subsequent progressive degeneration of the entire eye. Therefore, our data support the idea 523 524 that the first steps of eye development constitute an absolute developmental constraint to morphological evolution. To the best of our knowledge, the closest to a counter-example is 525 526 the medaka mutant eyeless, a temperature-sensitive rx3 mutant line in which optic vesicles evagination does not occur. However, the homozygous eyeless fish either die after hatching 527 (Winkler et al., 2000) or, for the 1% which can be raised to adulthood in laboratory, are sterile 528 529 probably due to anatomical hypothalamic or hypophysis defects (Ishikawa et al., 2001), still 530 reinforcing the hypothesis of a strong developmental constraint.

531

532 Conclusion

533 This work proposes conclusions and hypotheses regarding defective cavefish eye 534 morphogenesis, and interprets cavefish eye development in the frame of the current

knowledge on the topic. Further analyses including the tracking, characterization and comparison of cell movements occurring in both morphs will refine the current scenario. For example, quantifying the extent of the rim movement, the curvature of the optic cup and the amount of cells entering the optic vesicle through time would be informative. These data also pave the way for experiments aiming at understanding the defective molecular or signalling mechanisms in cavefish eye morphogenesis, using the *Zic1:hsp70:GFP* knock-in lines.

541 Methods

542

543 Animals

544 Laboratory stocks of A. mexicanus surface fish and cavefish were obtained in 2004 from the Jeffery laboratory at the University of Maryland. The surface fish were originally collected 545 from San Solomon Spring, Texas and the cavefish are from the Pachón cave in Mexico. Surface 546 fish are kept at 26°C and cavefish at 22°C. Natural spawns are induced after a cold shock (22°C 547 548 over weekend) and a return to normal temperature for surface fish; cavefish spawns are induced by raising the temperature to 26°C. Embryos destined for in situ hybridization were 549 550 collected after natural spawning, grown at 24°C and staged according to the developmental staging table (Hinaux et al. 2011) and fixed in 4% paraformaldehyde. After progressive 551 dehydration in methanol, they were stored at -20°C. Embryos destined to transgenesis or live 552 imaging were obtained by in vitro fertilization. Embryos were raised in an incubator until 1 553 month post fertilization for the surface fishes and two month post fertilization for the cavefish. 554 555 They were kept at low density (15/20 per litre maximum) in embryo medium, in 1 litre plastic tanks with a soft bubbling behind the strainer. Larvae were fed from day 5 with paramecium 556 557 and transitioned to artemia nauplii from day 10-15. Artemia were given twice a day except for 558 the weekends (once a day) and carefully removed afterward to avoid polluting the medium. At least two thirds of the medium were changed every day and dead larvae removed. After 559 one month for the surface fish and two months for the cavefish, juveniles were taken to the 560 561 fish facility where they were fed dry pellets (Skretting Gemma wean 0.3) and quickly moved 562 to bigger tanks in order to allow their fast growth.

563 Animals were treated according to French and European regulations of animals in research. 564 SR' authorization for use of animals in research is 91-116, and Paris Centre-Sud Ethic 565 committee authorization numbers are 2012-52 and 2012-56.

566

567 In situ hybridization

Some cDNAs were available from our cDNA library : *Zic1* (FO290256), *Zic2a* (FO320762) and
 Rx3 (FO289986), *FoxD1* (FO380710); others were already cloned in the lab : *Lhx2* (EF175737)

and *Lhx9* (EF175738) (Alunni et al. 2007); obtained from other labs (*Vax1* : Jeffery lab,
University of Maryland,(Yamamoto et al. 2004)); or cloned for the purpose of this work in
pGEMT-Easy (Promega) :

- FoxG1: forward primer CTGACGTTCATGGACCGAGC; reverse primer
- 574 CAGTCTGCTTCCTGTGGATGT.
- Tbx5a: forward primer GCCTTCATTGCGGTCACTTC; reverse primer
 CCCTCGTTCCATTCAGGCAT.
- Vax2: forward primer GGGCAAAACATGCGCGTTA; reverse primer
 CAGTAATCCGGGTCCACTCC.
- Pax2a: forward primer AGCTGCATAACCGAGGCGA; reverse primer
 CTCCATTAGAGCGAGGGATTCCGA
- Bhlhe40: forward primer : GCACTTTCCCTGCGGATTTC; reverse primer :
- 582 TGGAGTCTCGTTTGTCCAGC

cDNAs were amplified by PCR, and digoxygenin-labelled riboprobes were synthesized from 583 PCR templates. Embryos were rehydrated by graded series of EtOH/PBS, then for embryos 584 585 older than 24hpf, proteinase-K permeabilization at 37°C was performed for 36hpf embryos 586 only (10 µg/ml, 15 min) followed by a post-fixation step. Riboprobes were hybridized for 16 hr at 65°C and embryos were incubated with anti-DIG-AP (Roche, dilution 1/4000) overnight 587 at 4°C. Colorimetric detection with BCIP/NBT (Roche) was used. Mounted embryos were 588 imaged on a Nikon Eclipse E800 microscope equipped with a Nikon DXM 1200 camera running 589 under Nikon ACT-1 software. Brightness and contrast were adjusted using FIJI, some of the 590 images used for illustration purpose were created from an image stack, using the extended 591 depth of field function of Photoshop CS5. Area, distance and angle measurements were 592 593 performed using FIJI (Schindelin et al., 2012; Schneider, Rasband and Eliceiri, 2012).

594

595 In vitro fertilization (IVF) and injections

Surface and cavefish were maintained in a room with shifted photoperiod (light: 4pm – 7am,
L:D 15:11) in order to obtain spawns during the working day (*Astyanax* spawn at night (Simon et al., 2019)). Fish activity was monitored after induction and upon visible excitation or when
first eggs were found at the bottom of the tank, fish were fished. Females were processed first

600 to obtain eggs: they were quickly blotted on a moist paper towel and laid on their side in a petri dish. They were gently but firmly maintained there while their flank was gently stroked. 601 If eggs were not released immediately, the female was put back in the tank. Once eggs were 602 collected, a male was quickly processed similarly to females, on the lid of the petri dish to 603 collect sperm. The sperm was then washed on the eggs with 10-20mL of tank water 604 (conductivity ~500µS) and left for a few moments (30s to 2 min approximatively), after which 605 embryo medium was added in the petri dish. Fertilised eggs were quickly laid on a zebrafish 606 injection dish containing agarose grooves. They were injected with a Picospritzer III (Parker 607 Hannifin) pressure injector. 608

609

610 CRISPR injections and Knock-In lines

sgRNA were designed to target the low-conservation regions between elements 1 and 2 and 611 between elements 3 and 4. Two sgRNA were initially designed per region and sgRNA2 was 612 found to efficiently cut the targeted region (Fig. S7). The mix contained Cas9 protein 613 generously provided by TACGENE and sgRNA2 with the following targeting sequence: 614 CCCAATTCACCAGTATACGT (synthesized with AMBION T7 MEGAshortscript[™] T7 transcription 615 kit). Concentrations were kept with a 1:1.5 Cas9 to sgRNA molar ratio and varied between 616 617 0.71μ M (25ng/µL) and 5.67µM (200ng/µL) of sgRNA 2, mostly 2.84 and 1.42µM were used. The donor construct contained a HSP70 promoter used as a minimal promoter, a GFP cDNA 618 and SV40 poly-adenylation signal, flanked by I-Scel meganuclease cutting sites. I-Scel was used 619 to generate sticky ends and was either detached by 7 min at 96°C or injected with the 620 621 construct. Concentrations of the repair construct varied between 3.33 and 10.92nM but were mostly used at 10.71nM. 622

623

624 mRNA injection

Transgenic embryos used for live imaging were injected in the cell or yolk at 1 cell stage with a H2B-mCherry fusion mRNA at a concentration of $50 \text{ ng}/\mu\text{L}$.

627

628 Imaging

Transgenic embryos were obtained by IVF with wild-type eggs and transgenic sperm and were immediately injected with H2B-mCherry mRNA for nuclear labelling. Injected embryos were screened for GFP and mCherry fluorescence under a Leica M165C stereomicroscope around 10-11hpf, when GFP reporter fluorescence first becomes detectable.

Selected embryos were immediately mounted in a phytagel tube (SIGMA, CAS Number: 71010-52-1) molded with Phaseview Teflon mold (1.5mm of diameter) and maintained in position with 0.4% low melting point agarose (Invitrogen UltraPure[™] Low Melting Point Agarose). The tube containing the embryo was placed horizontally into the chamber containing 0.04% Tricaine in embryo medium (Sigma, CAS Number: 886-86-2). The tube was rotated under the microscope so that the embryo would face the objective.

Live imaging was performed approximately from 10.5-11hpf to 24hpf every 2.5min-3min, using a Phaseview Alpha³ light sheet apparatus, coupled with an Olympus BX43 microscope and using either a 20X/NA 0.5 Leica HCX APO objective or a 20X/NA 0.5 Olympus objective. Images were acquired using QtSPIM software (Phaseview), which controlled a Hamamatsu ORCA-Flash4.0 Digital sCMOS camera.

Room temperature was maintained at 24°C by air conditioning and the chamber temperature
was further controlled by a BIOEMERGENCES-made thermostat. Medium level was
maintained by a home-made perfusion system and an overflow to renew the medium.

647

648 Imaging analysis

649 Images obtained were visualized with Arivis Vision4D software and re-oriented to adopt a 650 similar optical section plane, cutting through the middle of the lens and the optic stalk at all 651 time-steps. On one time-step per hour, measurements were performed on the re-oriented image: optic vesicle/optic cup length (at the widest), optic vesicle size increase (calculated by 652 653 subtracting the length at the onset of furrow formation to the length at time t), optic stalk width, distance between the anterior optic cup and the lens, distance between the posterior 654 optic cup and the lens, distance between the optic cup edges, position of the lens relative to 655 anterior optic vesicle (=distance between center of the lens and anterior OV / (distance 656 between center of the lens and anterior OV + distance between center of the lens and 657 posterior OV)) (see schemes on Figures). 658

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856 Figures and Legends

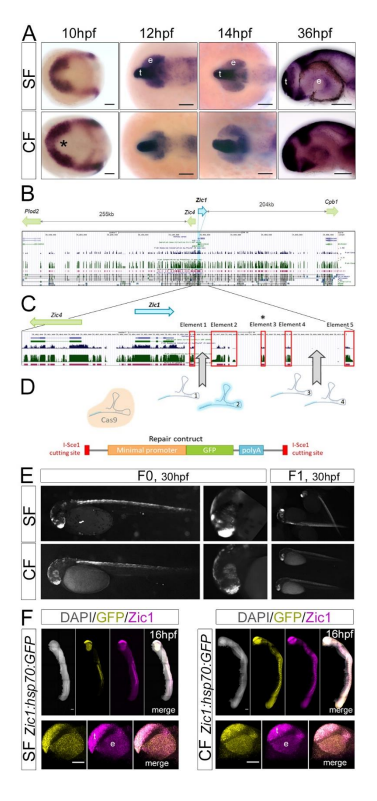
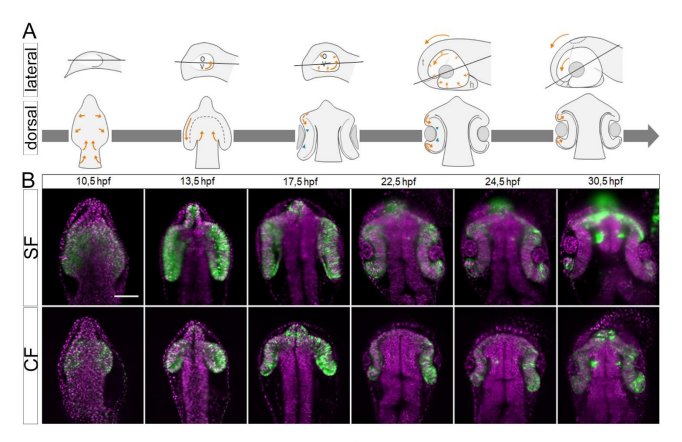


Figure 1





- (A) Zic1 expression at indicated stages in SF and CF. Anterior is to the left. Dorsal views at 10,
- 12 and 14hpf, lateral views at 36hpf. Asterisk: larger indentation in the CF eyefield.
- (B) Zebrafish Zic1 genomic region in UCSC genome browser (2010 assembly). Green blue peaks
- as well as magenta and black elements correspond to high conservation, showing the complexity of the region.
- (C) Close-up on *Zic1*. Red boxes highlight conserved elements; element 3 is not conserved in *Astyanax* (asterisk).
- (C) sgRNA were designed to target the low-conservation regions between elements 1 and 2,
- and 3 and 4. SgRNA2 (pale blue) efficiently generated cuts. It was co-injected together with
- the Cas9 protein and the linear repair construct containing a minimal Hsp70 promoter and the
- 871 GFP.
- (D) Zic1-like GFP fluorescence in mosaic FOs and stable F1s, both for SF and CF.
- 873 (E) Double-fluorescent in situ hybridization at 16hpf for Zic1 (magenta) and GFP (yellow)
- showing that the transgene recapitulates the endogenous Zic1 pattern, both for SF and CF
- 875 lines. The top panels show entire embryos and the bottom panels show close-ups on the head,
- including the *Zic1*-expressing telencephalon (t) and eye (e). Lateral views.
- 877 Scale bars=100μm.



879



880

Figure 2: Live imaging of surface fish and cavefish eye morphogenesis.

(A) Schematic drawings of the principal steps of eye morphogenesis in fish, in lateral (top) and
dorsal views (bottom). Orange arrows indicate cell and tissue movements; green arrowheads
show initiation of basal constriction. The grey line indicates the optical section plane used in
the pictures in B, which follows an optic stalk to lens center axis and accompanies the anterior
rotation illustrated by the arrows. All subsequent measures were realized on these planes.

(B) Still images of time-lapse acquisitions from 10.5hpf to 30.5hpf on SF and SF *Zic1:hsp70:GFP* lines (green, GFP; magenta, nuclear mCherry). Representative steps of eye morphogenesis

889 illustrating CF/SF differences are shown. Dorsal views, anterior to the top.

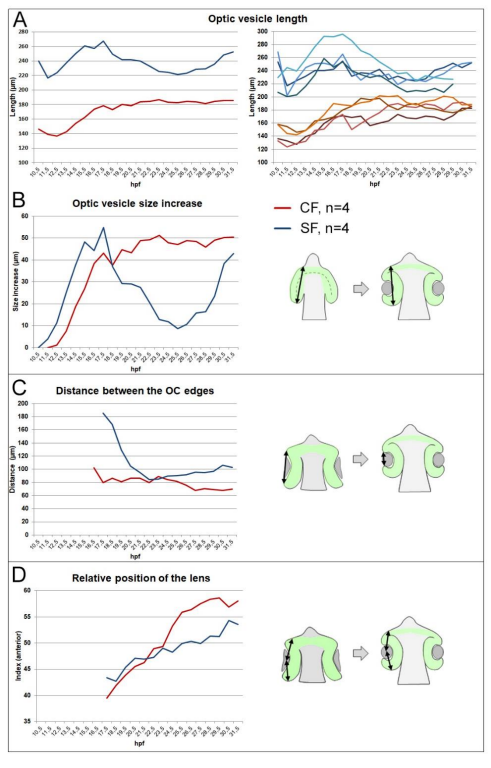
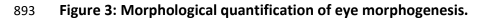
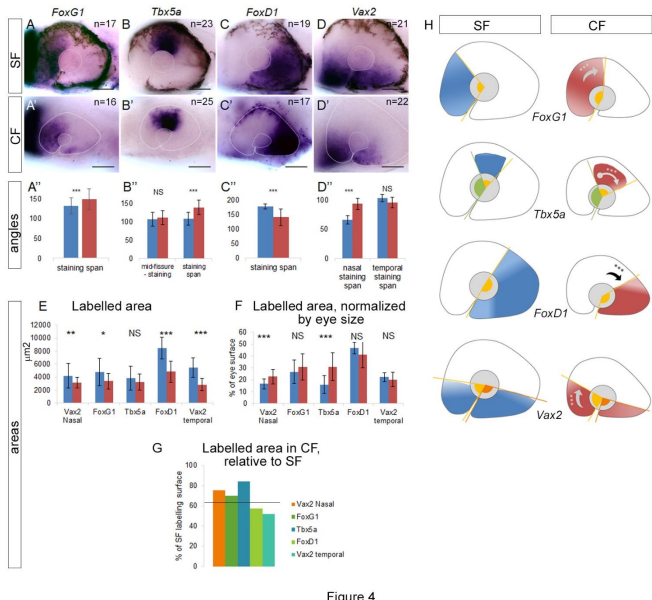


Figure 3

891



- 894 Measures are illustrated on the diagrams on the right in each box.
- (A) Optic vesicle length. The left graph shows the mean of n=4 eyes in each morph (blue, SF;
- red, CF); the right graph displays the trajectories of individual eyes to show the reproducibility
- 897 of the results.
- 898 (B) Optic vesicle size increase.
- 899 (C) Distance between the two optic cup edges.
- 900 (D) Position of the lens relative to anterior optic vesicle, showing that the lens is progressively
- shifted anteriorly between 25hpf and 30hpf.



902

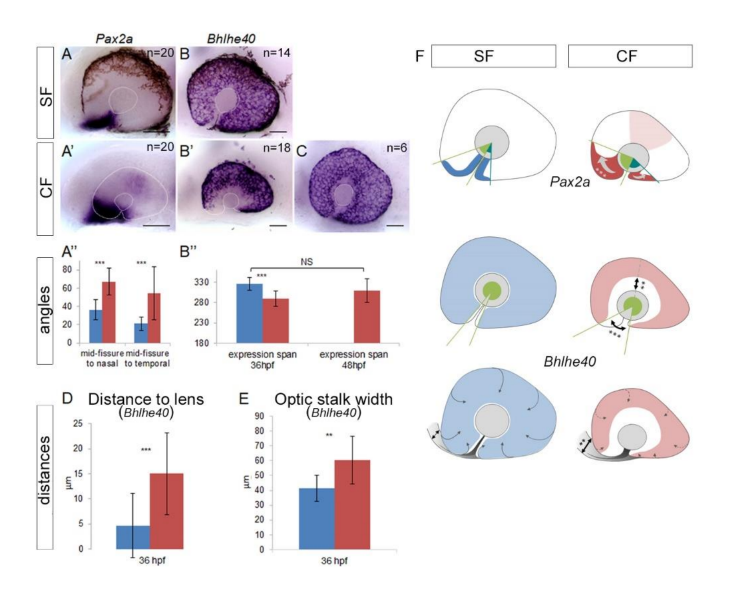


Figure 4: regionalization of the eye. 903

(A-D') In situ hybridization for the indicated markers on 36hpf SF and CF embryos. Sample sizes 904

- 905 are indicated.
- (A"-D") Quantification of angles of expression and position with regards to the choroid fissure. 906
- All the angles measured are represented in H. SF, blue; CF, red. 907
- (E) Absolute area of expression, in μm^2 . 908
- (F) Area of expression normalized by eye size. 909
- (G) Area of expression in CF, expressed as percentage of the expression area in SF. Horizontal 910
- line shows that the CF eye area is 64% of the SF eye. 911
- Statistical test: Mann-Whitney; * p<0.05; ** p<0.01; *** p<0.001. 912

- 913 (H) Summary diagram of the expression patterns. The angles measured are represented.
- 914 Expression patterns are represented in blue for SF and red for CF. The angles measured are
- 915 between the middle of the choroid fissure and the borders of the expression domains.



917

Figure 5

918 **Figure 5: tissue identity.**

- 919 (A-B') In *situ* hybridization for the indicated markers on 36hpf SF and CF embryos. (C) *Bhlhe40*
- 920 expression in 48hpf CF. Sample sizes are indicated.
- 921 (A") Angle between the optic fissure and the limit of Pax2a domain (see F for angles
- 922 measured).
- 923 (B") Angles of *Bhlhe40* expression at 36 and 48hpf.
- 924 (D) Distance between *Bhlhe40*-expressing cells and the lens at 36hpf.
- 925 (E) Width of the *Bhlhe40* expression gap at the back of the eye. SF, blue; CF, red.
- 926 Statistical test: Mann-Whitney; * p<0.05; ** p<0.01; *** p<0.001.
- 927 (F) Summary diagram of expression patterns. Angles measured are represented.

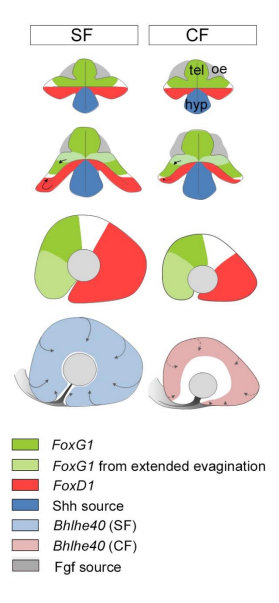


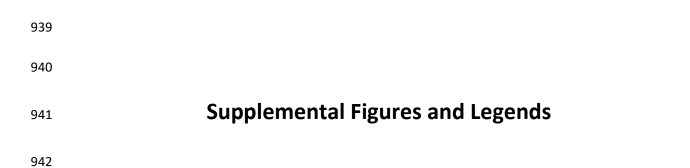
Figure 6

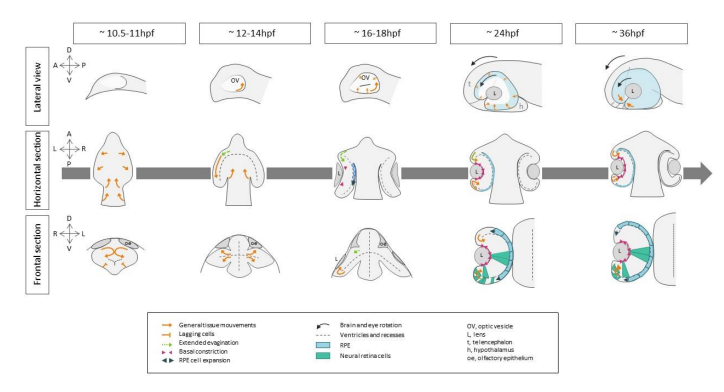
928 929

930 Figure 6: summary and hypothetical scenario for cavefish eye development defects.

Combining data from live imaging and gene expression patterns, we propose that the small size of the optic vesicles, correctly patterned, explains the decrease in temporal quadrant size. Unaltered extended evagination movements bring new cells into the prospective nasal retina, partially rescuing its size (pale green). Finally, the expansion of the RPE to engulf the retina which is concomitant with the rim movement is delayed, leaving the optic stalk wide and optic fissure margins away from each other.

937 oe, olfactory epithelium; tel, telencephalon; hyp, hypothalamus.

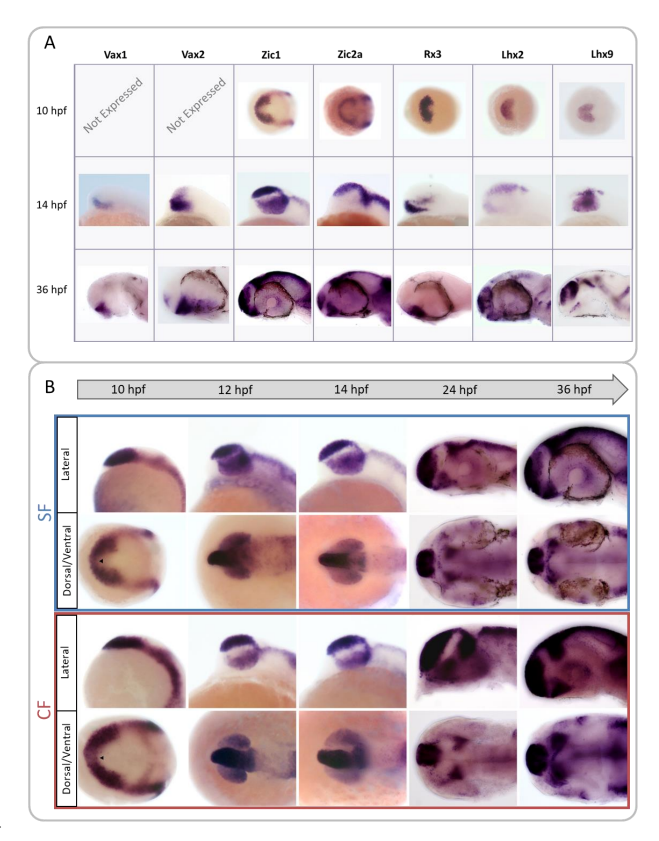




943

944 Supplemental Figure 1: eye morphogenesis in fish.

- 945 Schemes depicting the principal steps of eye morphogenesis in fish models, summarized
- 946 from the available literature cited in Introduction.
- 947 Stages and orientations are indicated.
- 948 Orange arrows show general cell and tissue movements.
- 949 Black arrows show the anterior-wise rotation of the eye and brain.
- 950 Green arrows show the contribution of extended evagination.
- 951 Pink arrowhead show cellular basal constriction.
- 952 The blue color depicts the RPE cells, while the green color depicts retina neuroepithelium cells
- 953 changing shape.



954

955 Supplemental Figure 2: choosing a candidate gene for transgenesis.

956 (A) Mini-screen of candidate genes by *in situ* hybridization at different stages of interest on

957 surface fish embryos (anterior is to the left). Dorsal views at 10hpf; lateral views at 14hpf and

36hpf. The eyes were dissected out for *Vax1* and *Lhx9* (as no eye expression was detected foreither of them) to allow better visibility of the inner tissue.

960 (B) Detailled analysis of *Zic1* expression pattern at 5 different stages in surface (SF) and 961 cavefish (CF). Anterior is to the left, at 10, 12 and 14hpf, bottom pictures are taken in dorsal 962 view; at 24 and 36hpf, bottom picture are taken in ventral views. Arrowheads indicate an 963 indentation in the eyefield.

964 Description of expression patterns:

Vax1 expression was detectable from 12hpf in the presumptive ORR (between the optic
vesicles) and additionally in the dorsal hypothalamus (according to brain axis (Puelles &
Rubenstein, 2015), closest to the ORR) and quite faintly in the ventral telencephalon.

968 Vax2 expression was very similar to Vax1 both in terms of onset of expression and pattern, 969 with the addition of the ventral quadrant of the eye. Although Vax2 had a very interesting 970 ventral pattern, we discarded it as a candidate for transgenesis for its expression onset was 971 very late. Moreover, in Vax2 enhancer trap zebrafish line (Kawakami Laboratory), the GFP 972 fluorescence is only visible at 18hpf (personal observation, data not shown).

973 *Rx3* expression showed a typical eyefield expression pattern at 10hpf but progressively faded 974 away during optic vesicle stages and was finally not expressed anymore at 24hpf. Conversely, 975 an anterior and ventral expression in the presumptive hypothalamus was detectable from 976 12hpf and remained throughout the stages examined. At 36hpf, it was clear that only the 977 dorsal half of the hypothalamus, closest to the ORR, was labelled. Due to the rapid fading of 978 its optic vesicle expression, we did not consider *Rx3* as a valid candidate.

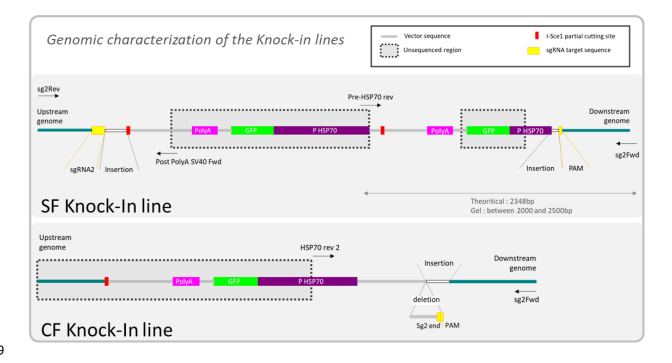
Lhx2 and *Lhx9* were both already known to be expressed in the eyefield at neural plate stages
in *Astyanax* (Pottin et al., 2011). *Lhx2* expression showed very dim expression, if any, in the
optic vesicles at 12 and 14hpf but was expressed both in the prospective telencephalon and
more faintly in the prospective hypothalamus. Later on at 36hpf, *Lhx2* was expressed strongly
in the telencephalon and the olfactory epithelia; lighter expression was also visible in the ORR,
hypothalamus and sometimes eyes. Additional expression in the pineal, optic tectum and in
the hindbrain was also present.

Lhx9 staining was strong in the optic vesicles at 12hpf (during evagination) and slightly lighter
at 14hpf. Moreover dorsal and ventral lateral labelling at the border of the neural keel and the

988 optic vesicles appeared, possibly prefiguring respectively the strong telencephalic staining 989 visible at 24 and 36hpf and the hypothalamic cluster at the limit of the ORR already described 990 in a previous publication (Alié et al., 2018). At these late stages, we could not detect *Lhx9* 991 expression in the eye anymore. Salt and pepper staining was visible in the olfactory epithelia; 992 a band of expression outlining the optic tectum and lateral discrete marks in the hindbrain 993 were present. We did not choose *Lhx2* or *Lhx9* because of the rapid decay of their eye 994 expression.

995 At 10hpf, Zic2a was expressed at the border of the neural plate and almost entirely 996 surrounding the eyefield except for a medial posterior gap. Faint staining in the bilateral eyefield could also be seen on some embryos. At 12 and 14hpf, there was a strong Zic2a 997 expression in the telencephalon and a faint staining in the eye or distal part of the eye could 998 often be seen. Strong staining was generally visible throughout the dorsal-most brain. At 999 1000 24hpf, Zic2a expression remained strong in the telencephalon and was also now strongly visible at the border of the eye, in the ORR or optic stalk but without reaching the midline. 1001 Faint staining in the eye remained. At 36hpf, the expression pattern was similar, with the 1002 1003 ORR/optic stalk staining reaching much closer to the midline. The eye expression was now 1004 more focused around the lens, probably in the CMZ. Roof plate staining persisted throughout development. Because Zic2a was never strongly expressed in the eye, we did not favour it as 1005 a candidate for transgenesis. 1006

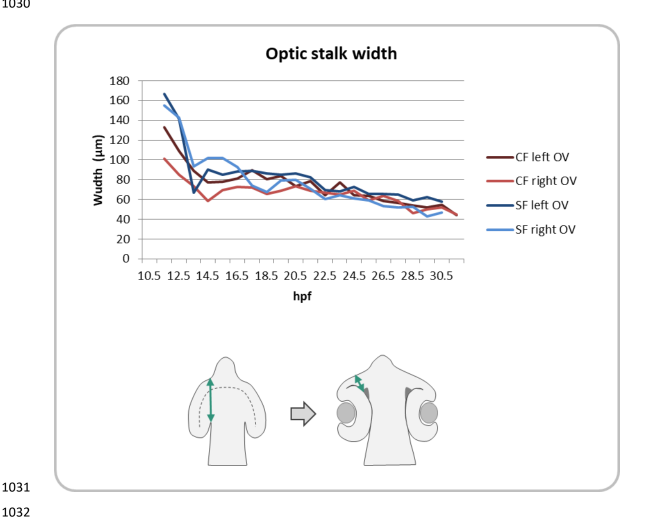
1007 Zic1 was strongly expressed at 10hpf in the neural plate border and in the anterior neural plate, at the level of the eyefield. At 12 and 14hpf, Zic1 expression was consistently found in 1008 1009 the optic vesicles and between them (prospective ORR and optic stalk). A strong staining was also present throughout the telencephalon. More posteriorly, the roof plate of the midbrain 1010 1011 and hindbrain was stained. The somites were also labelled. The pattern was very similar at 1012 24hpf and 36hpf with a strong telencephalic expression and a milder ORR expression (mainly laterally and posterior to the optic recess)/optic stalk and eye staining (widely around the 1013 lens). Roof plate and somites expression remained. Even though its pattern of expression was 1014 1015 complex and encompassed a region wider than the optic region of interest, Zic1 was chosen for transgenesis due to its early and persistent expression throughout the eye and the 1016 1017 ORR/optic stalk regions.



1019 1020

1021 Supplemental Figure 3: genomic characterization of the Knock-in lines.

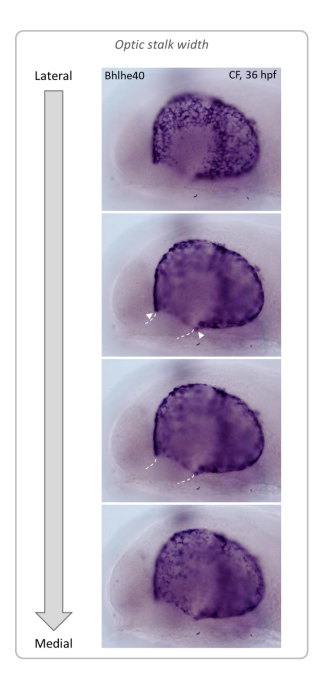
Knock-In insertions, based on partial sequencing. Dotted boxes indicate un-sequenced
regions, leaving uncertainties. For example, in the surface fish line, there is at least a partial
insertion of the repair construct, containing a truncated Hsp70 promoter and at least another
insert in the same direction (but potentially several). Of note, the surrounding genomic region
is very rich in T and A (GC content around 35%) with many repeats, making PCRs sometimes
challenging.
The data show that for both lines the transgenes are inserted at the correct targeted site.



1032

Supplemental Figure 4: optic stalk width. 1033

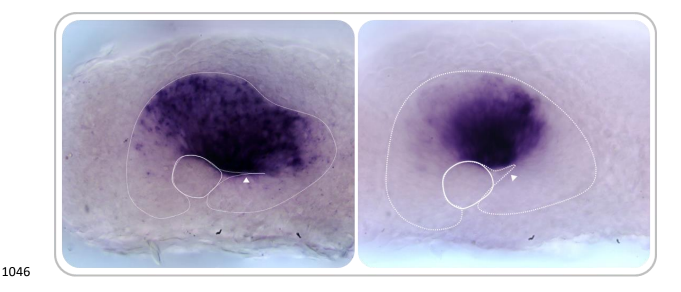
The size of the optic stalk (in a wide meaning: the connection between the optic vesicle and 1034 the neural tube) is smaller in cavefish during early development due to the smaller size of the 1035 optic vesicles but rapidly becomes indistinguishable from the optic stalk of the surface fish. 1036





Supplemental Figure 5: illustration of the gap of *Bhlhe40* staining, which we interpret as the optic stalk width.

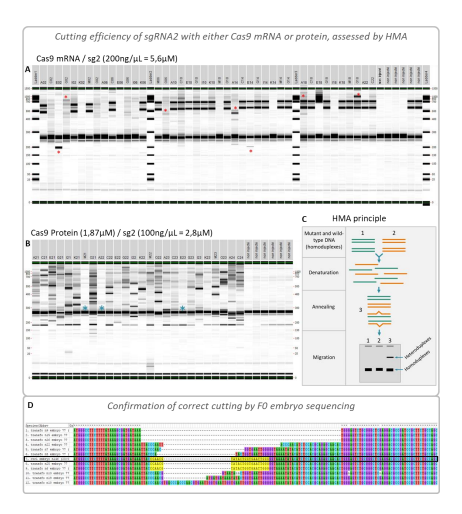
1042 Different focuses of the same embryo, from lateral to medial on the felt eye. Arrowheads 1043 indicate the limits of the staining gap we measured. Dotted lines show the exit trajectory of 1044 the optic stalk.



1047

Supplemental Figure 6: retinal folds in cavefish. 1048

Illustration of the retina folds sometimes observed in cavefish: an "extreme case" on the left 1049 and a more moderate case on the right. 36hpf cavefish embryos, mounted laterally, anterior 1050 is to the left. 1051



1053 1054

1055 Supplemental Figure 7: cutting efficiency of sgRNA 2

(A) Assessment of sgRNA 2 cutting efficiency when injected with Cas9 mRNA by heteroduplex mobility assay (HMA, explained in (C)). Each column is an individual F0 embryo. Embryos with strong additional bands are labelled with a red asterisk; additional light bands can be seen in several individuals, indicating cuts and imprecise repairs. Note that the 2 heavy bands seen on many embryos are also present in some of the un-injected controls (the 6 columns on the right) indicating a polymorphism in this region in the wild-type fish (not on the sgRNA target sequence).

(B) Assessment of sgRNA 2 cutting efficiency when injected with Cas9 protein, note the strong presence of additional band compared to the 6 control embryos on the right. Embryos without any visible cuts are labelled with a blue asterisk. Additional bands are seen much more frequently and are much more important than with the Cas9 mRNA injection, probably indicating more frequent but also more precocious cut and repair events in the embryo, so that many cells share the same sequence. (C) Principle of the heteroduplex mobility assay: in an electrophoresis, heteroduplexes are
slowed down compared to homoduplexes so that they form additional bands that can be seen
even if the polymorphism is only a single substitution. In short, the DNA fragments are
denatured and renatured to form heteroduplexes. An electrophoresis is then performed (here
with a LabChip, PerkinElmer) to detect the presence of polymorphism.

1074 (D) Different cutting and repair events in a single injected embryo. A PCR was performed on one injected embryo (100ng/ μ L sgRNA2, Cas9mRNA) around the sgRNA2 target site and the 1075 1076 product was cloned into pGEM-T Esay (Promega) and transformed into One shot TOP10 competent bacteria (Thermo Fischer). Plasmidic preparations from individual colonies were 1077 then sequenced. Various sequences were obtained, evidencing different cut and repair events 1078 in one single embryo. sgRNA2 target sequence is highlighted in yellow whenever intact. This 1079 1080 FO fish harbours both insertions and deletions around the cutting site of sgRNA2. A noninjected control fish sequence is included, outlined in black. 1081

1082