

1 **Glucose inhibits haemostasis and accelerates diet-induced hyperlipidaemia in zebrafish larvae**

2 Simone Morris<sup>1</sup>, Pradeep Manuneedhi Cholan<sup>1†</sup>, Warwick J Britton<sup>1,2</sup>, Stefan H Oehlers<sup>1,3 \*</sup>

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

5 <sup>2</sup> Department of Clinical Immunology, Royal Prince Alfred Hospital, Camperdown, NSW 2050,  
6 Australia

7 <sup>3</sup> The University of Sydney, Discipline of Infectious Diseases & Immunology and Marie Bashir  
8 Institute, Camperdown NSW 2050 Australia

9 <sup>†</sup> Co-corresponding author email: [pradeep.manuneedhicholan@mq.edu.au](mailto:pradeep.manuneedhicholan@mq.edu.au)

10 <sup>\*</sup> Corresponding author email: [s.oehlers@centenary.org.au](mailto:s.oehlers@centenary.org.au)

11

12 **ABSTRACT**

13 Hyperglycaemia damages the microvasculature in part through the reduced recruitment of immune  
14 cells and interference with platelet signalling, leading to poor wound healing and accelerated lipid  
15 deposition in mammals. We investigated the utility of zebrafish larvae to model the effect of  
16 glucose on neutrophil and macrophage recruitment to a tail wound, wound-induced haemostasis,  
17 and chicken egg yolk feed challenge-induced hyperlipidaemia by supplementing larvae with  
18 exogenous glucose by immersion or injection. Neither method of glucose supplementation affected  
19 the recruitment of neutrophils and macrophages following tail transection. Glucose injection  
20 reduced thrombocyte retention and fibrin plug formation while only thrombocyte retention was  
21 reduced by glucose immersion following tail transection. We observed accelerated lipid  
22 accumulation in glucose-injected larvae challenged with high fat chicken egg yolk feeding. Our  
23 study identifies conserved and divergent effects of high glucose on inflammation, haemostasis, and  
24 hyperlipidaemia in zebrafish larvae compared to mammals.

25

26 **INTRODUCTION**

27 Hyperglycaemic damage to the microvasculature is hypothesised to underpin much of the pathology  
28 associated with diabetes in mammals, including perturbations to leukocyte biology, haemostasis,  
29 and the accumulation of lipid laden macrophages in the vessel wall <sup>1-4</sup>. Previous mammalian  
30 research has demonstrated that hyperglycaemia damages the microvasculature resulting in reduced  
31 expression of endothelial adhesion molecules for immune cell recruitment <sup>2,4</sup>. As a result, fewer  
32 neutrophils and macrophages are recruited to diabetic wounds <sup>1,2,5</sup>. Of the macrophages and  
33 neutrophils that eventually arrive, there is a skewing of differentiation towards an inflammatory  
34 phenotype owing to the inflammatory nature of the diabetic wound microenvironment <sup>6,7</sup>.

35

36 Mammals with hyperglycaemia demonstrate perturbed coagulation and platelet signalling, causing  
37 disruption of haemostasis <sup>3,8</sup>. Reduced efficiency of haemostasis results in unstable and ineffective  
38 clots within diabetic foot ulcers <sup>3,9</sup>. Treatments for diabetic foot ulcers can involve platelet and  
39 fibrin therapy, indicating an important role for inadequate fibrin clot production in the ulceration  
40 process <sup>10,11</sup>.

41

42 Hyperglycaemia-induced damage to the microvasculature also increases vascular lipid  
43 accumulation in conjunction with hyperlipidaemia <sup>12</sup>. Hyperlipidaemia and hyperglycaemia are  
44 compounding risk factors for the development of Type 2 diabetes, associated with the ‘Western  
45 Diet’ consisting of fat and sugar alongside limited exercise <sup>13,14</sup>. There are multiple mechanisms by  
46 which hyperglycaemia damages the microvasculature: through the formation of advanced glycation  
47 products <sup>3</sup>, the induction of oxidative stress <sup>15</sup>, interfering with nitric oxide production <sup>16</sup>, and  
48 inducing macrophages to form lipid-laden foam cells <sup>17</sup>. The degradation of the endothelial  
49 structural integrity increases the rate of lipid deposition by providing a physical niche for lipid  
50 infiltration <sup>12</sup>. This can ultimately lead to the development of atheroma, which are common in  
51 diabetic patients <sup>17</sup>.

52

53 These phenotypes have not been previously investigated in fish. Our study investigates the effect of  
54 high glucose environments on inflammation, thrombosis and hyperlipidaemia using a zebrafish  
55 model. Zebrafish are established models for investigating each of these processes in isolation <sup>18-21</sup>.  
56 Pertinent to this study, zebrafish have similar clotting, metabolic, and immune systems to mammals,  
57 and these have been used to provide insight into the shared function of these systems across  
58 vertebrates <sup>19,20,22</sup>. Here we have combined these models to investigate the role of high glucose  
59 environments in disrupting thrombosis and lipid accumulation, but not immune cell recruitment to a  
60 wound, in zebrafish larvae.

61

## 62 METHODS

### 63 Zebrafish Husbandry

64 Zebrafish embryos were produced through natural spawning (Sydney Local Health District Animal  
65 Welfare Committee Approval 17-036). The strains used were *Tg(lyzC:DsRed)<sup>nz50</sup>* to visualise  
66 neutrophils <sup>23</sup>, *Tg(mfap4:turquoise)<sup>xt27</sup>* to visualise macrophages <sup>24</sup>, *Tg(itga2b:gfp)<sup>la2</sup>* to visualise  
67 thrombocytes <sup>25</sup>, and *Tg(fabp10a:fgb-gfp)<sup>mi4001</sup>* to visualise fibrin deposition <sup>26</sup>. From 1-5 days post  
68 fertilisation (dpf) embryos were kept in a dark incubator at 28°C.

69

### 70 Induction of High Glucose Concentrations in the Zebrafish

71 Injection method: Eggs were injected with approximately 15 nmol of glucose, or an equal volume  
72 of PBS as a control within four hours of fertilisation. Immersion method: Dechorionated 2 dpf  
73 embryos were immersed in 5% mannitol (osmolarity control) or glucose dissolved in E3 media,  
74 media was changed daily to reduce microbial growth.

75

### 76 Glucose Oxidase Assay

77 Larvae were snap frozen and stored at -20°C. Larvae were lysed in the buffer solution from the  
78 Amplex Red Oxidase kit (Sigma: A22188), lysates were sedimented by centrifugation, and the

79 Amplex Red Glucose oxidase kit was used to measure the glucose content of the supernatant in  
80 accordance with the manufacturer's instructions.

81

## 82 **Tail Transection Assays**

83 Caudal fin amputations were performed on larvae at 5 dpf. Larvae were anesthetised with 2.5%  
84 (v/v) ethyl-3-aminobenzoate methanesulfonate (tricaine) (Sigma, E10521), wounded posterior to  
85 the notochord using a sterile scalpel and kept in a 28°C incubator to recover as previously described  
86 <sup>27</sup>. Wounded larvae were imaged at 6 hours (neutrophil and macrophages) or 2.5 hours (fibrin and  
87 thrombocytes).

88

## 89 **Hyperlipidaemia Assay**

90 Post 5 dpf, larvae were transferred to a 28°C incubator with a 14/10 hour light/dark cycle. Larvae  
91 were placed in an E3 solution containing 0.05% of emulsified chicken egg yolk from 5 dpf. Each  
92 day, a random sample of larvae were removed, euthanised, and fixed in paraformaldehyde. Larvae  
93 were stained with Oil Red O to quantitate lipid accumulation, as previously described <sup>21,28,29</sup>.

94

## 95 **Imaging and Image Analysis**

96 Larvae were imaged using a Leica M205FA fluorescent microscope. ImageJ software was used to  
97 quantify fluorescent pixel count within 100 µm of the wound site for transgenic wound assays as  
98 previously described <sup>27,30</sup>.

99

100 Oil Red O staining was quantified in ImageJ by adjusting the colour threshold to eliminate non-red  
101 signal. The image was then converted to a binary mask, and the tail region posterior to the swim  
102 bladder was selected to measure the number and area of particles <sup>29</sup>.

103

## 104 **Statistical analysis**

105 Outliers were excluded using a 1% ROUT test. Statistical testing was carried out by ANOVA with  
106 correction for multiple comparisons or Student's *t*-tests as appropriate using GraphPad Prism. Data  
107 are expressed as mean  $\pm$  SD. Every datapoint represents a single embryo unless otherwise noted.

108

109 **RESULTS**

110 **Exogenous glucose exposure increases glucose in zebrafish larvae**

111 To establish the efficacy of the injection and immersion techniques to increase glucose  
112 concentrations in 5 dpf zebrafish larvae, we conducted a glucose oxidase assay. Consistent with past  
113 literature, we observed an increase in the glucose concentration contained within the glucose-  
114 injected and -immersed larvae compared to controls (Figure 1A and 1B)<sup>31,32</sup>.

115

116 Glucose immersion caused sporadic microbial overgrowth and we observed reduced growth of  
117 larvae immersed in either glucose or mannitol, but not in glucose-injected larvae, as measured by  
118 total body area (Figure 1C and 1D) or eye area (Figure 1E and 1F).

119

120 We next analysed the effect of glucose supplementation on the development of key innate immune  
121 cells. We estimated the quantity of macrophages in transgenic *Tg(mfap4:turquoise)*<sup>xt27</sup> larvae,  
122 where macrophages are marked by turquoise fluorescent protein, and found similar numbers of  
123 macrophages in larvae that were injected with or immersed in glucose (Figure 1G and 1H). We  
124 estimated the quantity of neutrophils in *Tg(lyzC:DsRed)*<sup>nx50</sup> larvae, where neutrophils are marked by  
125 DsRed fluorescent protein, and found reduced numbers of neutrophils in glucose-injected larvae but  
126 similar numbers of neutrophils in glucose-immersed larvae (Figure 1I and 1J).

127

128 **Glucose does not affect neutrophil and macrophage recruitment to wounds in zebrafish  
129 larvae.**

130 Altered innate immune cell recruitment to wounds is a conserved feature of hyperglycaemia in  
131 mammals<sup>4,5,33-35</sup>. To determine if this phenomenon was conserved in zebrafish larvae, we utilised  
132 the tail transection wound model which causes reproducible leukocyte recruitment<sup>27</sup> (Figure 2A).  
133 We first performed this assay using transgenic *Tg(mfap4:turquoise)<sup>xz27</sup>* larvae to quantify  
134 macrophage recruitment to the tail wound (Figure 2B). Surprisingly, we observed no difference in  
135 macrophage recruitment between the glucose-injected and control larvae at 6 hours post wounding  
136 (hpw) (Figure 2C). We also observed no difference in macrophage recruitment in the glucose  
137 immersion model (Figure 2D).

138

139 We then used *Tg(lyzC:DsRed)<sup>nz50</sup>* larvae to quantify the recruitment of neutrophils to the tail wound  
140 (Figure 2E). We observed great variability between experiments with two out of four experiments  
141 finding significantly reduced neutrophil recruitment between the control and glucose-injected larvae  
142 at 6 hpw and two out of four experiments finding significantly increased neutrophil recruitment  
143 (Figure 2F). Consistently, we did not observe any difference in neutrophil recruitment in the  
144 glucose immersion model (Figure 2G).

145

146 Together, these results indicate that zebrafish neutrophil and macrophage recruitment is not affected  
147 by exogenous glucose supplementation in zebrafish larvae.

148

#### 149 **Glucose impedes haemostasis in zebrafish larvae**

150 Hyperglycaemia perturbs coagulation and platelet activation in mammals, resulting in ineffective  
151 haemostasis<sup>3,8</sup>. To visualise the effects of high glucose on haemostasis in zebrafish larvae, we first  
152 used the *Tg(itga2b:gfp)<sup>la2</sup>* line to visualise thrombocyte plug formation in the severed blood vessel  
153<sup>25,36</sup>. Following tail transection (Figure 3A), glucose-injected *Tg(itga2b:gfp)<sup>la2</sup>* larvae demonstrated  
154 reduced thrombocyte accumulation (Figure 3B-C). This effect was replicated in glucose-immersed  
155 embryos compared to control embryos (Figure 3D).

156

157 To determine if the reduction in thrombocyte recruitment was mirrored by perturbed clotting, we  
158 conducted tail transections on *Tg(fabp10a:fgb-gfp)<sup>mi4001</sup>* larvae (Figure 3E), which express  
159 fluorescently tagged fibrinogen and allow visualisation of clots<sup>26</sup>. Glucose-injected  
160 *Tg(fabp10a:fgb-gfp)<sup>mi4001</sup>* larvae had reduced fibrin accumulation following tail transection  
161 compared to PBS-injected larvae (Figure 3F). We did not observe any effect of glucose immersion  
162 on the deposition of fluorescently tagged fibrinogen (Figure 3G).

163

164 Together, these results demonstrate that a high glucose environment inhibits the thrombocyte  
165 component of haemostasis in zebrafish larvae.

166

### 167 **High glucose accelerated hyperlipidaemia in zebrafish larvae challenged with a high fat diet**

168 Hyperglycaemia and hyperlipidaemia are intimately associated in mammals<sup>37</sup>. To determine if this  
169 interaction is conserved in zebrafish larvae, we fed glucose-injected larvae a high fat diet consisting  
170 of emulsified chicken egg yolk from 5-7 dpf (Figure 4A). Glucose- and PBS-injected larvae had  
171 similar Oil Red O vascular staining prior to the initiation of chicken egg yolk feeding at 5 dpf  
172 (Figure 4B-C). Quantification of Oil Red O staining revealed glucose-injected larvae had increased  
173 vascular lipid content at one and two days post feeding (Figure 4D-E).

174

175 These results recapitulate the interaction between hyperglycaemia and hyperlipidaemia seen in  
176 mammals<sup>15</sup>, demonstrating that zebrafish larvae are a robust model to study this conserved  
177 interaction.

178

### 179 **DISCUSSION**

180 In this study, we explored the effect of high glucose levels on inflammation, thrombosis, and lipid  
181 accumulation in the zebrafish embryo model. We found that high glucose reduces haemostasis

182 around wound sites. In addition, we determined that high glucose accelerates lipid accumulation  
183 when larvae are challenged with a high fat diet.

184

185 The reduced size of glucose and mannitol immersed larvae compared to control larvae is a major  
186 caveat when interpreting datasets generated with the glucose immersion model. We thus preferred  
187 the injection method for the chicken egg yolk challenge experiments and for the interpretation of  
188 the effects of glucose on larval zebrafish immunity and haemostasis. Despite daily changes of  
189 media, we experienced further difficulty rearing zebrafish larvae from 2 to 5 dpf in 5% glucose  
190 solutions due to microbial overgrowth which may have slowed larval development by consuming  
191 oxygen or damaging larval mucosal surfaces. Although the glucose injection model did not achieve  
192 as high a fold change in glucose at 5 dpf, we did not observe developmental delays and  
193 haematopoiesis was largely comparable to PBS-injected control larvae.

194

195 Hyperglycaemia in mammals causes vascular dysfunction that restricts the recruitment of a broad  
196 range of immune cells to wounds <sup>16</sup>. It was therefore surprising that high glucose appeared to have  
197 no effect on the recruitment of leukocytes to the tail wound in this study. It is possible that high  
198 glucose in zebrafish larvae does not affect abluminal crawling of leukocytes along blood vessels,  
199 since wound-responsive leukocytes have been demonstrated to move predominantly through  
200 interstitial tissue in zebrafish larvae <sup>5,38</sup>. Overall our findings suggest exogenous supply of glucose  
201 to zebrafish larvae may not be a suitable platform for studying the impact of high glucose on  
202 leukocyte biology.

203

204 The relationship between hyperglycaemia and hyperlipidaemia has been previously reported in  
205 various mammalian species <sup>15,32,39</sup>, and a recent study by Wang *et al.* has demonstrated a similar the  
206 interaction of a high cholesterol diet with glucose immersion on vascular lipid accumulation in  
207 zebrafish after 10 days of feeding <sup>32</sup>. Our study using chicken egg yolk as a high fat diet challenge

208 demonstrates a dramatically accelerated accumulation of lipid after just one day of feeding. This  
209 chicken egg yolk feeding-based model, therefore provides a more rapid model to investigate lipid  
210 accumulation.

211

212 In summary, we report glucose-supplemented zebrafish larvae as a tractable platform to investigate  
213 the conserved interactions between glucose and haemostasis, and glucose and diet-induced  
214 hyperlipidaemia.

215

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220

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226

## 227 **COMPETING INTERESTS**

228 The authors declare no competing interests.

229

## 230 **DATA AVAILABILITY STATEMENT**

231 The datasets generated during the current study are available from the corresponding author on  
232 reasonable request

233

234 **REFERENCES**

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329

330 Figure legends

331 Figure 1: Injection and immersion methods increase glucose levels in zebrafish larvae.

332 A. Relative concentration of glucose in 5 dpf larvae that had been injected with 15 nmol glucose as  
333 eggs. Statistical testing by t-test, each data point is representative of a group of n = 10-30 larvae.

334 B. Relative concentration of glucose in 5 dpf larvae immersed in 5% solutions of glucose or  
335 mannitol from 2 dpf. Statistical testing by ANOVA, each data point is representative of a group of n  
336 = 10-30 larvae.

337 C. Total body area calculated from lateral images of 5 dpf glucose-injected larvae. Statistical testing  
338 by t-test. Data are representative of 2 biological replicates.

339 D. Total body area calculated from lateral images of 5 dpf larvae immersed in 5% solutions of  
340 glucose or mannitol from 2 dpf. Statistical testing by ANOVA. Data are representative of 2  
341 biological replicates.

342 E. Eye area calculated from lateral images of 5 dpf glucose-injected larvae. Statistical testing by t-  
343 test. Data are representative of 2 biological replicates.

344 F. Eye area calculated from lateral images of 5 dpf larvae immersed in 5% solutions of glucose or  
345 mannitol from 2 dpf. Statistical testing by ANOVA. Data are representative of 2 biological  
346 replicates.

347 G. Quantification of total macrophage number from lateral images of 5 dpf glucose-injected larvae.  
348 Statistical testing by t-test. Data are representative of 2 biological replicates.

349 H. Quantification of total macrophage number from lateral images of 5 dpf larvae immersed in 5%  
350 solutions of glucose or mannitol from 2 dpf. Statistical testing by ANOVA. Data are representative  
351 of 2 biological replicates.

352 I. Quantification of total neutrophil number from lateral images of glucose-injected 5 dpf larvae.  
353 Statistical testing by t-test. Data are representative of 2 biological replicates.

354 J. Quantification of total neutrophil number from lateral images of 5 dpf larvae immersed in 5%  
355 solutions of glucose or mannitol from 2 dpf. Statistical testing by ANOVA. Data are representative  
356 of 2 biological replicates.

357

358 Figure 2: High glucose does not affect neutrophil and macrophage recruitment to a tail wound.

359 A. Schematic of experiment to measure immune cell recruitment to a tail wound.

360 B. Representative images of macrophage (red) recruitment to a tail wound in glucose-injected  
361 larvae.

362 C. Quantification of macrophage recruitment following tail transection in the glucose injection  
363 model.

364 D. Quantification of macrophage recruitment following tail transection in the glucose immersion  
365 model.

366 E. Representative images of neutrophil (red) recruitment to a tail wound in glucose-injected larvae.

367 F. Quantification of neutrophil recruitment following tail transection in the glucose injection model.

368 Each paired data point represents the average of an biological replicate with  $n > 10$  embryos per  
369 condition.

370 G. Quantification of neutrophil recruitment following tail transection in the glucose immersion  
371 model.

372 Scale bars represent 100  $\mu\text{m}$ . Statistical testing by t-test. Data are representative of 3 biological  
373 replicates.

374

375 Figure 3: High glucose reduced thrombocyte and fibrin accumulation at a tail wound.

376 A. Schematic of experiment to visualise haemostasis following tail transection.

377 B. Representative overlay of thrombocytes (red) at 2.5 hours after tail transection in glucose-  
378 injected larvae.

379 C. Quantification of thrombocyte plug size following tail transection in the glucose injection model.

380 D. Quantification of thrombocyte plug size following tail transection in the glucose immersion  
381 model.

382 E. Representative images of fibrinogen deposition (red) at 2.5 hours after tail transection in glucose-  
383 injected larvae.

384 F. Quantification of fibrin clot size following tail transection in the glucose injection model.

385 G. Quantification of fibrin clot size following tail transection in the glucose immersion model.

386 Scale bars represent 100  $\mu$ m. Statistical testing by t-test. Data are representative of 3 biological  
387 replicates.

388

389 Figure 4: Glucose-injected larvae have increased lipid accumulation following a high fat diet.

390 A. Schematic of the high fat feeding challenge assay to measure lipid accumulation in glucose-  
391 injected larvae.

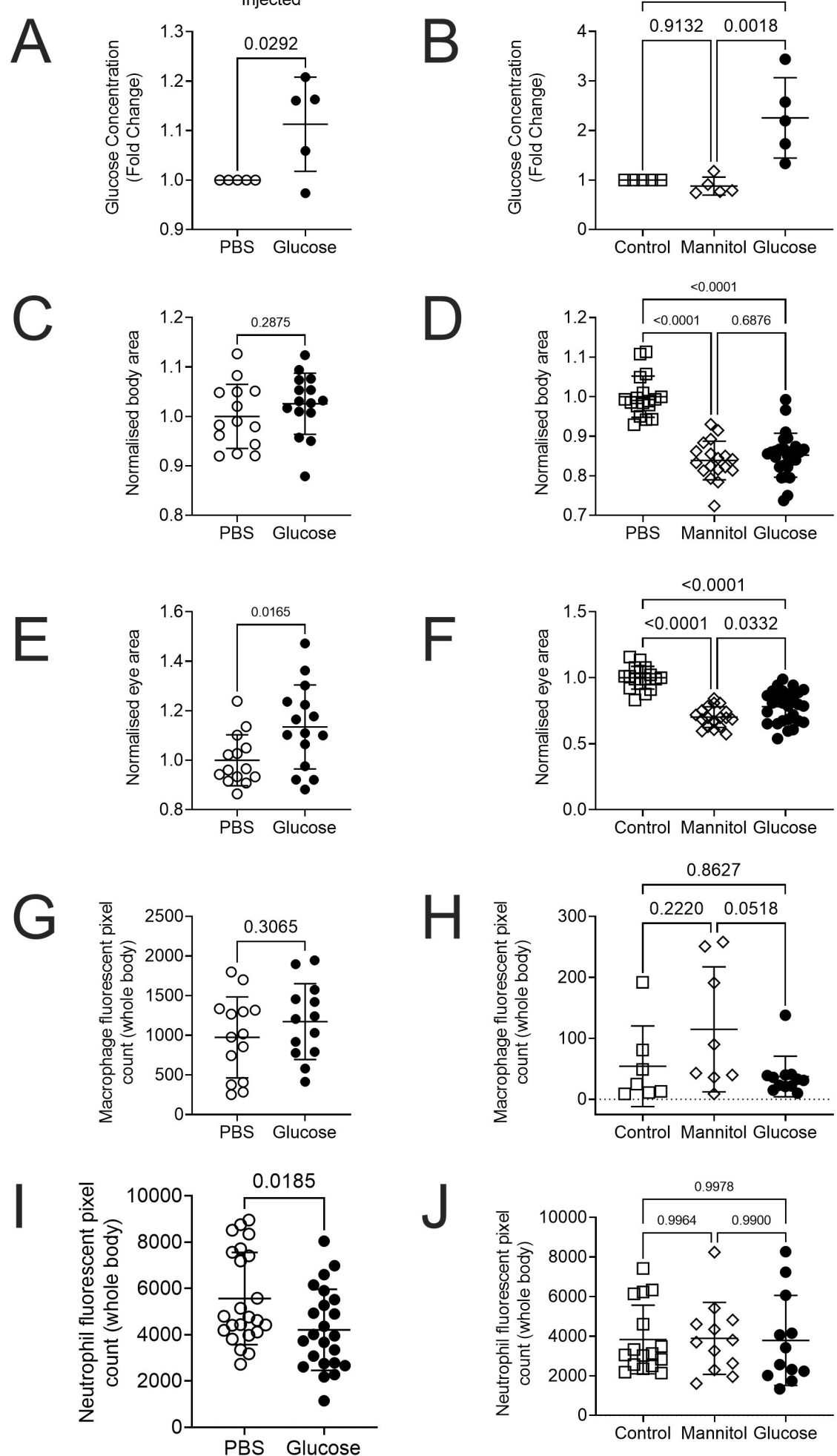
392 B. Bright field images of 6 dpf Oil Red O-stained larvae, demonstrating darker vascular staining in  
393 glucose-injected larvae. Box indicates location of inset, arrowheads indicate stained intersegmental  
394 vessels in inset, asterisk indicates intestinal lumen which was excluded from analysis.

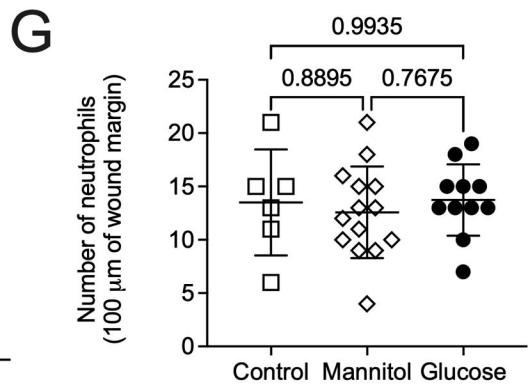
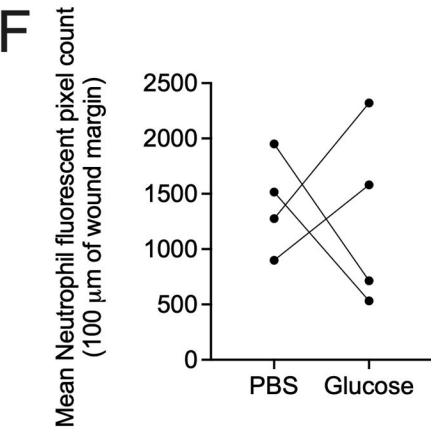
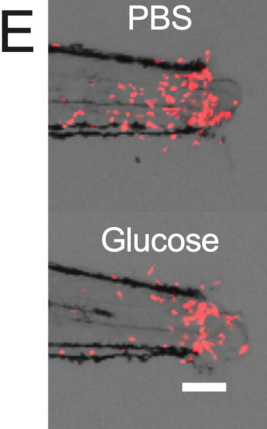
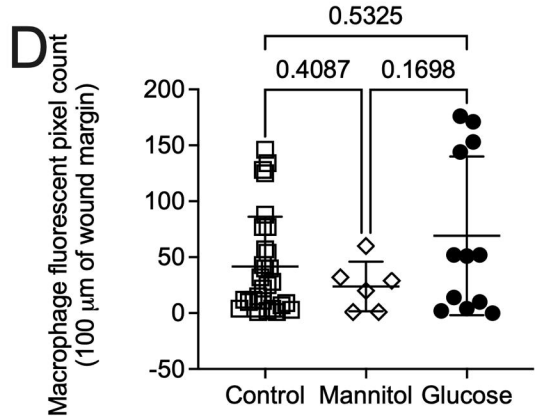
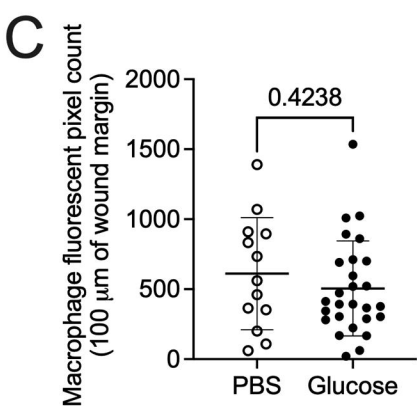
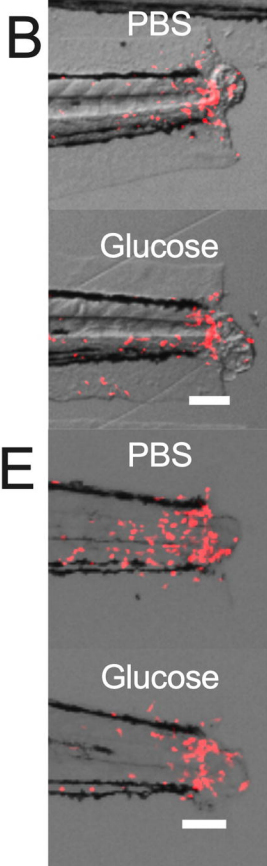
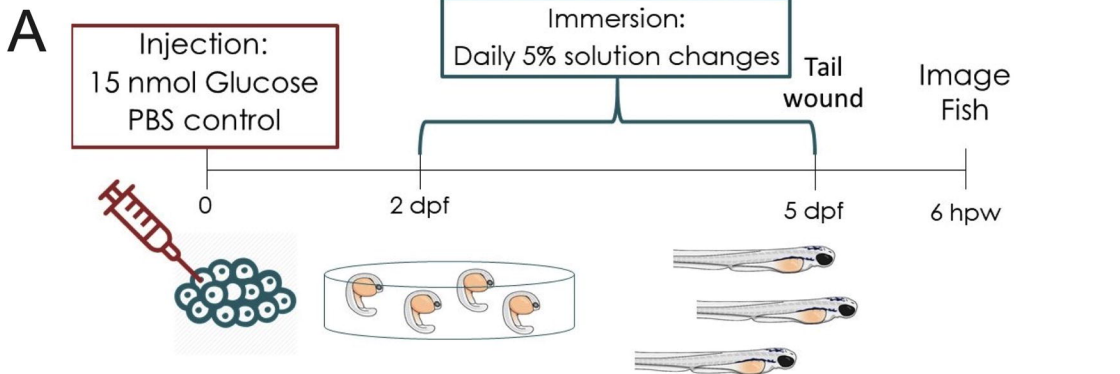
395 C. Quantification of lipid accumulation in 5 dpf glucose-injected larvae.

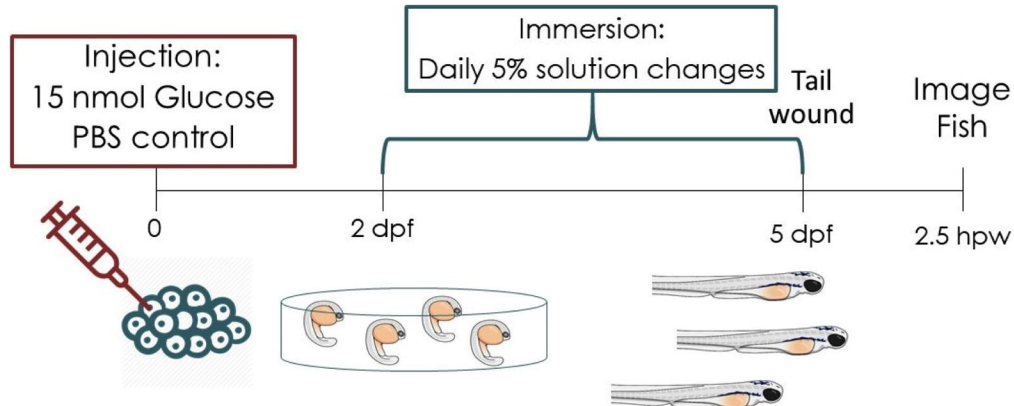
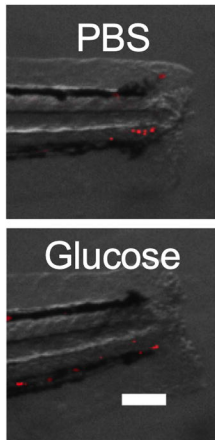
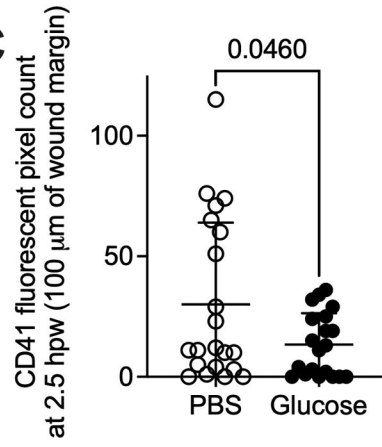
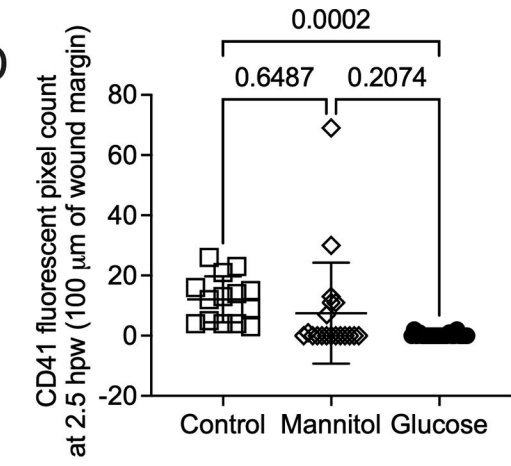
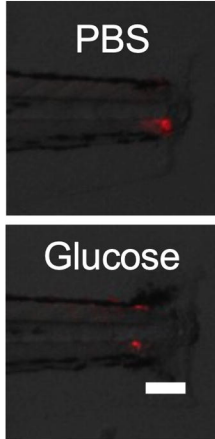
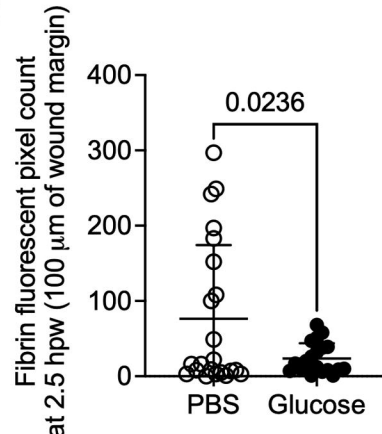
396 D. Quantification of lipid accumulation in 6 dpf glucose-injected larvae challenged with a high fat  
397 diet from 5 dpf.

398 E. Quantification of lipid accumulation in 7 dpf glucose-injected larvae challenged with a high fat  
399 diet from 5 dpf.

400 Statistical testing by t-test. Data are representative of 2 biological replicates.





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