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Retinoic acid temporally orchestrates colonization of the gut by vagal neural crest cells

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3	neural crest cells
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13	Running head: Retinoic acid temporally regulates ENS formation
14	Key words:
15	Retinoic Acid, neural crest, meis3, zebrafish, enteric nervous system
16	
17 18	Summary
19	The enteric nervous system arises from neural crest cells that migrate as chains into and
20	along the primitive gut, subsequently differentiating into enteric neurons and glia. Little is
21	known about the mechanisms governing neural crest migration en route to and along the
22	gut in vivo. Here, we report that Retinoic Acid (RA) temporally controls zebrafish enteric
23	neural crest cell chain migration. In vivo imaging reveals that RA loss severely

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24 compromises the integrity and migration of the chain of neural crest cells during the 25 window of time window when they are moving along the foregut. After loss of RA, enteric progenitors accumulate in the foregut and differentiate into enteric neurons, but 26 27 subsequently undergo apoptosis resulting in a striking neuronal deficit. Moreover, 28 ectopic expression of the transcription factor meis3 and/or the receptor ret, partially rescues enteric neuron colonization after RA attenuation. Collectively, our findings 29 suggest that retinoic acid plays a critical temporal role in promoting enteric neural crest 30 chain migration and neuronal survival upstream of Meis3 and RET in vivo. 31 anusci 32 33 34 35 Impact statement: A novel stage-specific role for Retinoic Acid in enteric neural 36 crest migration and neuronal survival along the developing gut in vivo 37 ccef 38 39 40 41 Introduction 42 43 The enteric nervous system (ENS) is a network of thousands of interconnected ganglia, located within the myenteric and submucosal plexuses of the gut wall, that 44 regulates peristalsis, secretions and water balance (Furness, 2006; Bergeron et al., 45 46 2013). As the largest division of the peripheral nervous system, the mature ENS contains hundreds of millions of neurons, surpassing the number found in the spinal cord 47

48 (Furness, 2006). In particular, the ENS is the only major population of neurons outside of the central nervous system with circuits that perform autonomous reflex activity. In 49 humans, defects in ENS formation lead to pediatric disorders such as *Hirschsprung* 50 51 Disease (HSCR), also known as gut aganglionosis, which is characterized by a paucity of 52 ganglia along variable lengths of the gut. Colonic aganglionosis is the most common 53 form of HSCR, occurring every 1 in 5000 births (Obermayr et al., 2012; Bergeron et al., 54 2013). The current treatment is surgical resection of the aganglionic intestinal segment (Bergeron et al., 2013); however eventual outcomes of patients varies greatly and most 55 56 exhibit functional enteric defects throughout life-highlighting the need for alternative treatments and understanding the ontogeny of both normal and abnormal ENS 57 58 development.

During development, the ENS is largely derived from "vagal" neural crest cells 59 that arise dorsally in the post-otic neural tube at early stages of central nervous system 60 formation (Le Douarin et al., 1973; Epstein et al., 1994; Kuo and Erickson, 2011). Neural 61 62 crest cells are a group of multipotent stem cells that undergo an epithelial to mesenchymal transition (EMT) and embark on migratory routes in order to reach their 63 64 final and often distant destinations throughout the embryonic body (Saint-Jeannet, 2006; Green et al., 2015). During development, neural crest cells give rise to various types of 65 specialized tissues, including craniofacial cartilage, peripheral neurons and glia, pigment 66 67 cells and dental tissues (Green et al., 2015). During their initial journey, vagal neural crest emigrate from the neural tube at axial levels adjacent to somites 1-7, migrate 68 69 ventrally and, after entering the foregut mesenchyme, migrate in chains along its 70 rostrocaudal extent until they reach the hindgut (Le Douarin and Teillet, 1973; Peters-Van Der Sanden et al., 1993; Burns et al., 2000; Tucker et al., 1986; Anderson et al., 2006). 71

72 The process of gut invasion by vagal neural crest is largely conserved across jawed vertebrates, though the ENS is simpler in zebrafish than in amniotes (Shepherd et 73 al., 2004; Heanue et al., 2016; Ganz et al., 2016). By 32 hours post fertilization (hpf), 74 75 zebrafish vagal neural crest cells commence migration medially toward the foregut as two chains and migrate caudally along the left and right sides of the gut tube to fully colonize 76 the hindgut by 72 hpf (Kelsh and Eisen, 2000; Dutton et al., 2001; Shepherd et al., 2001; 77 Elworthy et al., 2005; Olden et al., 2008; Uribe and Bronner, 2015). Within the gut 78 environment, neural crest cells are referred to as 'enteric neural crest'. To ensure proper 79 80 numbers of enteric progenitors, enteric neural crest proliferate extensively and undergo terminal differentiation to give rise to glia and numerous classes of enteric neurons-such 81 as motor, sensory and interneurons (Furness, 2006; Sasselli et al., 2012). Ultimately, 82 83 enteric neurons terminally differentiate into subtypes, such as serotonergic (5HT), catecholaminergic and Nitric Oxide (NO)-containing neurons (Sasselli et al., 2012; 84 Uesaka et al., 2016). 85

Mutations have been identified in several genes that contribute to HSCR and are 86 required for ENS development (Zimmer and Puri, 2015; Bergeron et al., 2013). These 87 88 include the receptor tyrosine kinase *Ret*, expressed on vagal neural crest cells, and its ligand, Glial Cell-line Derived Neurotrophic Factor (GDNF). Ret mutations cause total 89 intestinal aganglionosis and are known to be the most common cause of HSCR in 90 91 humans; the conservation of this function has been demonstrated in numerous model organisms, such as mouse and zebrafish (Natarajan et al., 2002; Pachnis et al., 1998; 92 Jain et al., 2004; Heanue et al., 2008). Similar to Ref^{-/-}, GDNF^{/-} mice lack almost all 93 94 enteric ganglia (Taraviras and Pachnis, 1999; Sanchez et al., 1996; Moore et al., 1996; Pichel et al., 1996). Additionally, loss of genes that encode conserved transcription 95

96 factors *Sip1*, *Sox10*, *Foxd3*, *Tfap2α* and *Pax3* cause reduction or loss of vagal neural
97 crest cells, along with defects in the ENS and other parts of the peripheral nervous
98 system (Southard-Smith et al., 1998; Kapur, 1999; Lang et al., 2000; Wakamatsu et al.,
99 2001; Knight et al., 2003; Montero-Balaguer et al., 2006; Carney et al., 2006).

100 While these studies identify some transcription factors and receptors/ligands 101 critical for ENS development, events that control the timing of enteric neural crest 102 migration and differentiation along the gut are less well understood. Outstanding 103 questions remain regarding the temporal signals and their downstream mediators that 104 regulate enteric neural crest migration en route to and along the gut. These events are 105 difficult to investigate in amniotes since the gut is semi-opague and problematic to visualize. While studies of gut explants have made it possible to analyze the speed, 106 107 directionality and frontal expansion of the enteric neural crest during both their individual 108 and collective cell migration (Druckenbrod and Epstein, 2005; Young et al., 2014), far 109 less is known about the molecular or temporal mechanisms underlying this migration. 110 One signal that has been suggested to influence enteric neural crest development is Retinoic Acid (RA). For example, in mouse embryos, knockout of 111 retinaldehyde dehydrogenase Aldh1a2 (formerly Raldh2), the enzyme responsible for 112 113 generation of RA, results in complete failure of ENS formation due to the catastrophic 114 loss of vagal neural crest cells (Niederreither et al., 2003). Consistent with this, studies in 115 cell and explant cultures have shown that RA treatment affects enteric neural crest migration (Simkin et al., 2013; Fu et al., 2010), proliferation and/or neuronal differentiation 116 117 (Sato and Heuckeroth, 2008). However, it remains unclear when RA is affecting enteric 118 neural crest migration or what downstream transcriptional effectors are involved. These

119 events are particularly difficult to study in amniotes due to the inaccessibility of intact gut 120 tissue to live imaging.

121 To address these questions in vivo, we have turned to zebrafish embryos due to 122 their ease of manipulation and optical clarity that enables visualization of the process of 123 enteric neural crest migration in real time. Utilizing pharmacological and genetic modulation of the RA pathway, combined with live imaging and epistasis experiments, we 124 125 examine the temporal role of the RA pathway during early ENS development. The results reveal a novel mechanism whereby RA functions to temporally influence migration 126 of enteric neural crest and neuronal survival upstream of the transcription factor Meis3 127 namu and the RET receptor in vivo. 128

129

130 Results

131

RA pathway components are expressed in the gut during enteric neural crest 132

133 migration

As a first step in investigating the role of RA signaling in zebrafish enteric 134 135 nervous system development *in vivo*, we examined the distribution of RA pathway 136 components involved in RA synthesis and turnover along the gut during enteric neural crest migration. To this end, we performed in situ hybridization to examine the spatial 137 138 distribution of the following transcripts at 48 hpf: aldehyde dehydrogenase 1 family, 139 member A2 (aldh1a2), retinol binding protein 1a, cellular (rbp5) and cellular retinoic acid 140 *binding protein 2, a (crabp2a).* While retinol-binding proteins are involved in the transfer 141 of retinol, the RA precursor also known as Vitamin A, aldehyde dehydrogenase enzymes 142 are responsible for the synthesis of RA from retinol and Crabp proteins balance the 143 intracellular levels of RA to mediate feedback and degradation of RA (Schilling et al.,

144 2012; Barber et al., 2014). In particular, we focused on their expression along the

145 foregut, midgut and hindgut, as depicted schematically in **Figure 1A**.

146 *aldh1a2* was detected at all levels of the gut at 48 hpf (**Fig. 1B**; arrows).

147 Similarly, *rbp5* was observed along the entire gut at 48 hpf (**Fig. 1C**), while *crabp2a* was

148 primarily localized to the foregut (Fig. 1D). In transverse sections through the foregut, we

149 confirmed localization of Aldh1a2 protein to the gut mesenchymal layer through which

150 neural crest cells migrate, as well as in neural crest cells, using an antibody specific to

151 zebrafish Aldh1a2. Signal was observed surrounding the gut endoderm, as marked by

152 sox17:GFP (Fig. 1E-E") and in neural crest visualized by sox10:GFP at 48 hpf (Fig. 1F-

153 F", arrows). Altogether, these results show that RA pathway components are expressed

154 in the gut during enteric neural crest gut migration, consistent with possible functional

155 roles therein.

156

157 Temporal addition of RA during colonization is sufficient to enhance enteric

158 neuron numbers in vivo

159 Tissue culture studies have shown that RA can enhance enteric neural crest migration (Sato and Heuckeroth, 2008; Fu et al., 2010; Simkin et al., 2013), suggesting a 160 role in enteric nervous system formation. To test the time during which RA plays a role in 161 enteric neural crest migration, we tested the effects of exogenous application of RA 162 163 during the migratory phase en route to and along the zebrafish gut. To this end, 164 zebrafish embryos were incubated in 1 μ M RA or with DMSO alone, as a control, from 24-48 hpf or from 26-52 hpf. Migration of neural crest cells was visualized via expression 165 166 of the pan neural crest marker *crestin* by *in situ* hybridization (**Fig. 2A-D**). When compared with control embryos, we observed an expansion of *crestin*⁺ neural crest cells 167

along the vagal and anterior trunk level of the embryo in RA-treated embryos (Fig. 2A,B; 168 169 arrows). Histological examination showed an expanded domain of *crestin*⁺ cells localized 170 near the vicinity of the foregut, when compared with control embryos (Fig. 2C.D: arrows). 171 As a secondary means of analyzing the distribution of neural crest cells, examination of the Tg(sox10:GFP) line revealed GFP⁺ cells surrounding the foregut in 172 transverse sections at 52 hpf in control embryos (Fig. 2E), and an expansion of GFP⁺ 173 neural crest cells near the foregut in RA-treated embryos (Fig. 2F; arrows). To test 174 whether this early enhanced localization of neural crest cells later resulted in altered 175 176 distribution of enteric neurons at larval stages, we performed whole mount immunochemistry with antibodies to the pan-neuronal marker Hu, as well as 5HT to 177 identify serotonergic neurons in control and RA-treated larvae guts at 96 hpf. RA-treated 178 larvae exhibited an increase in the total numbers of Hu⁺ enteric neurons along the gut, 179 with an average of 106, compared with control larvae, which had an average of 87 Hu⁺ 180 neurons (Fig. 2G,H-H',I-I'; p=.011). Similarly, RA-treated larvae exhibited increased 181 182 numbers of 5HT⁺ neurons, with an average of 23, compared with control larvae, which had an average of 19 5HT⁺ neurons along the length of the gut (Fig. 2G,H-H",I-I"; 183 p=.001). However, the percentages of 5HT neurons were similar between control (21.8% 184 185 5HT⁺ neurons) and RA-treated (21.7% 5HT⁺ neurons) larvae indicating that RA addition 186 increased the overall numbers of neurons. Collectively, these experiments suggest that 187 addition of RA at the time when enteric neural crest cells invade and initiate migration 188 along the rostrocaudal extent of the gut is sufficient to enhance the numbers of enteric precursors that subsequently leads to an increase in the numbers of enteric neurons in 189 190 vivo.

Loss of RA during gut invasion prevents caudal colonization of the gut by enteric neural crest

194 In order to test the role of RA during early ENS development, we examined the 195 effects of inhibiting RA pathway activity on the progression of enteric neural crest cell 196 migration along the gut. Importantly, we inhibited RA signaling after 24 hpf to avoid early 197 effects of RA on embryonic patterning and focus on its temporal role during ENS 198 development. To this end, zebrafish embryos were treated with 10 µM DEAB (N,N-199 diethylaminobenzaldehyde), an aldehyde dehydrogenase inhibitor (Morgan et al., 2015), or DMSO for control, from 24-52 or 24-72 hpf. Such DEAB treatment did not result in 200 global defects in neural crest localization at cranial or trunk levels, as assayed by 201 202 examining the neural crest marker crestin in DEAB treated and control embryos at 52 hpf 203 (Fig. 3A,B; arrows), nor did it alter the differentiation of neural crest-derived 204 melanophores (Fig. 3L).

205 At 72 hpf, the localization of *crestin*⁺ neural crest cells was evident within the 206 hindgut in control larvae, while neural crest cells in DEAB treated larvae were restricted to the foregut (Fig. 3C,D; arrowhead), indicating delayed migration. To investigate the 207 208 effects of RA inhibition on enteric neural crest localization along the gut in live larvae, we 209 utilized the Tg(-8.3phox2bb:Kaede) line, which expresses the fluorescent protein Kaede 210 under control of a 8.3 kilobase (kb) phox2bb enhancer (Harrison et al., 2014), expressed 211 in enteric neuronal precursors and differentiated neurons. The results showed that DEAB treated larvae exhibited delayed enteric neural crest gut colonization. Whereas control 212 213 larvae contained enteric neural crest at the level of the hindgut at 72 hpf, enteric neural 214 crest cells were restricted to the foregut of DEAB treated larvae (Fig. 3E,F).

To assess possible effects of RA depletion of mesodermal tissue integrity along the gut, we examined larvae from the *TgBAC(hand2:GFP)* line (Yin et al., 2010) which labels gut mesenchyme along the gut tube. Using confocal z-stack imaging, we found no difference between control and DEAB-treatment in the anterior-posterior presence of gut mesenchyme in live larvae (**Fig. 3G,H**), demonstrating that gut mesodermal tissue is present along the gut tube following temporal reduction of RA.

221 Next, we investigated whether the delay in neural crest colonization of the gut after RA loss might be due to a reduction in the number of neural crest cells. To this end, 222 we quantified numbers of neural crest cells along and near the foregut in transverse 223 cryosections. Interestingly, DEAB treated sox10:GFP⁺ larvae did not exhibit a decrease 224 in the number of neural crest cells with direct gut contact at 52 hpf, compared with 225 controls (Fig. 3I-K; white arrows). However the number of *sox10*:GFP⁺ neural crest in 226 the ventral mesenchyme surrounding the gut was increased (Fig. 3I-K; yellow arrows). 227 228 These results suggest that neural crest cells accumulate in the vicinity of the foregut, but 229 then fail to migrate along the gut following temporal loss of RA.

230 To confirm the effects of loss of RA on enteric development using an additional approach, we utilized the transgenic line Tg(hsp70:dnRAR-GFP) (Kikuchi et al., 2011), 231 232 which expresses a dominant negative zebrafish Retinoic acid receptor, Raraa-GFP, under control of the heat shock promoter *hsp70*. Tg(*hsp70*:dnRAR-GFP)^{-/+} embryos 233 234 were subjected to a single one-hour heat shock at 24 hpf and allowed to to develop to 48-235 52 hpf. Heat shock control embryos (GFP) and heat shock-positive embryos (GFP) were analyzed by in situ hybridization against crestin (Fig. S1). Although crestin 236 237 localization was not altered in the cranial or trunk neural crest regions (Fig. S1A,B), we observed a significant delay in enteric neural crest progression along the foregut in the 238

dnRAR embryos. When control embryos were examined in dorsal view, *crestin*⁺ neural
crest cells were present as two chains emanating from the post-otic vagal regions and
migrating along the foregut, just past the level of the fin buds (Fig. S1A'; arrows). In
contrast, the foregut was just beginning to be colonized by neural crest in dnRAR-GFP⁺
embryos, with neural crest chains evident rostral to the level of the fin buds (Fig. S1B';
arrows).

To examine enteric neural crest cell localization along the gut, Tg(hsp70:dnRAR-245 GFP) fish were crossed with Tg(sox10:mRFP) fish and mRFP⁺ progeny were heat 246 shocked at 24 hpf and allowed to develop until 52 hpf. The front of enteric neural crest 247 migratory chains was directly visualized by live confocal microscopy along the gut in 248 GFP⁻ and GFP⁺ embryos as depicted in **Fig. S1C**. In control embryos, *sox10*:mRFP⁺ 249 250 neural crest cells were present as a smooth, collective chain along the level of the midgut (Fig. S1D,E). By contrast, in dnRAR-GFP⁺ embryos, the enteric neural crest front was 251 252 delayed in the foregut and appeared less organized and more dispersed, although they 253 still remained in a migratory chain (Fig. S1F,G). These observations suggest that dampening RA pathway activity during the gut invasion stage is sufficient to delay 254 migration of enteric neural crest cells along the gut in vivo. These results phenocopy the 255 observations with DEAB treatment, leading to gut colonization defects (Fig. 3), consistent 256 257 with a model in which RA is temporally required for the efficient migration of enteric 258 neural crest along the foregut and midgut in vivo.

259

Loss of RA leads to stalled, disorganized migration of enteric neural crest along
 the foregut *in vivo*

262 To test the hypothesis that RA is required for the sustained and directed 263 migration of enteric neural crest along the foregut, DMSO and DEAB treated sox10:mRFP⁺ larvae were live imaged during the second day of development using 264 265 confocal time-lapse microscopy, as shown in Fig. 4A. In particular, we focused on the migratory front of the enteric neural crest chains in our time-lapse analysis. In control 266 larvae, the sox10:mRFP⁺ enteric neural crest chain front was observed migrating 267 268 caudally along the level of the midgut in a smooth, collective chain (Fig. 4B,B', Movie 1). In contrast, the DEAB treated sox10:mRFP⁺ enteric neural crest chain front was detected 269 more rostrally along the foregut; although initially in a chain, the cells failed to progress 270 caudally along the gut. Instead, enteric crest cells began to detach from one another at 271 both the leading edge and along the trailing edge of the chain (Fig. 4C,C'; arrows, Movie 272 273 2). Interestingly, not only did cells detach from one another, but they also began turning away from the chain both dorsally and ventrally (Fig. 4C,C'; arrows). Confirming the lack 274 275 of progress along the gut, cell tracking analysis to follow leading cells over the course of 3 hours showed that control enteric neural crest cells collectively migrate caudally, while 276 DEAB treated enteric crest fail to progress along the gut (Fig. 4D,E). Strikingly, the 277 migratory chain front in DEAB treated larvae dissociated later at 72 hpf, when compared 278 with controls (Fig. 4F). These data demonstrate that loss of RA as enteric neural crest 279 280 migrate along the foregut leads to adverse consequences on neural crest migratory 281 progress and the catastrophic loss of enteric neural crest chain integrity—leading to loss 282 of directed chain migration along the gut (**Fig. 4G**). These results reveal a novel 283 mechanism by which RA modulates enteric neural crest chain migration and suggest that 284 RA is temporally required to ensure collective migration of the enteric neural crest chain along the foregut in vivo. 285

286

Temporal loss of RA does not affect enteric neuron differentiation, but leads to cell death and subsequent intestinal aganglionosis

289 We next examined if loss of RA, in addition to affecting migration, might have an 290 effect on enteric neuron differentiation. During normal development, enteric neural crest 291 cells migrate to the caudal end of the hindgut by ~72 hpf and begin to terminally 292 differentiate into distinct enteric neuronal subtypes thereafter (Olsson et al., 2008; Olden et al., 2008; Uyttebroek et al., 2010; Taylor et al., 2016; Heanue et al., 2016). To 293 investigate whether DEAB-treatment resulted in loss of differentiated neurons along the 294 295 foregut, larvae were processed for whole mount immunochemistry using antibodies against Hu for differentiated neuronal cell bodies, acetylated tubulin for axons, or the 296 neurotransmitter serotonin (5HT). In control larvae, Hu⁺, 5HT⁺ and acetylated tubulin⁺ 297 neurons and axons were detected along the entire foregut (Fig. 5A,C; arrows; Movie 3); 298 however in DEAB treated larvae, neurons were generally missing (Fig. 5B,D; Movie 4). 299 300 Only one small cluster of ~3 neurons in the foregut of one DEAB treated larval fish was detected (Fig. 5B; arrow), indicating that temporal loss of RA results in intestinal 301 aganglionosis. 302

During enteric neurogenesis in zebrafish, the first Hu⁺ neurons are detected between ~52-54 hpf along the foregut (Olsson et al. 2008; Olden et al. 2008; Taylor et al. 2016). To determine if DEAB treatment alters the production of enteric neurons along the foregut at 52 hpf, and thus the initiation of enteric differentiation, *-8.3phox2bb*:kaede⁺/Hu⁺ neurons were examined in transverse section using immunohistochemistry in control and DEAB treated larvae. In control larvae, there was an average of 5.25 -

309 8.3phox2bb:kaede⁺ enteric progenitors along the left and right sides of the foregut, of

which an average of 1.5 were Hu⁺ (Fig. 6A-A", C, D), 28.5% of the Kaede⁺ cells. In 310 311 DEAB treated larvae, an average of 13 -8.3phox2bb:kaede⁺ enteric progenitors was 312 detected; however they were distributed adjacent to the gut and around the ventral 313 mesenchyme (Fig. 6B-B", arrows) rather than in the foregut, in agreement with our previous finding using the sox10:GFP line (Fig. 3K-M). Among the -8.3phox2bb:kaede⁺ 314 enteric progenitors in DEAB treated larvae, an average of 6 were Hu⁺(Fig. 6D), 46.2% of 315 316 the Kaede⁺ cells, demonstrating an overall increase in the number and proportion of 317 neurons present.

318 The above results show that neurons differentiate properly in DEAB treated embryos and in fact are increased in number in the foregut when compared with controls. 319 However, at later stages, neurons are missing. A possible explanation is that enteric 320 321 neurons fail to survive after differentiation. To test this possibility, we performed 322 immunostaining against activated-Caspase3 in control and DEAB- treated larvae at 70 hpf. As expected, in larvae examined in whole mount using confocal z-stack imaging, we 323 324 observed elevated cell death within the enteric neural crest after loss of RA compared with controls (Fig. 6E-E", F-F", Movie 5, Movie 6). As secondary confirmation, we used 325 immunohistochemistry on transverse sections through the foregut to examine the 326 presence of apoptotic cells. The results confirm the presence of Caspase3⁺ neural crest 327 328 cells near the gut and in surrounding ventral mesencyhme, compared with controls (Fig. 329 **6G-G**", **H-H**"; **arrows**). These observations indicate that enteric cells were undergoing cell death by this point, leading to a paucity of enteric neurons. 330

Collectively, these data demonstrate that temporal loss of RA results in
 accumulation of enteric neuronal progenitors along the foregut, which is later
 accompanied by cell death and a dramatic loss of enteric neurons resulting in HSCR-like

intestinal aganglionosis. Taken together with cell migration defects noted in Figs. 3, 4

and 5, these results reveal a previously unrecognized stage-dependent role for RA during

the ontogeny of enteric neuron formation along the gut *in vivo*.

337

338 The transcription factor Meis3 functions downstream of RA in the neural crest to

339 affect colonization of the gut

340 Next, we asked what transcriptional regulatory events might function downstream of RA signaling to regulate enteric colonization. One candidate is the transcription factor 341 Meis3, a known RA-target gene in the zebrafish mesoderm (Waxman et al., 2008). We 342 previously showed that its reduction resulted in colonic aganglionosis due to inefficient 343 migration of enteric neural crest cells along the gut (Uribe and Bronner, 2015). To 344 345 determine if meis3 expression was altered following modulation of RA, we performed in 346 situ hybridization after either temporal DEAB treatment, heat shock in hsp70:dnRAR-GFP embryos or exogenous RA treatment (Fig. 7). The results show that expression of meis3 347 348 was missing in the vagal region and diminished in the foregut of DEAB treated embryos, compared with controls as viewed in whole mount (Fig. 7A,B; arrow). Similarly, following 349 heat shock attenuation, the expression of meis3 was restricted to a smaller domain along 350 the foregut of GFP⁺ embryos, when compared with heat shock controls, (Fig. 7C,D; 351 352 arrowheads). Conversely, RA-treated embryos displayed an expansion of meis3 353 expression within the foregut (Fig. 7F; arrow) as well as a rostral expansion in the 354 hindbrain (Fig. 7F; bracket). By contrast, meis3 expression in control embryos was 355 restricted to the vagal-level hindbrain and foregut (Fig. 7E; bracket). These data 356 demonstrate that *meis3* expression is altered following modulation of RA levels, 357 suggesting that *meis3* may lie functionally downstream of RA.

358 To directly test a possible epistatic relationship between RA and Meis3 in enteric 359 colonization, we asked whether ectopic expression of meis3 was sufficient to rescue gut 360 colonization and enteric neuronal differentiation following temporal loss of RA. To this 361 end, -8.3phox2bb:Kaede embryos were injected with 25 or 40 pg of meis3 mRNA at the 1-cell stage, developed until 24 hpf and incubated in 10µM DEAB, or DMSO for control, 362 363 until 73 hpf. To assay for possible differences between enteric gut colonization and 364 differentiation, whole mount immunochemistry was performed to detect the presence of -365 8.3phox2bb:Kaede⁺/Hu⁺ neurons along the gut. The number of embryos from each condition exhibiting "normal colonization" (full gut), "partial colonization" (up to level of 366 midgut) or "no colonization" (loss of neurons along whole gut) were scored. Control 367 larvae possessed enteric neurons along the entire length of the gut (Fig. 8A), with 100% 368 exhibiting normal colonization (Fig. 8I). Injection of meis3 alone, at 25 pg (Fig. S2) and 369 at 40 pg (Fig. 8B) had no effect on enteric neuron distribution along the gut, with 100% of 370 larvae displaying normal colonization (Fig. 8I). In contrast, DEAB treated larvae 371 372 exhibited neuronal loss along the entire length of the gut (Fig. 8E), with 100% of larvae exhibiting no colonization (Fig. 8I), consistent with our earlier experiments (Fig. 3,5). 373 Injection of *meis3* mRNA together with DEAB-treatment led to a rescue in enteric neuron 374 localization through the midgut (partial colonization) in ~67% of injected larvae (Fig. 8F,I; 375 Fig. S2). 376

To determine whether *meis3* functions downstream of RA specifically in the neural crest, rather than in the mesenchyme, we ectopically expressed *meis3* within neural crest by creating a construct *sox10:meis3-P2A-mcherry*. Its effects were compared with the control construct, *sox10:turq-P2A-mcherry*, which expresses the fluorophore Turquoise2. Injection of the control and *meis3* constructs were performed as

schematized in **Fig. 9A**. Embryos exhibiting mosaic mCherry⁺ expression in vagal neural 382 383 crest cells at 24 hpf were sorted (Fig. 9B,C) and incubated in either DMSO, for control, or DEAB until 75 hpf. Control larvae expressing the control or meis3 construct had Hu⁺ 384 385 enteric neurons along all levels of the gut (Fig. 9D,E). In contrast, DEAB treated larvae expressing the control construct, sox10:turg-P2A-mcherry, displayed little or no 386 387 colonization, with the exception of a single neuron (Fig. 9F,H, arrow). Conversely, DEAB treated larvae expressing the construct sox10:meis3-P2A-mcherry contained Hu⁺ enteric 388 neurons in a mosaic manner throughout the gut at all levels (Fig. 9G,I; yellow arrows). 389 These results show that ectopic expression of *meis3* within neural crest is sufficient to 390 partially rescue colonization of the gut following RA loss, confirming that meis3 functions 391 392 downstream of RA signaling in the neural crest to affect enteric colonization of the gut in 393 vivo.

394

395 Ectopic ret expression is sufficient to rescue gut colonization after RA loss

Another factor known to be important for ENS development is the receptor 396 tyrosine kinase, RET, where its loss in both mouse and zebrafish leads to the 397 catastrophic death of vagal neural crest prior to gut entry (Natarajan et al., 2002; Pachnis 398 et al., 1998; Jain et al., 2004; Heanue et al., 2008). In avian explant cultures, RA has 399 400 been shown to induce the expression of *Ret* in vagal neural crest cells (Simkin et al., 401 2013). To examine whether ectopic expression of ret alone or in combination with meis3, was sufficient to rescue gut colonization following RA attenuation, we assayed enteric 402 403 neuron localization along the gut following injection of ret, or ret co-injected with meis3, in 404 DMSO or DEAB treated larvae. Ectopic expression of ret (Fig. 8C), or ret plus meis3 (Fig. 8D), did not affect colonization of the gut, with 100% of injected larvae exhibiting 405

normal colonization (Fig. 8I). In contrast, injection of *ret*, or *ret* plus *meis3*, in DEAB
treated embryos led to rescue of enteric neuron localization through the midgut.
Approximately 54% of *ret* injected larvae and 75% of *ret* + *meis3* injected larvae
displayed colonization caudally to the midgut (Fig. 8G,H,I). These data show that *ret*and/or *meis3* were sufficient to rescue enteric colonization following temporal DEAB
treatment. Moreover, their co-expression increases the percentage of larvae exhibiting
midgut colonization after loss of RA (54% versus 75%).

413 Quantification of cell number along the rescued larval guts revealed that 414 expression of meis3, ret, or meis3 + ret was sufficient to partially rescue enteric neuron 415 number following loss of RA (Fig. 8J). Whereas control larvae exhibited an average of 81.5 neurons, DEAB treated larvae exhibited an average of less than one neuron, 0.75 416 417 (Fig. 8J, p<.0001). In contrast, DEAB treated larvae expressing meis3 contained an 418 average of 20.75 neurons (p<.0001), larvae expressing ret contained an average of 22.75 neurons, and larvae expressing meis3 + ret contained 26.25 neurons (p<.001) 419 (Fig. 8J). Neuron number was slightly increased when comparing larvae expressing 420 meis3 versus meis3 + ret following DEAB treatment, but not significantly (20.75 vs. 421 26.25, p=.1373). 422

423 Collectively, these epistatic rescue data show that *meis3* and *ret* function
424 downstream of the RA signaling pathway during early ENS formation, thus expanding the
425 known signaling network underlying enteric nervous system formation.

426

427 **RA** affects colonization of the gut during early foregut invasion stages

Finally, we examined the precise time period during which RA functions by
treating embryos for discrete time windows: from 28-36, 36-48 or 48-73 hpf with DMSO

430 or DEAB. Embryos were allowed to develop to 73 hpf to assay for the presence of Hu⁺ 431 neurons along the gut. Control embryos displayed neurons along all levels of the gut 432 (Fig. 10A, E), with 100% exhibiting normal colonization. In comparison, larvae treated 433 from 48-73 hpf also exhibited neurons along all levels of the gut (Fig. 10D,E), with 95% of 434 larvae exhibiting normal colonization. Thus, the presence of RA during later phases of 435 enteric migration is not required for colonization. In contrast, treatment with DEAB from 28-36 hpf resulted in loss of colonization along the whole gut in 76% of larvae, with the 436 other 24% exhibiting partial colonization (Fig. 10B,E). Treatment from 36-48 hpf lead to 437 438 64% of larvae with partial colonization of the gut, 17% with no colonization, and 19% 439 appearing largely normal (**Fig. 10C,E**). Taken together, these time-course experiments indicate that RA is required during the early neural crest foregut invasion stage plus 440 441 subsequent migration of neural crest cells along the foregut, but is less important thereafter. These results are concordant with our previous results suggesting that RA is 442 critical for gut invasion and neural crest migration along the foregut (Fig. 3, 4, 5), playing 443 444 a stage-specific role in ENS formation.

445

446 **Discussion**

We have identified a critical role for retinoic acid in orchestrating the collective chain migration and survival of enteric neural crest cells along the developing gut *in vivo*. Using zebrafish as a model system, our findings reveal that RA functions in a temporally important manner; when enteric neural crest cells migrate into and along the foregut, to modulate the efficient collective chain migration of enteric neural crest cells to ensure complete colonization of the gut (**Fig. 10F**). Specifically, we found that loss of RA during the migratory phase caused enteric neuronal progenitors to stall in their migratory

454 progress and accumulate near the foregut, subsequently dissociating from collective 455 chains into detached cells (Fig. 3, 4, 6). Although following RA reduction we observed no 456 effect on ability to differentiate into enteric neurons, they subsequently underwent 457 apoptosis, leading to intestinal aganglionosis in which the gut lacked nearly all neurons 458 (Fig. 5, 6). Conversely, application of exogenous RA during enteric neural crest 459 migration enhanced the number of enteric neural crest and increased the total number of differentiated neurons along the gut in vivo (Fig. 2). Moreover, we find that the 460 transcription factor Meis3, as well as the receptor RET, can partially rescue the enteric 461 colonization defects caused by loss of RA (Fig. 8), thus identifying molecular effectors 462 that influence ENS development downstream of RA signaling. Cumulatively, these 463 results provide novel insights into the cellular and molecular mechanisms by which the 464 465 early ENS is created and suggest that Vitamin A/RA-coupled susceptibility to HSCR may occur precisely within a narrow time window during embryonic development. 466

Our results are consistent with previous literature suggesting that retinoid 467 468 availability and abundance during ENS development can lead to adverse consequences on ENS formation. For example, it was found that depletion of endogenous RA (via 469 knockout of Aldh1a2) caused a failure in ENS formation due to the severe loss of vagal 470 neural crest cells prior to gut entry (Neiderreither et al., 2003), achieved by rescuing early 471 embryonic lethality in Aldh1a2^{-/-} mice with RA. However, the partially rescued embryos 472 473 exhibited a myriad of other embryonic defects, making the precise role of RA in the ENS difficult to parse. Similarly, dietary Vitamin A (retinol) deficiency, coupled with genetic 474 predisposition, has been shown to cause HSCR in a mouse model (Fu et al. 2010). In 475 this study, the authors showed that while Rbp4^{-/-} mice displayed largely normal ENS 476 development, added restriction of maternal dietary Vitamin A during ENS development 477

478 depleted retinoid levels and led to colonic aganglionosis in more than half of the mice 479 analyzed. These results suggest that optimal levels of Vitamin A are necessary for gut 480 colonization in genetically predisposed mice; however how Vitamin A deficiency 481 functionally affects enteric neural crest migration, proliferation or differentiation in vivo 482 was not fully addressed. In vitro and gut explant studies further support a role for RA in 483 enteric neural crest migration, proliferation and differentiation. For example, incubation in excess RA caused expansion of isolated enteric neural crest cells cultured from E12.5 484 mice, a time point in which enteric neural crest have migrated into the colon, and an 485 increase in their neuronal differentiation (Sato and Heuckeroth, 2008). In slice cultures of 486 487 E12.5 mid-small gut, application of a RAR inhibitor, BMS493, reduced net migration distance traveled by enteric neural crest in response to a GDNF gradient and reduced the 488 489 density of enteric neural crest along colon tissue in gut explant, contributing to colonic hypoganglionosis (Fu et al., 2010). Additionally, RA application to quail vagal neural 490 491 crest cells prior to and during gut entry conferred ability to migrate efficiently in chains 492 along aneural gut explant tissue (Simkin et al., 2013).

493 The present work in zebrafish extends these findings in a system where it is possible to examine enteric neural crest development in vivo, in a temporally defined 494 495 manner. We discovered that blockade or enhancement of RA signaling, using 496 pharmacological (DEAB and RA) and genetic means (hsp70:dnRAR), during enteric 497 neural crest migration stages resulted in altered enteric neural crest development in the foregut: exogenous RA application led to increased enteric neuron numbers during larval 498 499 stages, whereas temporal attenuation of RA caused loss of colonization (aganglionosis) 500 of the gut. Live time-lapse confocal analysis of migrating enteric neural crest cells 501 revealed compromised chain migration and integrity, leading to the accumulation of

502 enteric crest within the foregut. Although this had no effect on enteric neurogenesis,

subsequently enteric neuronal progenitors became ectopically localized in the ventral
 mesenchyme and underwent apoptosis that ultimately led to intestinal aganglionosis.

505 Of particular interest, we identified Meis3 as a functional player downstream of 506 RA in the developing ENS. Meis3 is a known RA response gene in the zebrafish lateral 507 plate mesoderm (Van der Velden et al., 2012; Waxman et al., 2008), in the neural 508 ectoderm (Kudoh et al., 2002) and in developing mouse limbs (Qin et al., 2002). Meis3 509 has also been discovered to function with RA to pattern the early frog hindbrain (Dibner 510 et al., 2001). We showed that loss of Meis3 results in decreased enteric precursor 511 migration along the gut that leads to colonic aganglionosis (Uribe and Bronner, 2015). The present results suggest that Meis3 functions downstream of RA to influence enteric 512 513 neural crest cell migration and development. Our results show that Meis3 is downregulated after loss of RA and upregulated with excess RA. Moreover, ectopic 514 515 expression of *meis3* within neural crest partially rescues gut colonization after loss of RA, 516 suggesting that meis3 is downstream of RA in the neural crest (Fig. 9). These experiments demonstrate a vital role for meis3 as a RA effector important for enteric 517 colonization. To date, this is the first report to functionally describe a transcription factor 518 downstream of RA to influence ENS development in any system. 519

In addition to *meis3*, our data suggest that RA functions to affect the RET-Gfra1Gdnf pathway. Gfra1 is a glycosyl phosphatidylinositol (GPI) attached receptor
expressed on enteric neural crest cells along with Ret (Worley et al., 2000; Airaksinen
and Saarma, 2002; Natarajan et al., 2002; Shepherd et al., 2004). GDNF is expressed in
the gut mesenchyme that enteric neural crest cells migrate through (Worley et al., 2000;
Natarajan et al. 2002; Reichenbach et al., 2008). Following loss of *Ret* or *GDNF*, vagal

526 neural crest cells undergo apoptosis before and as they enter the foregut environment, 527 leading to loss of enteric progenitors (Schuchardt et al., 1994; Sanchez et al., 1996; 528 Moore et al., 1996; Pichel et al., 1996). Notably, application of RA to the early neural 529 tube in chicken embryos (Robertson and Mason, 1995) and in culture (Simkin et al., 530 2013) expands the domain and/or upregulates the expression of *Ret.* Consistent with this, we find that ectopic expression of ret mRNA is sufficient to rescue midgut 531 532 colonization following loss of RA and to partially rescue enteric neuron number (Fig. 8), 533 suggesting that RET is downstream of the RA during ENS development in vivo. In 534 addition, ectopic expression of ret plus meis3 increased the percentage of embryos that 535 exhibited midgut colonization rescue following DEAB treatment (75%) compared with meis3 alone (67%) or ret alone (54%), suggesting that ret and meis3 may function 536 537 together to affect caudal colonization of the gut downstream of RA signaling. In contrast, 538 ret plus meis3 did not significantly enhance the number of enteric neurons following 539 DEAB treatment (Fig. 8J) compared with *meis3* alone or *ret* alone. This suggests that the 540 extent to which cells have migrated caudally may be separable from the total numbers of enteric neurons present in the whole gut, with respect to meis3 and ret. In light of these 541 542 data, our results suggest that ret and meis3 may act in a shared pathway to influence enteric neural crest migration, but not total cell number, along the gut. 543

Ret mutations are known to be a major cause of HSCR in humans and the conservation of this function has been demonstrated in numerous model organisms, such as mouse and zebrafish (Natarajan et al., 2002; Pachnis et al., 1998; Jain et al., 2004; Heanue et al., 2008). Due to its significance in the etiology of HSCR, much research has focused on elucidating the molecular mechanisms underlying its function, regulation and spatial distribution during development. Recently, a study examining the conserved

regulatory enhancers that modulate the expression of *Ret* in humans, and often mutated in human HSCR patients, has identified that Rarb directly binds an enhancer element, RET -7, in conjunction with two other enhancers, RET -5.5 and RET +3, bound by the transcription factors Gata2 and Sox10, respectively (Chatterjee et al., 2016). This study implicates RA signaling in modulation of *Ret* expression via *Rarb*, therefore suggesting that RA may be functionally important for development of the ENS in humans and the etiology of HSCR.

557 Examination of gut mesenchyme using a BAC transgenic reporter of hand2 558 expression revealed no qualitative difference in gut mesenchyme tissue along the 559 anterior-posterior length of the gut tube in live larvae treated with DEAB versus control (Fig. 3). However, we have not ruled out the possibility that differentiation of the intestinal 560 561 mesenchyme may be altered following RA inhibition. For example, following loss of hand2 in zebrafish, smooth muscle fails to differentiate, coincident with a reduction of 562 enteric neuron colonization and differentiation along the gut (Reichenbach et al., 2008). 563 564 Nonetheless, Seiler et al. discovered that differentiation of smooth muscle in zebrafish occurred after enteric neuron terminal differentiation (Seiler et al., 2010), suggesting that 565 566 enteric colonization and differentiation does not depend upon differentiated gut mesenchyme. Corroborating these data, previous studies in amniotes have shown that 567 568 differentiation of foregut (stomach) mesenchyme depends upon and follows the arrival of 569 enteric neural crest cells (Faure et al., 2016; Bourret et al., 2017); loss of enteric neural crest cells via tissue-specific ablation impairs the Notch signaling pathway in stomach 570 571 mesenchyme and perturbs smooth muscle development (Faure et al., 2016). Thus, it 572 may be difficult to parse whether effects on the mesenchyme are direct or indirect via alterations to the neural crest. 573

574 Our live imaging and perturbation results *in vivo* suggest that retinoid depletion 575 during enteric migratory phases can lead to adverse consequences on enteric neural 576 crest migration and chain formation during ENS development. In the gut environment, RA 577 pathway components are expressed in the mesenchyme, within the epithelium and in 578 neural crest cells. Between E12.5 and E14.5 in the mouse gut, pathway components involved in RA response are present, such as Rara, Rarg and Rxrg, as well as those 579 involved in its synthesis and degradation, such as Aldh1a1-3 and Cyp26a1 (Sato and 580 581 Heuckeroth, 2008). Aldh1a2 is detected within the gut mesenchyme, in agreement with 582 our findings in zebrafish tissue (Fig. 1). Live time-lapse analysis shows that chains of 583 enteric neural crest collectively migrate caudally in a precise and organized manner along the gut tube (Fig. 4) (Harrison et al., 2014; Uribe and Bronner, 2015). By contrast, we 584 585 discovered that RA inhibition stalled enteric neural crest migration into and along the foregut and cells along the leading and trailing edge veered dorsally and ventrally, but did 586 not progress caudally. By 72 hpf, the enteric neural crest chain dissociated along the 587 588 foregut, when compared with control (Fig. 4). These results suggest that RA is necessary for the propelled caudal advance of enteric neural crest along the gut and 589 implicate it in promoting efficient chain migration. Our results in live embryos are 590 591 consistent with previous findings in explant culture, where application of RA to vagal 592 neural crest was sufficient to enhance chain migration and RA inhibition disrupted chain 593 formation of enteric crest in aneural gut tissue (Simkin et al., 2013). RA also has been shown to be important for the formation of lamellapodia on enteric progenitors and to 594 595 prevent accumulation of the phosphatase PTEN within leading edge cells (Fu et al., 596 2010) by modulation of the Rac1-Rho signaling pathway, which influences actin-597 cytoskeletal dynamics (Fukuda et al., 2002; Hall, 2005; Vohra et al., 2007). Therefore, it

is possible that localization of polarized intracellular effectors of cell migration is affectedfollowing loss of RA.

600 In mouse gut explants, enteric neural crest cells migrate as a web of chains 601 caudally along the growing gut tube (Young et al., 2014), with solitary cells roaming 602 chaotically in a random walk near the vicinity of the chains. Net advancement of the 603 wave front occurred more rapidly than the trailing edge ensuring colonization of the hindgut. In contrast, in our time-lapse movies of the zebrafish gut, we did not detect 604 evidence for solitary enteric neural crest near the collective chain. However, following RA 605 606 inhibition, we did identify solitary enteric neural crest in the ventral mesenchyme 607 surrounding the gut (Fig. 4, 6). It could be that RA normally inhibits enteric neural crest 608 dispersal along the gut and maintains the sole presence of a collective chain. Zebrafish 609 enteric neural crest, therefore, may represent a simplified method of chain migration in response to cues and provides a tractable model to study detailed mechanisms of this 610 611 particular behavior in vivo.

612 Taken together, our results are the first in vivo study to show a temporal requirement for Retinoic Acid in enteric neural crest migration in vivo in any species. The 613 use of zebrafish as a model further enables us to examine the effects of RA loss using 614 live imaging. We find that enteric neurogenesis commences in the absence of RA, but 615 then is followed by neuronal death. These data suggest a model in which vagal neural 616 617 crest cells residing within the post-otic vagal region respond to local RA signaling cues and commence migration toward the foregut entrance. Once within the gut 618 619 mesenchyme, enteric neural crest cells are exposed to RA and Meis3-RET signaling as 620 they migrate caudally in response to a GDNF gradient. Sustained RA signaling and its downstream activation of Meis3 ensures collective enteric neural crest chain integrity as 621

they navigate through the gut mesenchyme and ensures the survival of nascent enteric neurons (Fig, 10F). Future studies focused on whether RA pathway components function to affect the expression of Ret and other HSCR-associated genes directly, or via a conserved regulatory element, using a combination of mouse and zebrafish models, will greatly enhance our understanding of the molecular and genetic mechanisms underlying construction of the ENS *in vivo*.

- 628
- 629 Materials and Methods
- 630

631 Zebrafish Maintenance and lines

632 Zebrafish (*danio rerio*) were maintained at 28.5°C on a 13-hour light/11 hour dark cycle.

633 Animals were treated in accordance with California Institute of Technology IACUC

634 provisions. The following zebrafish lines used include: Wild-type AB (Zebrafish

635 International Resource Center), Tg(-4.9sox10:GFP; ba2Tg) (Carney et al., 2006),

636 Tg(*sox10*:mRFP; *vu234Tg*) (Kucenas et al., 2008), Tg(-8.3phox2bb:Kaede; *em2Tg*)

637 (Harrison et al., 2014), TgBAC(hand2:GFP; pd24Tg) (Yin et al., 2010), Tg(sox17:GFP;

638 s870Tg) (Sakaguchi et al., 2006) and Tg(*hsp70*:dnraraa-GFP; *pd18Tg*) (Kikuchi et al.

639 2011).

640

641 Pharmacological treatments

642 Embryos were de-chorionated and incubated in 1 μM Retinoic Acid (RA) (190269 MP

643 Biomedicals) or 10 μM N,N-diethylaminobenzaldehyde (DEAB) (D86256 Sigma Aldrich)

644 from 24-48 hpf onwards, depending on experiment as described in the Results. Embryos

645 were then either removed from treatment and immediately fixed and processed for

646 downstream experiments or allowed to recover in egg water depending upon

647 experimental endpoint as described in the Results section.

648

655

649 Rescue construct cloning, mRNA synthesis and microinjections

650 The coding sequence of *ret* was amplified using the following primers from 35 hpf cDNA:

651 forward 5' GGCTCCTTTCGCTCGAATCA 3', reverse 5' GCCGTTAGCACAATCACAGC

652 3', ligated into pGEM-Teasy vector (Promega), and sequence verified. ret cDNA was then

653 subcloned into pCS2⁺ vector to create pCS2-*ret*, which was linearized with Clal and used

as template to generate *ret* capped mRNA using the T3 mMessage RNA synthesis kit

656 a template to generate *meis3* capped mRNA with the Sp6 mMessage RNA synthesis kit

(Ambion). pCS2-meis3 (Uribe and Bronner, 2015) was linearized with Notl and used as

657 (Ambion). Tg(-8.3phox2bb:Kaede) embryos were injected with 25 or 40 pg of meis3

658 mRNA and/or 50 pg of ret mRNA at the one-cell stage. Injected embryos and uninjected

659 controls were incubated in either DMSO or DEAB from 24-73 hpf and immediately fixed

660 for anti-Hu, anti-Kaede immunohistochemistry as described below. The number of

661 embryos from each condition exhibiting "normal colonization" (full gut), "partial

colonization" (up to level of midgut) or "no colonization" (loss of neurons along whole gut)
was counted and the percentages represented in bar graph format using Excel software
(Microsoft).

For neural crest tissue-specific rescue experiments, the zebrafish *sox10* promoter (Carney et al., 2006) was subcloned into the Gateway 5' entry vector p5E, to create p5E-*sox10*, via HindIII/SpeI sites. The coding sequence of *meis3* with no stop codon was PCR amplified using the following primers, to add KpnI/SacII sites: 5' CGGGTACCATGGATAAGAGGTATGAG 3', 5'

670 CGCCGCGGTGTGGGCATGTATGTCAAG 3'. The construct pME-Turquoise (Turq)

671 (Oehlers et al., 2014) was used as template to PCR amplify the coding sequence of the

672 fluorophore Turq with no stop codon using the following primers, to add Kpnl/SacII sites:

673 5' ATGGTACCCCATGGTGAGCAAGGGCG 3', 5'

674 CGCCGCGGtCTTGTAGAGCTCGTCCAT 3'. The Gateway middle entry vector pME was

675 PCR amplified to add Kpnl/SacII sites with the following primers: 5'

676 CGCCGCGGACCCAGCTTTCTTGTACA 3', 5' CGGGTACCGGGTCCCCAAACTCACCC

3', creating pME-KS. *meis3* no stop and *turq* no stop PCR fragments were then

678 directionally cloned into pME-KS to create pME-meis3 no stop and pME-turq no stop,

679 respectively. For final construction reactions, p5E-sox10, pME-meis3 no stop, or pME-

680 turq no stop, and p3E-P2A-mCherry (Villefranc et al., 2013; addgene # 26031) was then

681 used in a Gateway LR Clonase II plus reaction (Invitrogen) using the destination vector

pDestTol2pA2 (Kwan et al., 2007) to create pDest-Tol2-sox10-meis3-P2A-mcherry-pA2

and pDest-Tol2-sox10-turq-P2A-mcherry-pA2. Constructs were injected individually into

the 1-cell stage embryo at 25 pg each, along with 30 pg of Tol2 transposase mRNA.

Tol2 mRNA was created by linearizing the construct pCS2-transposase (Kwan et al.,

686 2007) with Not1 and transcribing mRNA using the Sp6 mMessage RNA synthesis kit

687 (Ambion). pDest-sox10-meis3-P2A-mcherry-pA2 and pDest-sox10-turg-P2A-mcherrypA2

688 injected embryos were sorted for mCherry signal at 24 hpf and incubated in DMSO or the

inhibitor DEAB until 75 hpf, then accordingly fixed and subjected to anti-Hu

690 immunohistochemistry to assay for neuronal colonization of the gut tube. Injection-

691 rescue experiments were repeated using three biological replicates, with n=10 for each

692 condition, therefore each replicate N=40; total N=120 across all three replicates.

693

694 in situ hybridization

695 Whole mount *in situ* hybridizations were performed essentially as described (Jowett and

696 Lettice, 1994). The following cDNA constructs were used as templates to generate anti-

697 sense probes: *meis*3 (Rauch et al., 2003), *crestin* (Luo et al., 2001), *rbp5* (formerly *rbp1a*;

698 cDNA #cb465, Zebrafish International Resource Center), crabp2a (cDNA #cb432,

699 Zebrafish International Resource Center), and *aldh1a2* (Feng et al., 2010). Following in

situ hybridization, embryos were processed into 75% glycerol and imaged whole-mount

701 or incubated in 15% sucrose, embedded in 7.5% gelatin and cryosectioned for

702 histological examination. Sections were imaged using a Zeiss Image.M2 Apoptome.2

703 microscope using the DIC image settings and a Plan-Apochromat/.8 20X objective. All

images were processed and cropped using Acrobat Adobe Photoshop CS6 software.

705

706 Immunohistochemistry and image analysis

Larvae were fixed and prepared for immunohistochemistry as previously described (Uribe 707 708 and Bronner, 2015). The following antibodies were used: rabbit anti-GFP 1:500 (Life 709 Technologies, A-11122), goat anti-GFP 1:500 (Abcam, ab6673), mouse anti-pH3 1:500 710 (Abcam, ab14955), rabbit anti-Aldh1a2 1:500 (Genetex, GTX124302), rabbit anti-Kaede 711 1:250 (MBL International, PM102M), mouse anti-HuC/D 1:200 (Invitrogen), rabbit anti-712 Caspase3 1:500 (R&D Systems, AF835) or rabbit anti-5HT 1:1000 (Immunostar). The 713 following secondary antibodies were used: Alexa Fluor Goat anti-Rabbit 488, Donkey anti-Goat 488, Goat anti-Rabbit 568 or Goat anti-Mouse 633 (Invitrogen) was used 1:700. 714 715 DAPI and/or Alexa Fluor 568 Phalloidin (Invitrogen) were included with some secondary 716 antibody incubations to visualize nuclei or F-actin, respectively, on cryosections. Whole 717 mount embryos were processed into 75% glycerol prior to imaging.

718

719 Cryosections were imaged using a Zeiss Image.M2 Apoptome.2 microscope and a Plan-720 Apochromat/.95 korr 40x oil objective and whole-mount embryos were imaged with a 721 Zeiss LSM 800 confocal laser-scanning microscope using a Plan-Apochromat/.8 20X 722 objective and Zen software. For confocal images, z-stacks were acquired laterally along 723 the gut encompassing the region where neural crest cells or enteric neurons were 724 detected, with z-stack intervals ranging from 18-30 microns thick and slices acquired 725 every 1-3 microns. Confocal images were processed and exported as maximum intensity projections using Imaris imaging software (Bitplane) in the form of .tif files and/or 726 rendered as 3D rotation animations in .mp4 format to view whole z-stack tissue.. For cell 727 counts on cryosections, cells were quantified in comparable axial locations along the 728 729 foregut in control and DEAB treated larvae. For counts in whole-mount guts, neurons were counted from maximum intensity Z-stack projections. The average number of cells 730 was depicted in bar graph format using Prism software (Graphpad). Statistical difference 731 between two conditions was determined using Student's t-test analysis with Prism 732 software (Graphpad). The significance threshold was set at .05. All final images were 733 processed and cropped using Acrobat Adobe Photoshop CS6 software. 734

735

736 Heat shock experiments

Tg(*hsp70*:dnraraa-GFP)^{+/-} fish were transferred from 28.5°C to 38°C preheated water at
24 hpf for one hour and then transferred back to 28.5°C to recover for 3 hours. Heat
shocked embryos were then examined and sorted for GFP fluorescence, accordingly
sorted into GFP⁻ and GFP⁺ groups, incubated until 48 or 52 hpf and processed for
immediate downstream experiments.

742

743 Live imaging

Tg(sox10:mRFP), Tg(phox2bb:kaede) or TgBAC(hand2:GFP) larvae were anesthetized 744 745 in low dose tricaine, embedded in 1% low melt agarose dissolved in egg water in a glassbottom imaging chamber (Lab-Tek, chambered #1.0 Borosilicate) and imaged using a 746 20X Plan-Apochromate/ .8 objective, in a 28.5°C heated imaging chamber on a Zeiss 747 LSM 800 confocal laser scanning microscope. For time-lapse analysis, control and 748 DEAB treated larval fish were oriented laterally against the bottom of the imaging 749 750 chamber and the foregut-midgut level of the gut was examined. Z-stacks ~30 microns thick, encompassing the area of gut where enteric neural crest chains were evident, were 751 acquired every 10 minutes. Acquired time-lapse movies were exported as .mp4 files, 752 753 time lapse stills were exported as .tif images and cell track analysis was used to manually track the first few cells of the enteric neural crest chain in control and DEAB treated 754 larvae over the course of 3 hours using an Imaris imaging software system (Bitplane) in 755 756 the Biological Imaging Facility (BIF), Caltech, Pasadena, Ca.

757

758 Author Contributions

R.A.U. and M.E.B designed the study; R.A.U. and S.S.H. performed experiments and
collected data; R.A.U. and M.E.B. analyzed data, drafted the manuscript and obtained
funding.

762

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- 775

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1000 Figure 1. RA pathway components are expressed along the gut during ENS

- 1001 development.
- 1002 (A) Cartoon illustration of a 1-dpf zebrafish embryo depicted laterally to reveal the
- 1003 location of the foregut, midgut and hindgut. ot-otic, d-dorsal, v-ventral, a-anterior, p-
- 1004 posterior
- 1005 (B) Whole-mount *in situ* hybridization against *aldh1a2* at 48 hpf reveals its localization
- 1006 along all levels of the gut tube.
- 1007 (C) Whole-mount in situ hybridization against rbp5 at 48 hpf reveals its localization along
- 1008 all levels of the gut tube.
- 1009 (D) Whole-mount in situ hybridization against crabp2a at 48 hpf reveals its localization
- 1010 along the foregut.
- 1011 (E-E") Cryosection through the level of the foregut depicts Aldh1a2 protein localization
- 1012 (red) in the gut mesenchyme (m), but not in the gut endoderm (e), marked by sox17:GFP.
- 1013 s-somite, yk-yolk
- 1014 (F-F") Cryosection through the level of the foregut depicts Aldh1a2 protein localization
- 1015 (red) in the gut mesenchyme (m) and within neural crest cells migrating in the gut
- 1016 mesenchyme (arrows), marked by *sox10*:GFP. s-somite, yk-yolk.
- 1017 Scale bars: 50 µM
- 1018

1019 Figure 2. RA treatment during enteric neural crest migration stages enhances

1020 enteric neuron numbers

- 1021 (A-B) A dorsal view following whole-mount in situ hybridization at 48 hpf for crestin in
- 1022 control (A) and RA-treated larvae (B) reveals neural crest localization along the embryo.
- 1023 RA-treated embryos display an expansion of neural crest, arrows (B), when compared to
- 1024 control (A). vg-vagal
- 1025 (C-D) Cryosections through the foregut of larvae at 52 hpf following in situ hybridization
- 1026 for crestin in (C) control and (D) RA-treated larvae shows an expanded distribution of
- 1027 *crestin*⁺ neural crest near the vicinity of the gut (arrows), when compared with control
- 1028 larvae. NT- neural tube
- 1029 (E-F) Transverse sections depicting the localization of *sox10*:GFP⁺ neural crest in (E)
- 1030 control and (F) RA-treated larvae shows an expansion of neural crest near the vicinity of
- 1031 the gut (arrows), when compared with control larvae. Scale bars, 50 µM
- 1032 (G) Bar graph to represent the average number of Hu⁺ and 5HT⁺ neurons along the gut in
- 1033 control and RA-treated larvae at 96 hpf. Error bars indicate +/- S.E.M., *n*=6 embryos for
- 1034 each condition.
- 1035 (H-I) Lateral views of the gut at 96 hpf following whole-mount antibody staining to detect
- 1036 Hu⁺/5HT⁺ enteric neurons in (H-H") control and (I-I") RA-treated larvae.
- 1037 (J) Zoomed in view of control larval fish depicted in H.
- 1038 (K) Zoomed in view of RA-treated larval fish depicted in I.
- 1039 Scale bars, 50 µM.
- 1040
- 1041 Figure 3. Temporal loss of RA stalls migration of enteric neural crest within the
- 1042 foregut without affecting enteric neural crest cell numbers
- 1043 (A-B) Whole-mount in situ hybridization against crestin in (A) control and (B) DEAB
- 1044 treated larvae at 52 hpf.

1045 (C-D) Whole-mount *in situ* hybridization against *crestin* in control and DEAB treated

1046 larvae at 72 hpf reveals that enteric neural crest are delayed along the foregut, when

1047 compared with controls. Arrows and yellow asterisk marks caudal end of enteric neural

- 1048 crest migratory front along the gut. yk-yolk
- 1049 (E-F) Live images of -8.3phox2bb:Kaede (E) control and (F) DEAB treated larvae at 72
- 1050 hpf reveals that enteric progenitors are delayed in migration along the foregut, when

1051 compared with control cells along the hindgut. yk-yolk

1052 (G-H) Live confocal projection images of hand2:GFP in (G) control and (H) DEAB treated

1053 larvae at 52 hpf reveals the presence of gut mesenchyme laterally along the gut.

1054 (I-J) Transverse cryosections show that sox10:GFP⁺ cells located near the foregut in (I)

1055 control and (J) DEAB treated larvae. When compared with control sections, DEAB

1056 treated larvae exhibit increased numbers of neural crest in the mesenchyme surrounding

1057 the gut (yellow arrows), while number of neural crest in direct gut contact (white arrows)

1058 are not affected. s-somite, yk-yolk

1059 (K) Bar graph depicting the average number of *sox10*:GFP⁺ neural crest with direct gut

1060 contact or in the surrounding ventral mesenchyme near the gut in control and DEAB

1061 treated larvae. *n*=9 embryos for each condition. Error bars indicate +/- S.E.M. *, p < .05

1062 with Student's *t*-test.

1063 (L) Bright field images of a control (top) and DEAB treated (bottom) larval fish at 48 hpf to 1064 reveal the distribution of melanophores.

1065 Scale bars in panels E-H, 100 μ M; scale bar in panel I-J, 30 μ M.

1066

1067 Figure 4. Temporal inhibition of RA results in stalled migration of enteric neural

1068 crest chains and loss of chain formation along the foregut

- 1069 (A) Cartoon schematic of a zebrafish larval fish to illustrate the timing and location of live
- 1070 imaging experiments in panels B-C. Enteric neural crest migratory front cells along the
- 1071 leading edge were imaged in control and experimental conditions.
- 1072 (B-B',C-C') Cropped time lapse stills over the course of 3 hours showing the enteric
- 1073 neural crest chain front along the foregut-midgut of a sox10:GFP⁺ control larval fish (B-B')
- 1074 and the foregut a DEAB treated (C-C') larval fish. The DEAB treated enteric neural crest
- 1075 chain exhibits delayed migration along the foregut and solitary cells detached from the
- 1076 chain (arrows), while the control enteric neural crest chain front was observed migrating
- 1077 along the midgut collectively.
- 1078 (D-E) Cell tracks of control (D) and DEAB treated (E) enteric neural crest show that
- 1079 DEAB treated enteric neural crest cells fail to progress caudally along the gut, when
- 1080 compared with control.
- 1081 (F) Whole mount immunochemistry against RFP reveals that at 72 hpf, control enteric
- 1082 neural crest chains are maintained along the gut, while DEAB treated enteric neural crest
- 1083 have dissociated along the foregut.
- 1084 (G) Cartoon schematic summarizing the neural crest chain migration phenotype observed
- 1085 following loss of RA along the gut. Red cells depict neural crest migrating in chains along
- 1086 the gut tube in control (top) and in DEAB treated fish (bottom). DEAB treated neural
- 1087 crest chains never make it past the foregut, where they become ectopically localized in
- 1088 the vicinity surrounding the gut.
- 1089 Scale bar in B,C,D and E: 40 μ M; scale bar in F: 50 μ M.

1090

1091 Figure 5. Temporal loss of RA causes total intestinal aganglionosis

- 1092 (A-B) Lateral view of the foregut of a control (A) and DEAB treated larval fish (B) following
- 1093 double immunochemistry against Hu (red) and 5HT (green) shows that control fish
- 1094 successfully differentiate enteric neurons, while DEAB treated fish lack almost all
- 1095 neurons. A small cluster of ~3 neurons was detected in the anterior foregut of the DEAB
- 1096 treated larval fish (arrow in B).
- 1097 (C-D) Lateral view of the foregut of a control (C) and DEAB treated larval fish (D)
- 1098 following immunochemistry against Acetylated Tubulin shows that control fish contain
- 1099 differentiated enteric neurons, along with axons (arrows), while DEAB treated larval fish

19

- 1100 do not.
- 1101 Scale bar in A and C: 30 μ M. Anterior to the left.
- 1102

1103 Figure 6. RA depletion causes accumulation of enteric progenitors in the ventral

- 1104 mesenchyme near the foregut and their apoptosis
- 1105 (A-A"-B-B") At 52 hpf, transverse sections through the foregut of control (A-A") and
- 1106 DEAB treated larvae (B-B") reveal the location of -8.3phox2bbb:Kaede⁺/Hu⁺ enteric
- 1107 progenitors (arrows),
- 1108 (C-D) Bar graphs depict the average number of Kaede⁺ enteric progenitors (C) and Hu⁺
- 1109 enteric neurons (D) in control and DEAB treated larval foregut sections. *n*=5 embryos for
- 1110 each condition. Error bars indicate +/- S.E.M. *, p < .05 with Student's *t*-test.
- 1111 (E-E"-F-F") Whole mount double immunochemistry against activated-Caspase3 (red)
- 1112 and GFP (green) in sox10:GFP control (E-E") and DEAB treated (F-F") larval fish.
- 1113 GFP⁺/Caspase3⁺ cells are present along the foregut of DEAB treated fish, however not
- 1114 detected in control embryos.

- 1115 (G-G", H-H") At 70 hpf, transverse sections through the foregut of control (G-G") and
- 1116 DEAB treated larvae (H-H") show the location of *sox10*:GFP⁺ and Caspase3⁺ cells (red),
- 1117 which reveals neural crest cells that are Caspase3⁺ surrounding the gut in DEAB treated
- 1118 fish, when compared to controls.
- 1119 Scale bar in A, B, G-H: 20 µM; scale bar in E,F: 60 µM
- 1120
- 1121 Figure 7. RA modulates the spatial expression of *meis3* in the vagal and foregut

1122 tissue domains

- 1123 (A-B) Following DEAB treatment from 24-52 hpf (B), the expression area of meis3 in the
- 1124 vagal (vg) region (yellow arrow) and foregut tissue (white arrow) is diminished when
- 1125 compared with control larvae (A).
- 1126 (C-D) Following heat shock induction, dnRaraa-GFP⁺ (D) larval fish exhibit a reduced
- 1127 expression domain of meis3 along the foregut (arrowhead) and vagal region, when
- 1128 compared to heat shock controls.
- 1129 (E-F) Following RA incubation from 24-48 hpf (F), the hindbrain expression of meis3 was
- 1130 rostrally expanded (brackets), as well as the foregut (arrow), when compared with control
- 1131 larvae (E).
- 1132

1133 Figure 8. Ectopic expression of *meis3* and/or *ret* is sufficient to partially rescue gut

- 1134 colonization in embryos temporally lacking RA
- 1135 (A-H) Maximum intensity confocal projection images show Hu⁺/-8.3phox2bb:Kaede⁺
- 1136 enteric neurons along the gut of (A) control larvae, (B) larvae expressing
- 1137 40pg of meis3, (C) larvae expressing 50pg of ret, (D) larvae expressing 40pg meis3 and
- 1138 50 pg ret; (E) DEAB treated larvae, (F) DEAB treated larvae expressing 40pg meis3, (G)

- 1139 DEAB treated larvae expressing 50 pg ret, (H) DEAB treated larvae expressing 40pg
- 1140 meis3 and 50pg ret.
- 1141 (E) Bar graphs depicting the percentage of larvae exhibiting normal colonization (neurons
- 1142 along whole length of gut), partial colonization (neurons present to the midgut) and no
- 1143 colonization (no neurons along the gut).
- 1144 (J) Bar graphs showing the average number of neurons for the rescue conditions shown
- 1145 in A-H. Error bars indicate S.E.M. **, p < .01 with Student's *t*-test.
- 1146 Scale bar in A-D: 60 µM.
- 1147
- 1148 Figure 9. Ectopic expression of *meis3* in the neural crest partially rescues gut
- 1149 colonization following RA inhibition
- 1150 (A) Cartoon schematic summarizing injection and treatment experiments.
- 1151 (B-C) Live images of phox2bb:Kaede/mCherry⁺ 24 hpf embryos injected with (B) pDest-
- 1152 sox10:turq-P2A-mcherry-pA2, or (C) pDest-sox10:meis3-P2A-mcherry-pA2. Lateral
- 1153 views, ot-otic.
- 1154 (D-I) Maximum intensity confocal projections following immunochemistry against mCherry
- 1155 (red) and Hu (cyan) at 75 hpf in larvae expressing (D, F, H) pDest-sox10:turq-P2A-
- 1156 mcherry-pA2, or (E, G, I) pDest-sox10:meis3-P2A-mcherry-pA2, following treatment with
- 1157 DMSO or DEAB, respectively.
- 1158 (H-I) Zoomed in view of the insets from F and G, respectively.
- 1159 Scale bars: 60 µM
- 1160
- 1161 Figure 10. Stage-specific disruption of RA during neural crest entry and migration
- along the foregut, but not thereafter, leads to colonization defects

1163 (A-D) Maximum intensity confocal projections reveal Hu⁺ neurons in lateral views of the

1164 gut at 73 hpf in (A) Control larvae (DMSO treated), and larvae treated with DEAB from

1165 (B) 28-36 hpf, (C) 36-48 hpf, and (D) 48-73 hpf. Scale bar: 70 µM

1166 (E) Bar graphs depicting the percentage of larvae exhibiting normal colonization (neurons

along whole length of gut), partial colonization (neurons present to the midgut) and no

1168 colonization (no neurons along the gut).

1169 (F) Cartoon illustration of the role of RA and Meis3 during enteric colonization of the gut.

1170 RA (cyan) is synthesized along the foregut mesenchyme (beige) concomitant with enteric

1171 neural crest (green) entry into the gut from the vagal neural crest domains. Meis3,

1172 functionally downstream of RA in the neural crest, and/or RET regulate caudal

1173 colonization of the gut during enteric nervous system development. The action of RA

1174 affects enteric colonization primarily during early foregut migration phases (28-48 hpf).

1175

1176 Movie 1: Neural crest migration along the control zebrafish larval gut.

1177 Lateral view of *sox10*:mRFP⁺ neural crest chains migrate caudally from the foregut

1178 through the midgut over the course of 9 hours. Neural crest chains successfully make

1179 their way from foregut to midgut in a smooth, collective chain.

1180

1181 Movie 2: Neural crest migration along the DEAB treated zebrafish larval gut.

1182 Lateral view of sox10:mRFP+ neural crest chain migrate caudally down the foregut,

1183 however the enteric neural crest chain fails to progress into the midgut and over the

1184 course of 3 hours dissociates along the foregut, with individuals neural crest cells

1185 becoming ectopically localized away from the gut tube.

1186

1187 Movie 3: 3D animation depicting enteric neurons along the control zebrafish larval

1188 **gut**.

1189 Rotation movie showing a lateral view of the 4dpf control larval fish foregut stained for Hu

- 1190 (red) and 5HT (green) positive neurons shown in Figure 5A.
- 1191
- 1192 Movie 4: 3D animation depicting enteric neurons along the DEAB treated zebrafish
- 1193 larval gut.

1194 Rotation movie showing a lateral view of the 4dpf DEAB treated larval fish foregut stained

- 1195 for Hu (red) and 5HT (green) positive neurons shown in Figure 5B.
- 1196
- 1197 Movie 5: 3D animation depicting a control larval foregut following anti-Caspase
- 1198 immunostaining.
- 1199 Rotation movie showing a lateral view of the 70 hpf control sox10:GFP⁺ larval fish foregut
- 1200 shown in Figure 6E stained for GFP (green) and activated Caspase3 (red).

1201

1202 Movie 6: 3D animation depicting a DEAB treated larval foregut following anti-

- 1203 Caspase immunostaining.
- 1204 Rotation movie showing a lateral view of the 70 hpf DEAB treated *sox10*:GFP⁺ larval fish
- 1205 foregut shown in Figure 6E stained for GFP (green) and activated Caspase3 (red).
- 1206

1207 Figure S1. Heat shock attenuation of RA signaling function leads to delayed

- 1208 colonization of the gut by enteric neural crest cells.
- 1209 (A-B) crestin whole-mount in situ hybridization in control heat shock (GFP⁻) larvae (A)
- 1210 and heat shock dnRaraa-GFP⁺ larvae (B) at 48 hpf, arrowhead points to anterior foregut

- 1211 region. Dorsal views reveal that neural crest cell entry along the foregut is delayed
- 1212 (arrows) in dnRaraa-GFP⁺ larvae (B'), when compared with control larvae (A'). vg-vagal,
- 1213 fb-fin bud
- 1214 (C) Cartoon schematic to depict genotype and time course of heat shock attenuation
- 1215 experiments shown in panels D-G.
- 1216 (D-G) Live confocal images showing *sox10*:mRFP^{-/+}; *hsp70*:dnRaraa-GFP^{-/+} larvae at 52
- 1217 hpf to reveal enteric neural crest chain localization along the gut. Lateral views show that
- 1218 enteric neural crest reside within the foregut following heat shock induction of dnRaraa-
- 1219 GFP, while control heat shock enteric neural crest are located within the midgut at the
- 1220 same time, asterisks denote caudal end of the migratory chain.
- 1221 Scale bar in D and F, 100 microns; in E and G, 50 microns.
- 1222
- 1223 Figure S2. Ectopic expression of 25 pg of *meis3* is sufficient to partially rescue gut

1224 colonization in embryos temporally lacking RA

- 1225 (A-D") Whole mount double immunochemistry shows Hu⁺/-8.3phox2bb:Kaede⁺ enteric
- 1226 neurons along the gut of (A-A") control, (B-B") DEAB treated larvae, (C-C") larvae
- 1227 injected with 25 pg meis3, and larvae injected with 25 pg meis3 and treated with DEAB
- 1228 (D-D"). While larvae treated with DEAB from 24-73 hpf exhibit a severe loss of enteric
- 1229 neurons (B-B"), embryos treated with DEAB also expressing 25 pg meis3 exhibit a partial
- 1230 rescue of enteric neuron localization along the midgut (D-D").
- 1231





























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