

1 Zebrafish heme oxygenase 1a is necessary for normal development and macrophage migration

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3 Kaiming Luo<sup>1,2</sup>, Masahito Ogawa<sup>2</sup>, Anita Ayer<sup>2,3</sup>, Warwick J Britton<sup>1,4</sup>, Roland Stocker<sup>2,3</sup>, Kazu  
4 Kikuchi<sup>2,5</sup>, Stefan H Oehlers<sup>1,6\*</sup>

5 <sup>1</sup> Tuberculosis Research Program at the Centenary Institute, The University of Sydney,  
6 Camperdown, NSW 2050, Australia

7 <sup>2</sup> Victor Chang Cardiac Research Institute, Darlinghurst NSW 2010, Australia

8 <sup>3</sup> The Heart Research Institute, Newtown, NSW 2042, Australia

9 <sup>4</sup> Department of Clinical Immunology, Royal Prince Alfred Hospital, Camperdown, NSW 2050,  
10 Australia

11 <sup>5</sup> National Cerebral and Cardiovascular Center, Suita, Osaka 564-8565, Japan

12 <sup>6</sup> The University of Sydney, Discipline of Infectious Diseases & Immunology and Marie Bashir  
13 Institute, Camperdown, NSW 2006, Australia

14

15 Corresponding author email: [stefan.oehlers@sydney.edu.au](mailto:stefan.oehlers@sydney.edu.au)

16

## 17 **Abstract**

18 Heme oxygenase function is highly conserved between vertebrates where it plays important roles in  
19 normal embryonic development and controls oxidative stress. Expression of the zebrafish heme  
20 oxygenase 1 genes are known to be responsive to oxidative stress suggesting a conserved  
21 physiological function. Here we generate a knockout allele of zebrafish *hmox1a* and characterize  
22 the effects of *hmox1a* and *hmox1b* loss on embryonic development. We find that loss of *hmox1a* or  
23 *hmox1b* causes developmental defects in only a minority of embryos, in contrast to *Hmox1* gene  
24 deletions in mice that causes loss of most embryos. Using a tail wound inflammation assay we find  
25 a conserved role for *hmox1a*, but not *hmox1b*, in normal macrophage migration to the wound site.

26 Together our results indicate zebrafish *hmox1a* has clearly a partitioned role from *hmox1b* that is  
27 more consistent with conserved functions of mammalian Heme oxygenase 1.

28

29 **Introduction**

30 Heme oxygenases are a highly conserved family of proteins that catalyze the breakdown of heme to  
31 carbon monoxide (CO), ferrous iron and biliverdin [1, 2]. In mammals, there are two isoforms of  
32 heme oxygenase (HMOX): the inducible HMOX1, and the constitutively expressed HMOX2 [3].  
33 Both isoforms contain a conserved 24-amino acid sequence known as the “heme binding pocket”,  
34 within which a conserved histidine residue acts as the ligand for heme iron [4, 5]. Heme is used as  
35 substrate and cofactor for both isoforms, but the physiological properties and regulation of the two  
36 proteins are different [4].

37

38 Under basal conditions in mammals, HMOX1 is highly expressed in tissues such as spleen, liver,  
39 and bone marrow, while it is almost undetectable in other tissues [6]. Expression of HMOX1 is  
40 induced by multiple stresses including oxidative stress, heme, iron starvation, inflammatory  
41 cytokines, and physical tissue injury [7, 8]. Conversely, mammalian HMOX2 is constitutively and  
42 highly expressed in the brain and testes and does not generally respond to stress conditions [4, 9,  
43 10].

44

45 The role of HMOX1 in regulating the immune response has been demonstrated in models of tissue  
46 injury or different diseases such as ischemic lung/liver injury, atherosclerosis, and non-  
47 hemochromatosis liver diseases [11-14]. In 1999, the first case of HMOX1 deficiency was reported  
48 in a young boy. He showed severe inflammatory phenotypes and accelerated atherosclerosis. The  
49 patient died when he was only six years old [15]. This case suggests a critical role of *Hmox1* in  
50 protecting humans against inflammation and atherosclerotic vascular disease. Consistent with this,  
51 irradiated mice transplanted with bone marrow cells from *Hmox1*<sup>-/-</sup> mice show enhanced

52 atherosclerotic lesions together with a greater macrophage content and increased inflammatory  
53 cytokines including monocyte chemotactic protein 1 [16].

54

55 The whole genome duplication event in the teleost lineage resulted in two homologs (*hmox1a* and  
56 *hmox1b*) of human *HMOX1* in zebrafish genome. Both genes are inducible under oxidative stress  
57 conditions. However, *hmox1a* has been shown to be expressed at a significantly higher level during  
58 the early developmental stage and in response to oxidative stress conditions than *hmox1b* [17, 18].

59

60 Until now, no *hmox1a* mutant zebrafish allele has been generated to study the function of heme  
61 oxygenase in zebrafish. Using a combination of CRISPR-Cas9 and TALEN mutagenesis  
62 techniques, we have studied the function of zebrafish *hmox1a* in development and leukocyte  
63 biology.

64

## 65 **Methods**

### 66 Zebrafish husbandry

67 Adult zebrafish were maintained at Centenary Institute and embryos were obtained by natural  
68 spawning followed and raised in E3 media at 28-32°C (Sydney Local Health District AWC  
69 Approvals: 16-037 and 17-036). All work with the mutant allele and the *Tg(mpeg1:gfp)* transgenic  
70 line were performed in a WT TU/AB genetic background, CRISPR work with the  
71 *Tg(mfap4:turquoise)* transgenic line was performed in a WT AB genetic background.

72

### 73 Generation of a *hmox1a* mutant allele by TALEN mutagenesis

74 The CHOPCHOP web tool was used to predict TALEN pairs targeting *hmox1a*. The selected  
75 TALEN information is listed in Supplementary Table 1. Constructs were assembled using Golden  
76 Gate assembly and cloned into pCS2TAL3DDD and pCS2TAL3RRR vectors [19-21].

77

78 Genome editing was performed as described [21]. Briefly, TALENs, used to generate mutant  
79 zebrafish, were constructed using the Golden Gate assembly method [19, 20]. mRNAs were  
80 synthesized using the Transcription Kits (Invitrogen) and diluted to 50 ng/µL for microinjection.  
81 Samples were prepared immediately before each injection. mRNAs were injected into zebrafish  
82 embryos at the early one-cell stage to 10% of the volume of the cell (~1 nL). Injected embryos were  
83 raised until they reached adulthood and then outcrossed to wild-type fish. Embryos from these  
84 crosses were used for screening to identify mutants. The resulting mutant line is *hmox1a*<sup>vc42</sup>.

85

86 After initial genotyping characterisation, experimental work was carried out on embryos derived  
87 from F3-F5+ generation adults.

88

#### 89 Genotyping

90 For *hmox1a*<sup>vc42</sup> genotyping, a piece of fin was cut off from larval or adult zebrafish for DNA  
91 extraction. PCR was performed with the *hmox1a* genotyping primers described in Supplementary  
92 Table 2 using an annealing temperature of 60°C and an extension time of 30 seconds. PCR products  
93 were digested by *HinfI* restriction enzyme at 37°C for 8-24 h and the digests were resolved by  
94 agarose gel electrophoresis.

95

#### 96 Phenotyping

97 Embryos showing no eyes, severe cyclopisa, small eyes, small trunk somites and heart, reduced  
98 notochord or missing floor plate were described as ‘abnormal’. Embryos showing no gross  
99 abnormalities were described as ‘normal’.

100

#### 101 CRISPR/Cas9 Gene Editing Technique

102 The gRNA target sites for each gene were designed using <https://www.crisprscan.org>, gRNA  
103 oligonucleotide sequences are listed in Supplementary Table 3. The gRNA templates were

104 amplified by PCR with scaffold reverse primer then were transcribed with HiScribe™ T7 High  
105 Yield RNA Synthesis Kit (NEB) [22]. One cell stage embryos were injected with 1 nL of a pre-  
106 incubated mixture containing 200 ng/µL of the four gRNAs, or eight gRNAs in the double  
107 knockdown, and 2 ng/µL Cas9.

108

109 Imaging

110 Imaging was carried out on Leica M205FA and DM6000B, and Deltavision Elite microscopes as  
111 previously described [23-25].

112

113 Drug treatments

114 Copper-protoporphyrin IX (CuPP) (Frontier) and tin-protoporphyrin IX (SnPP) (Frontier) were  
115 dissolved in 0.05M NaOH and then diluted in pH 8.0 Tris-HCl solution to a stock concentration of  
116 10 mM. All compounds were used at a final concentration of 10 µM. Embryos were treated with  
117 drugs within 3 h of egg fertilization and changed every three days for developmental studies or  
118 immediately after tail wounding and not changed for the duration of the macrophage migration  
119 assays.

120

121 Quantitative Real-time PCR

122 Total RNA was extracted from homogenates using Trizol (Thermofisher) and cDNA was  
123 synthesized with the High Capacity cDNA Synthesis Kit (Applied Biosystems). qPCR reactions  
124 were performed on a LightCycler® 480 System. Gene expression was quantified by the delta-delta  
125 C<sub>T</sub> method normalized to host *bact*. Sequences of primers are listed in Supplementary Table 2.

126

127 Tail wounding assay

128 Caudal fin amputation was performed on 4 dpf embryos. Zebrafish larvae were anaesthetized using  
129 tricaine for wounding and imaging experiments. Embryos were cut posterior to the notochord and

130 then recovered to fresh E3 and kept at 28°C. Macrophages were imaged at 6 and 24 h post  
131 wounding (hpw) after tail-fin incision of fluorescent reporter zebrafish larvae.

132

133 **Statistics**

134 All ANOVA and Student's *t* tests as appropriate for the number of comparisons were carried out  
135 using Graphpad Prism. Each data point indicates a single animal unless otherwise stated.

136

137 **Results**

138 **Generation of zebrafish *hmox1a*<sup>vc42</sup> mutant allele**

139 The traditional method to establish a mutant animal model is to induce a premature stop codon  
140 caused by nucleotide deletion. In several cases however, phenotypes in knockout mutant animals  
141 using this method have been shown to lack specificity compared to antisense knockdown methods  
142 [26-29]. Recently, it was determined that nonsense-mediated, decay-induced genetic compensation  
143 can cause different phenotypes between knockout mutants and knockdown methods [26]. Animal  
144 models established with in-frame mutation or mutant targeting at the gene promoter area have a  
145 reduced probability of genetic compensation [26]. To minimize the potential effects of genetic  
146 compensation by the paralogs *hmox1b*, *hmox2a*, and *hmox2b*, we generated an in-frame mutation  
147 allele of *hmox1a* targeting the ATG start codon in exon 2 (Fig. 1A).

148

149 An in-frame ATG was found in exon 3 at amino acid 37, which was predicted to be active when the  
150 initial ATG was deleted. The translated amino acid fragment from the second ATG start site in the  
151 mutant mRNA sequence was predicted to be non-functional as the heme-binding histidine is absent  
152 in the predicted amino acid sequence (Fig. 1A).

153

154 To generate the mutant allele, TALEN-injected embryos were raised to adulthood and three F0 fish  
155 with mutations in somatic tissue were individually outcrossed with wild-type (WT) fish. After

156 sequencing of F1 fish, we identified an in-frame 26 bp deletion mutant *hmox1a* allele with deletion  
157 of the original start codon, resulting in a predicted mutant protein lacking the functional histidine  
158 (Fig. 1B). The mutant allele was designated as *hmox1a*<sup>vc42</sup>, and offspring were propagated for  
159 analysis (Fig. 1C).

160

161 Characterization of *hmox1a*<sup>vc42</sup> expression

162 Genomic sequencing revealed the ‘AG’ splice acceptor site preceding exon 2 was deleted in  
163 *hmox1a*<sup>vc42</sup>. Because there was no splice acceptor attached at the beginning of exon 2 to define the  
164 termination of an intron in front of it, the mutant exon 2 would be treated as a part of a long intron  
165 till the next splice acceptor before exon 3 (Fig. 2A). Thus, we hypothesized that mature mutant  
166 *hmox1a*<sup>vc42</sup> mRNA would skip exon 2.

167

168 To explore this hypothesis, RNA was isolated from homozygous *hmox1a*<sup>vc42/vc42</sup> and WT zebrafish  
169 and cDNA was synthesized via reverse transcription. Primer F1 targeting the start of exon 1 was  
170 used as forward primer and primer R1 targeting the end of exon 7 was used as reverse primer (Fig.  
171 2A). The mutant *hmox1a* exon 1-7 amplicon was slightly smaller than WT as expected from the 41  
172 bp size of exon 2 (Fig. 2B). The exon skipping in *hmox1a* mRNA was further confirmed by  
173 sequencing the cDNA amplicon (Fig. 2C).

174

175 To confirm the effect of the *hmox1a*<sup>vc42</sup> allele on *hmox1a* transcripts, qPCR analysis was performed  
176 to examine the expression of sections of the *hmox1a* mRNA. Primers spanning the exon 1-2 and 2-3  
177 junctions were chosen for the first analysis (F2/R2; Fig. 2A). As expected, *hmox1a* exon 2 junction-  
178 containing mRNA was almost undetectable in the mutants (Fig. 2D).

179

180 Conversely, it was hypothesized that other exon sequences, including exon 1 and exon 3-7, should  
181 still maintained in *hmox1a*<sup>vc42</sup> transcripts. To investigate this hypothesis, primers (F3/R3; Fig. 2A)

182 targeting exon 4 and exon 5 regions were used for qPCR analysis at the same time. Mutant exons 4  
183 and 5 of *hmox1a* mRNA were sustained and not degraded (Fig. 2E), indicating that nonsense-  
184 mediated decay is not triggered by the *hmox1a*<sup>vcc42</sup> allele.

185

186 Morphologic analysis of *hmox1a*<sup>vcc42/vcc42</sup> zebrafish

187 The percentage of homozygous *hmox1a*<sup>vcc42/vcc42</sup> embryos generated from *hmox1a*<sup>vcc42</sup> heterozygote  
188 in crosses slightly decreased from 3 to 7 dpf and further gradually decreased to 30 dpf; from 30 dpf  
189 onwards, the percentage of homozygous mutant embryos was stable (Table 1).

190

191 To investigate the reason for the loss of homozygous mutants, 3 dpf embryos from sibling matched  
192 *hmox1a*<sup>WT/WT</sup> and *hmox1a*<sup>vcc42/vcc42</sup> adult in crosses were collected for morphologic analysis.  
193 Embryos showing no eye, severe cyclopia or small eyes, trunk somites and heart, reduced  
194 notochord and missing floor plate were described as 'abnormal' (Fig. 3A). Embryos that showed no  
195 gross abnormalities were described as 'normal'. While WT embryos did not display abnormal  
196 morphology (0%, n= 368), around 20% of *hmox1a*<sup>vcc42/vcc42</sup> embryos displayed an abnormal  
197 phenotype with shorter length at 3 dpf (Fig. 3B). The percentage of dead *hmox1a*<sup>vcc42/vcc42</sup> embryos  
198 at 11 dpf correlated with the proportion of abnormal *hmox1a*<sup>vcc42/vcc42</sup> embryos at 3 dpf (Fig. 3C,  
199 Table 2). This analysis suggests that a minority of zebrafish *hmox1a*<sup>vcc42/vcc42</sup> embryos undergo an  
200 aberrant morphogenetic process and are unable to survive embryogenesis.

201

202 It is known that embryo morphogenetic development is not only affected by gene expression but  
203 also depends on maternal factors that are required for processes prior to the activation of the zygotic  
204 genome. To identify whether *hmox1a* is a maternal factor, embryos from *hmox1a*<sup>vcc42/vcc42</sup> females  
205 crossed with *hmox1a*<sup>+/vcc42</sup> males and *hmox1a*<sup>+/vcc42</sup> females crossed with *hmox1a*<sup>vcc42/vcc42</sup> males  
206 were screened separately. No differences were observed between these crosses (Fig. 3D),  
207 suggesting that the morphological phenotype was independent of the paternal genotype.

208

209 The 20% of *Hmox1*-deficient mice that survive until adulthood display smaller size and less body  
210 weight [30]. To investigate whether zebrafish *hmox1a* deficiency also affects adult morphology,  
211 *hmox1a*<sup>vcc42/vcc42</sup> adult zebrafish were assessed for morphology, body length and weight. In contrast  
212 to *Hmox1*<sup>-/-</sup> mice, *hmox1a*<sup>vcc42/vcc42</sup> adult zebrafish developed without gross abnormalities compared  
213 to their WT clutch mates (Fig. 3E). Body length and weight was also comparable between  
214 *hmox1a*<sup>vcc42/vcc42</sup> and WT clutch mate adults (Fig. 3F). These results indicate zebrafish *hmox1a* may  
215 have a different impact on embryogenesis and development compared to mouse *Hmox1*, or that  
216 genetic compensation by another Hmox family member plays an important compensatory role  
217 during the development of *hmox1a*<sup>vcc42/vcc42</sup> zebrafish.

218

219 Zebrafish *hmox1b* partially compensates for loss of *hmox1a* during embryonic development

220 As a consequence of the whole genome duplication event in zebrafish, there are additional paralogs  
221 *hmox1b*, *hmox2a*, and *hmox2b* that may increase the survival of *hmox1a*<sup>vcc42/vcc42</sup> zebrafish  
222 compared to mice, where *Hmox2* is the only related gene to *Hmox1*.

223

224 We first examined the expression of *hmox1b*, *hmox2a* and *hmox2b* in *hmox1a*<sup>vcc42/vcc42</sup> zebrafish.  
225 Significant upregulation of *hmox1b* was detected in phenotypically normal *hmox1a*<sup>vcc42/vcc42</sup>  
226 embryos compared to WT embryos and *hmox1b* expression was even higher in abnormal  
227 *hmox1a*<sup>vcc42/vcc42</sup> embryos than the other phenotypically normal embryos of either genotype (Fig.  
228 4A). The zebrafish *hmox2* genes, *hmox2a* and *hmox2b*, were expressed at a slightly higher level in  
229 abnormal *hmox1a*<sup>vcc42/vcc42</sup> embryos relative to WT embryos, and there was no change in the  
230 expression of either *hmox2a* or *hmox2b* in normal homozygous *hmox1a*<sup>vcc42/vcc42</sup> embryos (Fig. 4A).  
231 Upregulation of *hmox1b*, but not of *hmox2a* or *hmox2b*, was also seen in *hmox1a* crisprants (Fig.  
232 4B).

233

234 Knockdown of *hmox1a* caused the appearance of visually similar developmental defects to those  
235 seen in homozygous *hmox1a*<sup>vcc42/vcc42</sup> embryos however there was a basal rate of pre-3 dpf embryo  
236 death caused by the microinjection that obscured any statistically significant difference in rate of  
237 developmental defects in *hmox1a* crispants (Fig. 4C, Table 2). Knockdown of *hmox1b* also caused  
238 some developmental defects similar to the loss of *hmox1a* but at a statistically similar rate to the  
239 embryo death in scrambled controls and overall developmental defects in *hmox1a* crispants (Fig.  
240 4C, Table 2). Double knockdown of *hmox1a* and *hmox1b* by CRISPR-Cas9 increased the rate of  
241 developmental defects compared to scramble control embryos across 3-7 dpf and increased the rate  
242 of developmental defects compared to *hmox1b* knockdown embryos at 3 dpf (Fig. 4C, Table 2).

243

244 Treatment of developing WT embryos with the Hmox inhibitor tin protoporphyrin (SnPP) caused  
245 increased mortality after 9 dpf compared to copper protoporphyrin (CuPP) controls (Fig. 4D, Table  
246 2). Together, these results demonstrate a conserved role for zebrafish Hmox1a and Hmox1b in  
247 embryonic development.

248

249 Conservation of Hmox1a regulation of macrophage migration

250 To investigate the role of *hmox1a* in macrophage biology, we crossed the *hmox1a*<sup>vcc42</sup> allele into the  
251 *Tg(mpeg1:GFP)*<sup>vcc7</sup> background.

252

253 As *Hmox1* has been reported to be essential for hematopoiesis [31, 32], the total fluorescent area of  
254 macrophages was measured by fluorescence microscopy in 4 dpf *Tg(mpeg1:GFP)*<sup>vcc7</sup> zebrafish  
255 embryos, where macrophages are marked by GFP. No difference was observed between the WT  
256 control and *hmox1a*<sup>vcc42/vcc42</sup> larvae (Fig. 5A).

257

258 Chemokine production plays a crucial role in the recruitment of leukocytes to sites of inflammation.  
259 The monocyte chemoattractant protein-1 (MCP-1) and its receptor CCR2 are regulated by HMOX1

260 in mouse models of mycobacterial infection [33, 34]. To determine if the relationship between  
261 Hmox1 and macrophage chemoattractant production is conserved in zebrafish, we performed tail  
262 wounding of zebrafish embryos (Fig. 5B). The expression of the chemokine *mcp-1* and receptor  
263 *ccr2* were examined at 6 h post wounding (hpw) by qPCR. Expression of *mcp-1* was reduced while  
264 *ccr2* expression was increased in *hox1a<sup>vcv42/vcv42</sup>* larvae compared to WT littermates (Fig. 5B).

265

266 To investigate the functional consequence of reduced *mcp-1* expression, we quantified macrophage  
267 migration to the wound. There was decreased macrophage accumulation at wound area in  
268 *hox1a<sup>vcv42/vcv42</sup>* homozygous larvae compared to their WT littermates at 24 hours post wounding  
269 (Fig. 5D).

270

271 CRISPR/Cas9 technology was used to knockdown *hox1a* expression in *Tg(mfap4:turquoise)<sup>xt12</sup>*  
272 zebrafish embryos, where the *mfap4* promoter drives macrophage-specific expression comparable  
273 to the *mpeg1* promoter [35]. The number of wound-associated *mfap4*:Turquoise<sup>+</sup> cells was  
274 significantly reduced in *hox1a* crisprants compared to scramble control larvae (Fig. 5E).

275

276 To confirm these genetic results, the HMOX inhibitor, SnPP, was used to treat tail wounded WT  
277 *Tg(mfap4:turquoise)<sup>xt12</sup>* larvae. SnPP decreased macrophage number at the wound site compared to  
278 CuPP-treated controls (Fig. 5F).

279

280 As *hox1b* knockdown had produced some of the developmental defects seen in *hox1a* depletion,  
281 we examined the effect of *hox1b* knockdown on macrophage migration to a wound. Knockdown  
282 of *hox1b* did not affect macrophage migration (Fig. 5G).

283

284 These results demonstrate depletion of *hox1a*, but not *hox1b*, decreases macrophage migration  
285 in zebrafish.

286

287 **Discussion**

288 We report the generation and characterization of a zebrafish with a *hmox1a* mutant allele with  
289 approximately 20% lethality in homozygotes during embryonic development. Morphologic analysis  
290 of *hmox1a*<sup>vcc42/vcc42</sup> embryos revealed the occurrence of developmental disorders from 3 dpf that  
291 progressed to lethality within 14 dpf. These developmental defects were reproduced in double  
292 *hmox1a* and *hmox1b* crispants, and mortality was phenocopied by Hmox1 inhibitor treatment.  
293 Using live imaging and gene expression analyses we demonstrate conservation of the link between  
294 *hmox1a* and macrophage migration in zebrafish. This *hmox1a*<sup>vcc42</sup> allele provides a platform to  
295 investigate the role of *hmox1a* in response to different environmental stresses in zebrafish.

296

297 The first *Hmox1*<sup>-/-</sup> mouse model was established by Poss *et al* in 1997 with total deletion of exons 3  
298 and 4, and partial deletion of exon 5. About 80% of *Hmox1*<sup>-/-</sup> mice succumbed to prenatal lethality  
299 and premature mortality [30]. Our *hmox1a*<sup>vcc42</sup> allele, which only contains deletion of exon 2,  
300 caused disorders of development and premature mortality in about 20% of *hmox1a*<sup>vcc42/vcc42</sup>  
301 embryos. This ~20% mortality rate was similar in *hmox1a* crispants suggesting that the different  
302 survival rates between *Hmox1*<sup>-/-</sup> mice and *hmox1a*-depleted zebrafish is not caused by residual  
303 peptide translated from the *hmox1a*<sup>vcc42</sup> transcript. A possible explanation for the difference in  
304 survival is the high expression of compensatory *hmox1b*, which exists in zebrafish but not in mice.  
305 Our double knockdown experiment where depletion of *hmox1a* and *hmox1b* by CRISPR-Cas9  
306 mutagenesis increased the rate of developmental abnormalities compared to control and *hmox1b*-  
307 depleted, but not *hmox1a*-depleted, embryos suggest *hmox1a* has a more important role in zebrafish  
308 development than *hmox1b*.

309

310 The upregulation of *hmox1b* in *hmox1a*<sup>vcc42/vcc42</sup> embryos might be caused by genetic compensation  
311 resulting from the genomic editing in the *hmox1a* gene locus. Although genetic compensation has

312 been described for a long time [36, 37], the nonsense-mediated decay pathway mechanism has only  
313 been recently described [26, 38]. Mutant alleles that avoid induction of the premature stop codon,  
314 such as the *hmox1a*<sup>vcc42</sup> allele, have been reported to reduce the probability of inducing genetic  
315 compensation [39, 40]. We also observed upregulation of *hmox1b* in *hmox1a* crispants. This  
316 suggests the upregulation of *hmox1b* in *hmox1a*<sup>vcc42/vcc42</sup> embryos is more likely to be caused by the  
317 stress within embryos secondary to the loss of Hmox1a function. Although we observed some  
318 developmental defects in *hmox1b* crispants, the lack of statistical significance compared to scramble  
319 control embryos and lack of significant additional developmental defects in double knockdown  
320 embryos compared to *hmox1a* knockdown alone suggests a biologically insignificant function of  
321 *hmox1b* in zebrafish embryogenesis.

322

323 About 80% of *hmox1a*<sup>vcc42/vcc42</sup> larvae survived until adulthood without gross abnormalities at 90  
324 dpf. This is in accordance with the mouse study where, in the minority of surviving *Hmox1*-  
325 deficient mice, no difference in body weight was reported between *Hmox1*-deficient homozygous  
326 and heterozygous mice until the age of 20 weeks when body size started to be lost in *Hmox1*-  
327 deficient mice [30]. Further morphological analysis of *hmox1a*<sup>vcc42/vcc42</sup> zebrafish over 90 dpf is  
328 necessary to investigate the conservation of this effect in zebrafish.

329

330 It has been reported that expression of MCP-1, also known as CCL2, was significantly increased in  
331 the liver, spleen, bronchoalveolar lavage fluid (BALF) and serum samples of *M. avium* infected  
332 *Hmox1*<sup>-/-</sup> mice compared to WT controls [34, 41]. However, inhibition of HMOX1 in *M. avium*-  
333 infected RAW 264.7 cells reduced the expression of *MCP-1* compared to untreated cells [34]. The  
334 different results from reported studies may be due to the examination of MCP-1 expression at  
335 different levels since in the two experiments using *Hmox1*<sup>-/-</sup> mice, MCP-1 expression was analyzed  
336 at protein level [34, 41], but in the macrophage infection experiment, the gene expression of *MCP-1*  
337 was analyzed [34]. In our study, decreased *mcp-1* transcript levels were detected in tail wounded

338 *hmox1a*<sup>vcc42/vcc42</sup> larvae, which was consistent with the *in vitro* macrophage study [34]. Taken  
339 together, these results suggest that *mcp-1* expression is dependent on *hmox1a* expression in  
340 zebrafish and raise the possibility that macrophage function is altered by increased alternative  
341 activation of macrophages driven by the Ccl2/Ccr2 axis.

342

343 Using live imaging in zebrafish embryos, we provide direct evidence linking Hmox1a to the  
344 regulation of macrophage migration. We found a macrophage migration defect in *hmox1a*<sup>vcc42/vcc42</sup>  
345 larvae that was recapitulated in *hmox1a* crispants and SnPP-treated larvae, but not in *hmox1b*  
346 crispants. This was similar to the smaller effect of *hmox1b* depletion on development and provides  
347 further evidence that *hmox1a* provides the most conserved functions expected of heme oxygenase 1  
348 in zebrafish embryos. Further work with loss of function alleles of the zebrafish *hmox* family are  
349 necessary to determine which gene or genes mediate the expected functions of heme oxygenase 1 in  
350 other biological processes.

351

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363

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456

457 **Figure legends**

458 **Figure 1. Creation of the *hmox1a*<sup>vc42</sup> allele. A.** Alignment of *HMOX1* homologs. Dark grey  
459 background indicates amino acid residues that are identified in all three proteins (Uniprot). Light  
460 grey background indicates amino acid residues that are similar in all four proteins (Uniprot). The

461 purple background indicates the functional histidine. Similarity, identity and functional histidine  
462 were predicted by Uniprot database (<https://www.uniprot.org/align/>). **B.** Partial sequence of the  
463 *hmox1a<sup>vcc42</sup>* allele. TALEN mRNA recognition sequences are blue, and the *HinfI* site in the spacer  
464 region is highlighted in green. Exons are indicated by grey-filled boxes with numbers. Black trace  
465 indicates guanine, blue trace indicates cytosine, green trace indicates adenine and red trace indicates  
466 thymine. **C.** Representative results from PCR-based genotyping using DNA extracted from the fins  
467 of *hmox1a<sup>vcc42</sup>* adult zebrafish followed by *HinfI* digests.

468

469 **Figure 2. Characterization of the *hmox1a<sup>vcc42</sup>* allele mRNA transcript.** **A.** Schematic diagram of  
470 the (predicted) splicing process. Primers used for whole-transcript PCR amplification are indicated  
471 as F1 and R1. Primers F2/R2 and F3/R3 were used for qPCR analysis. **B.** DNA gel electrophoresis  
472 visualization of the exon 1-7 amplicon of mutant *hmox1a* mRNA. **C.** Alignment of the mRNA  
473 transcript of the mutant *hmox1a<sup>vcc42</sup>* allele sequence (bottom) with WT zebrafish *hmox1a* sequence  
474 (top). **D.** Quantification of *hmox1a* exon 2 mRNA by RT-qPCR analysis. **E.** Quantification of  
475 *hmox1a<sup>vcc42</sup>* exons 4-5 mRNA by RT-qPCR analysis. Statistical testing by Student's *t* test, error  
476 bars represent one standard deviation, and each data point represents a biological replicate of pooled  
477 embryos.

478

479 **Figure 3. Morphologic analysis of *hmox1a<sup>vcc42/vcc42</sup>* embryos.** **A.** Images of 3 dpf embryos from a  
480 *hmox1a<sup>+/vcc42</sup>* intercross. Abnormal embryos displayed apparent shorter lengths. Scale bar  
481 represents 0.5 mm. **B.** Quantification of 3 dpf embryo length in WT and *hmox1a<sup>vcc42/vcc42</sup>* normal  
482 and abnormal embryos. Statistical testing carried out by ANOVA. Data is pooled from 3 WT and 4  
483 *hmox1a<sup>vcc42/vcc42</sup>* replicates. **C.** Proportions of abnormal embryo morphology and death during  
484 development. Raw counts are available in Table 2. Statistical testing was performed using Chi-  
485 squared test on raw counts with abnormal and dead embryos treated as one group. **D.** Proportions of  
486 abnormal embryos scored at 3 dpf from female *hmox1a<sup>+/vcc42</sup>*; male *hmox1a<sup>vcc42/vcc42</sup>* (F: +/vcc42; M:

487 vcc42/vcc42) and female *hox1a*<sup>vcc42/vcc42</sup>; male *hox1a*<sup>+/vcc42</sup> (F: vcc42/vcc42; M: +/vcc42)  
488 crosses. Statistical testing was performed using *t* test, and each data point represents a biological  
489 replicate of pooled embryos from an individual crossing. **E.** Representative images of adult (90 dpf)  
490 WT and *hox1a*<sup>vcc42/vcc42</sup> zebrafish from an *hox1a*<sup>+/vcc42</sup> intercross. Scale bar represents 0.5 cm. **F.**  
491 Quantification of the length (left) and weight (right) of adult (90 dpf) *hox1a*<sup>vcc42</sup> zebrafish from an  
492 *hox1a*<sup>+/vcc42</sup> intercross. Statistical testing was performed using *t* test, and error bars represent  
493 SEM.

494

495 **Figure 4. *hox1a* and *hox1b* have non-redundant roles in zebrafish development. A.**  
496 Quantification of *hox1b*, *hox2a* and *hox2b* expression in 3 dpf WT and *hox1a*<sup>vcc42/vcc42</sup>  
497 normal and abnormal embryos. Statistical testing was performed using the one-way ANOVA, error  
498 bars represent one standard deviation, and each data point represents a biological replicate of at  
499 least 10 pooled embryos. **B.** Quantification of *hox1b*, *hox2a* and *hox2b* expression in 3 dpf  
500 scramble control and *hox1a* crispant normal and abnormal embryos. Statistical testing was  
501 performed using the one-way ANOVA, error bars represent one standard deviation, and each data  
502 point represents a biological replicate of at least 10 pooled embryos. **C.** Quantification of abnormal  
503 embryo morphology and death during double *hox1a* and *hox1b* crispant development. Statistical  
504 testing was performed using Chi-squared test on raw counts with abnormal and dead embryos  
505 treated as one group. **D.** Quantification of abnormal embryo morphology and death during  
506 development in embryos treated with SnPP. Raw counts for panels C and D are available in Table  
507 2.

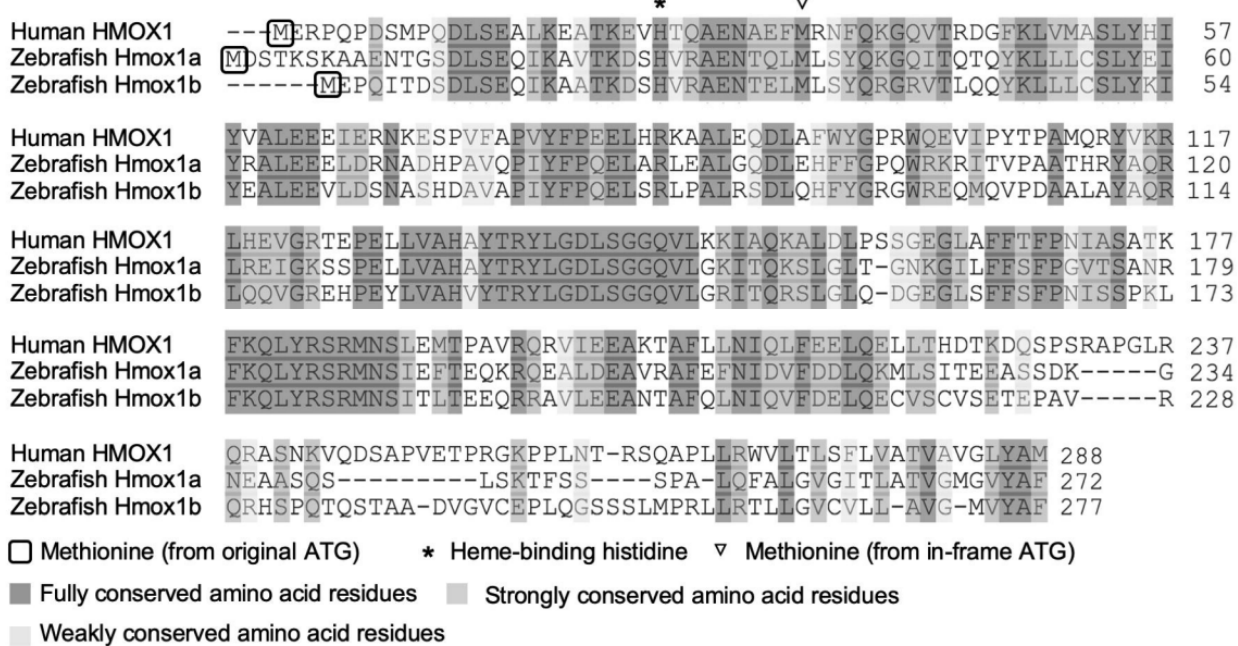
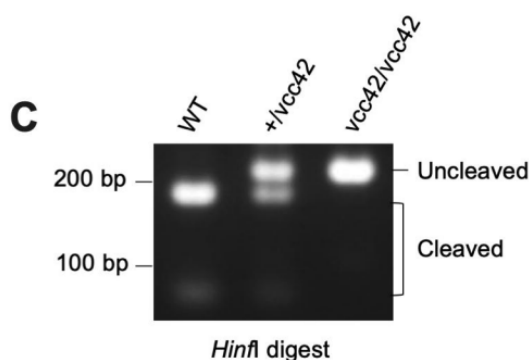
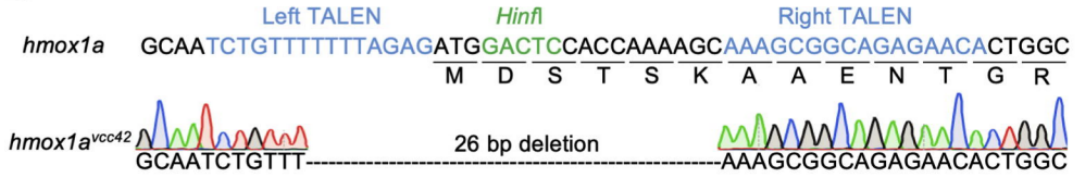
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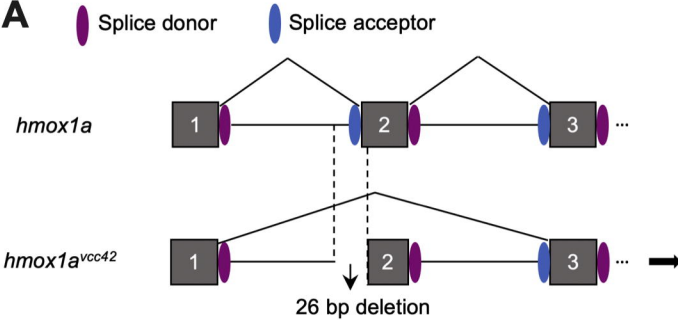
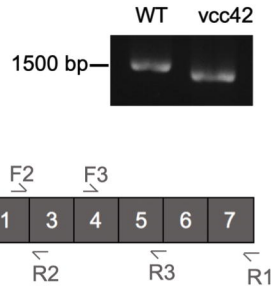
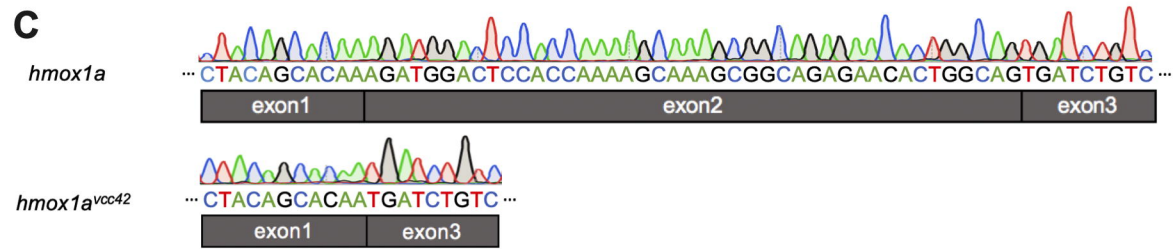
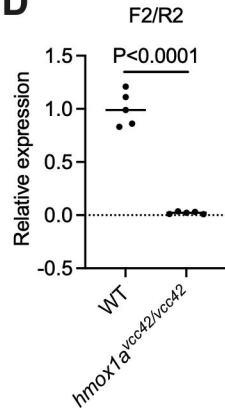
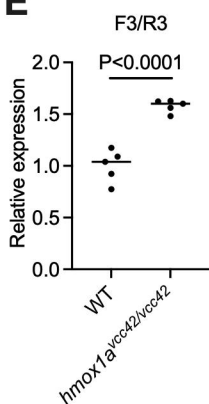
509 **Figure 5. Zebrafish *hox1a* aids macrophage migration. A.** Quantification of total macrophages  
510 of *hox1a*<sup>vcc42/vcc42</sup>, *hox1a*<sup>+/vcc42</sup> and WT larvae. Statistical testing was performed by one-way  
511 ANOVA. **B.** Images represent embryos at 4 dpf (top) and macrophage recruitment to wound site  
512 (bottom) at 6 hpw, red box indicates area measured for macrophage recruitment in panels D-G. **C.**

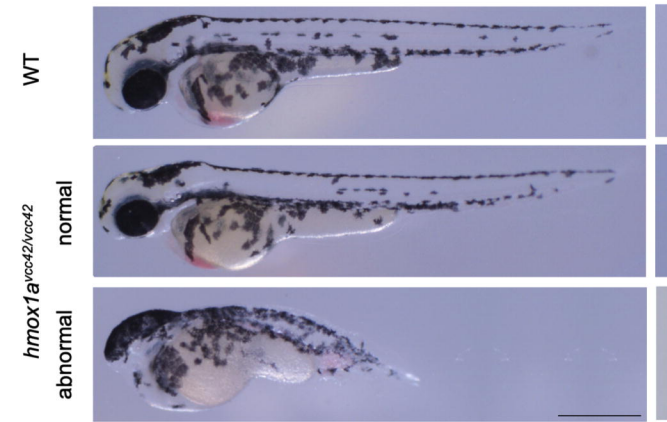
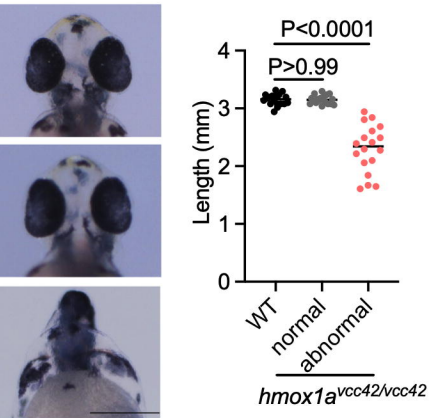
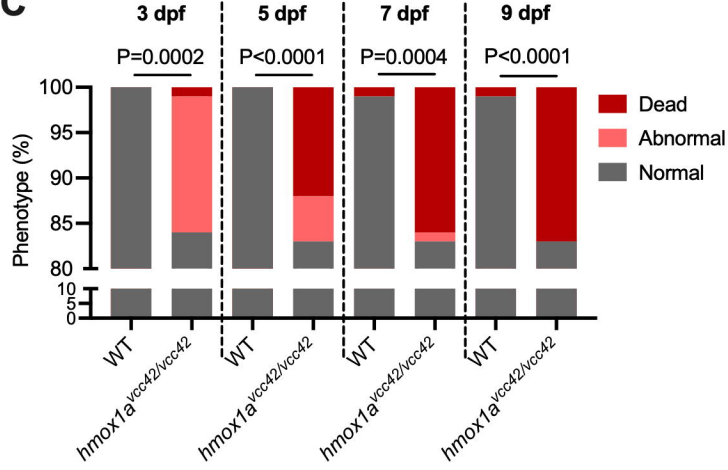
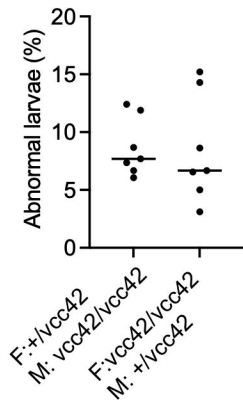
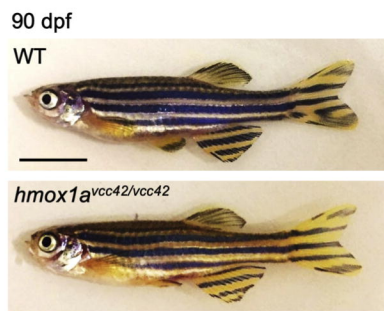
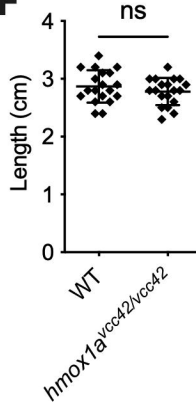
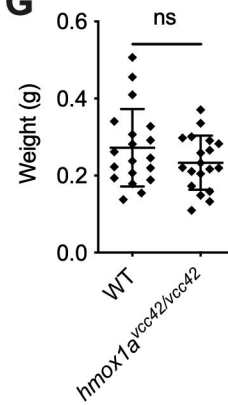
513 Relative quantity of *mcp-1* and *ccr2* transcripts measured by qPCR at 6 hpw, and each datapoint  
514 represents biological replicate of a pool of at least 10 embryos. **D.** Quantification of macrophage  
515 accumulation at wound site by normalized fluorescent area. Data are pooled from two biological  
516 replicates. Statistical testing was performed by one-way ANOVA. **E.** Quantification of  
517 macrophages at wound site by fluorescent area in scramble control larvae and *hmox1a* crispants. **F.**  
518 Quantification of macrophages at wound site by fluorescent area in PPIX or SnPP treated zebrafish  
519 larvae. **G.** Quantification of macrophages at wound site by fluorescent area in scramble control  
520 larvae and *hmox1b* crispants. Statistical testing was performed by *t* test unless otherwise stated,  
521 error bars represent one standard deviation, and data are from one experiment representative of at  
522 least two biological replicates unless otherwise stated.

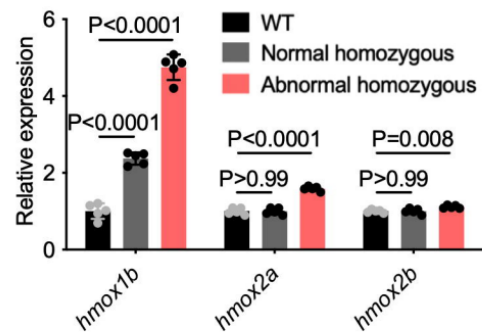
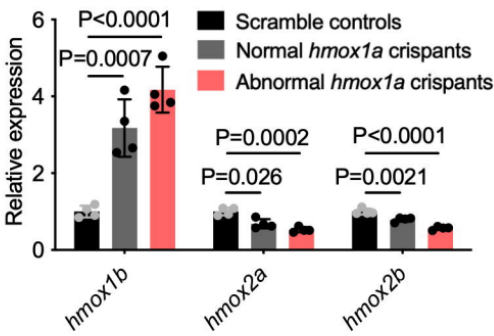
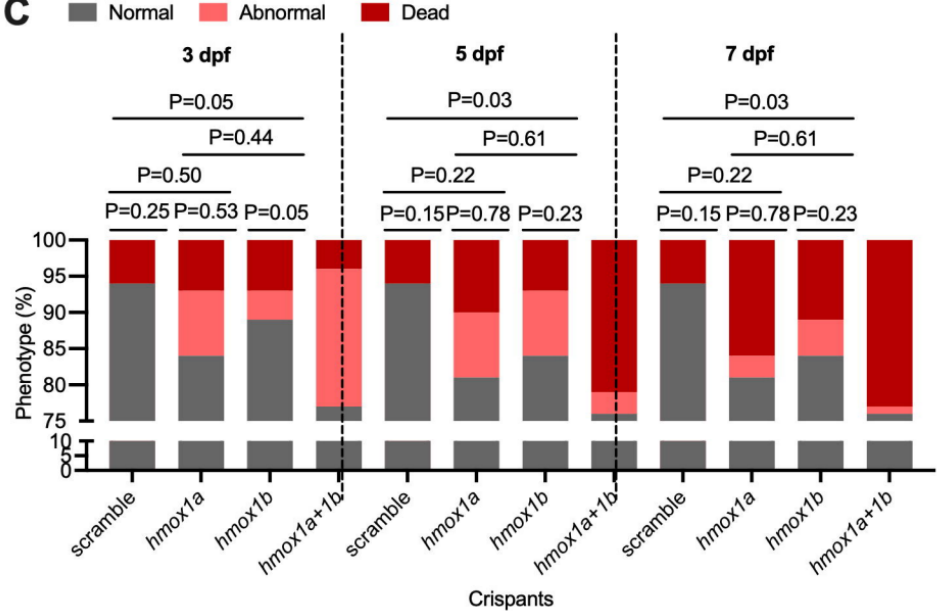
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524

**A****B**

**A****B****C****D****E**

**A** 3 dpf**B****C****D****E****F****G**

**A****B****C****D**