1	Early life glucocorticoid exposure modulates immune function in zebrafish (Danio rerio) larvae
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17 Abstract (number of words: 239)

18 In this study we have assessed the effects of increased cortisol levels during early embryonic 19 development on immune function in zebrafish (Danio rerio) larvae. Fertilized eggs were exposed to 20 either a cortisol-containing, a dexamethasone-containing (to stimulate the glucocorticoid receptor 21 selectively) or a control medium for 6 hours post-fertilisation (0-6 hpf). First, we measured baseline 22 expression of a number of immune-related genes (socs3a, mpeg1.1, mpeg1.2 and irg1l) 5 days post-23 fertilisation (dpf) in larvae of the AB and TL strain to assess the effectiveness of our exposure 24 procedure and potential strain differences. Cortisol and dexamethasone strongly up-regulated 25 baseline expression of these genes independent of strain. The next series of experiments were 26 therefore carried out in larvae of the AB strain only. We measured neutrophil/macrophage recruitment 27 following tail fin amputation (performed at 3 dpf) and phenotypical changes as well as survival 28 following LPS-induced sepsis (150 µg/ml; 4-5 dpf). Dexamethasone, but not cortisol, exposure at 0-6 29 hpf enhanced neutrophil recruitment 4 hours post tail fin amputation. Cortisol and dexamethasone 30 exposure at 0-6 hpf led to a milder phenotype (e.g. less tail fin damage) and enhanced survival 31 following LPS challenge compared to control exposure. Gene-expression analysis showed 32 accompanying differences in transcript abundance of tlr4bb, cxcr4a, myd88, il1ß and il10. These data 33 show that early-life exposure to cortisol, which may be considered to be a model or proxy of maternal 34 stress, induces an adaptive response to immune challenges, which seems mediated via the 35 glucocorticoid receptor.

36 Abbreviations used in the manuscript

- 37 ANOVA: analysis of variance; GR: glucocorticoid receptor; dpf: days post-fertilisation; GFP: green
- 38 fluorescent protein; hpa: hours post-amputation; hpf: hours post-fertilisation; HPI-axis: hypothalamus
- 39 pituitary interrenal axis; IL: interleukine; KMO: Kaiser-Meyer-Olkin; LPS: lipopolysaccharide;
- 40 **MANOVA**: multivariate analysis of variance; **MR**: mineralocorticoid receptor; **ns**: not significant; **PBS**:
- 41 phosphate buffered saline; **qPCR**: quantitative polymerase chain reaction; **SEM**; standard error of the
- 42 mean; TLR: Toll-like receptor; Tukey HSD: Tukey Honestly Significant Difference

43 **1. Introduction**

44 In teleosts, like zebrafsh (Danio rerio), cortisol is the main endogenous corticosteroid, which is 45 secreted when individuals perceive situations as stressful (Wendelaar Bonga, 1997). Like in other 46 vertebrate species, cortisol binds in teleosts to the mineralocorticoid receptor (MR) and the 47 glucocorticoid receptor (GR), which affect the transcription rates of genes following ligand binding 48 (Alderman and Bernier, 2009; Alsop and Vijayan, 2008; Nesan and Vijayan, 2013). Since the GR has 49 a lower affinity for cortisol than the MR, the GR mediates the actions of cortisol during stress, which 50 involves optimizing energy expenditure by tuning the balance between and within physiological 51 systems, like an organism's metabolism and its immune, cardiovascular and central nervous system 52 (e.g. Gorissen and Flik, 2016; Nesan and Vijayan, 2013; Wendelaar Bonga, 1997). Following long-53 term exposure to stress, baseline levels of cortisol are increased, reflecting the allostatic load that the 54 environment imposes on an organism (Gorissen and Flik, 2016; McEwen and Wingfield, 2003). 55 Cortisol has been shown to signal through the GR already during the very early stages of embryonic 56 development; in oocytes maternally deposited cortisol and GR mRNA are present (Pikulkaew et al., 57 2010, 2011; Wilson et al., 2013). These cortisol levels may reflect the allostatic load that the mothers 58 experience in their environment and it has therefore been hypothesized that these deposited cortisol 59 levels are important for preparing the offspring for the expected allostatic load that larvae will 60 encounter in the prevailing environment, thereby programming their cortisol secretion and the 61 functioning of physiological systems to meet expected demands (Nesan and Vijayan, 2013; van den 62 Bos et al., 2019). While in previous studies we focussed on the effects of cortisol exposure (between 0 63 and 6 hours post fertilisation (hpf)) on vigilance-related behaviour, baseline cortisol levels and 64 metabolism in larvae (van den Bos et al., 2019; van den Bos, 2019) in the present study we focussed 65 on the functioning of the immune system, in particular the innate immune system on which zebrafish 66 larvae rely (Meijer and Spaink, 2011). Cortisol exposure may be considered to be a model or proxy of 67 maternal stress (Best et al., 2017; Nesan and Vijayan, 2012, 2016; van den Bos et al., 2019). 68 Zebrafish is a highly suitable animal model to study early life events in the fields of biomedical 69 research, behavioural biology and eco-toxicology (e.g. Champagne et al., 2010; Dai et al., 2013; 70 Nesan and Vijayan, 2013; Steenbergen et al., 2011; Stewart et al., 2014). Fertilized eggs develop into 71 independently feeding larvae outside the mother, without parental care, and can easily be maintained 72 under different experimental conditions as well as pharmacologically manipulated. In zebrafish, it has

73 been demonstrated that mothers deposit cortisol and GR mRNA in oocytes (Pikulkaew et al., 2010, 74 2011; Wilson et al., 2013). These cortisol levels decrease over the first 24 hours post fertilisation, after 75 which zygotes start to produce cortisol by the then developing interrenal cells (Liu, 2007; Pikulkaew et 76 al., 2010, 2011; Wilson et al., 2013). After hatching (48-72 hpf) pituitary control over interrenal cortisol 77 production starts and it takes another 4-5 days before the hypothalamus-pituitary-interrenal (HPI) axis 78 is fully functionally mature (Alderman and Bernier, 2009; Alsop and Vijayan, 2008). Maternal GR 79 mRNA is present during the first 6 hpf, and at 8-9 hpf zygotic expression of the GR commences, while 80 the MR mRNA production starts at 24 hpf (Alsop and Vijayan, 2008; Pikulkaew et al., 2010, 2011).

81 In several studies the effect of cortisol exposure during early embryonic stages has been 82 investigated in zebrafish. In these studies cortisol levels were increased by injection of cortisol into the 83 yolk of one-cell stage embryos (e.g. Best et al., 2017; Nesan and Vijayan, 2012, 2016) or through 84 addition of cortisol to the medium (Hartig et al., 2016; van den Bos et al., 2019). These studies 85 showed that, as a result of the cortisol exposure during early embryonic stages, larval baseline levels 86 of cortisol were increased (Best et al., 2017; Hartig et al., 2016; Nesan and Vijayan, 2012; van den 87 Bos et al., 2019). For example, in a recent study, we have demonstrated that cortisol exposure 88 between 0 and 6 hpf increased baseline cortisol levels 5 days post fertilisation (dpf) and this effect was 89 stronger in larvae from the AB strain than in larvae from the TL strain (van den Bos et al., 2019).

90 Exposure to cortisol (0-5 dpf) has been shown to lead to an enhanced expression of immune-91 related genes in zebrafish larvae at 5 dpf (Hartig et al, 2016), suggesting that early cortisol exposure 92 increases the activity of the immune system. In the present study, we have first measured the 93 expression of a selected number of these up-regulated genes at 5 dpf (socs3a, mpeg1.1, mpeg1.2, 94 irg1/) following 0-6 hpf exposure to cortisol in zebrafish larvae of the AB strain to assess whether our 95 method produces similar effects. Tüpfel long-fin (TL) is another widely used zebrafish strain next to AB 96 and is characterized by spots rather than stripes as well as long fins rather than short fins. In previous 97 studies we have observed consistent differences between larvae of the AB strain and larvae of the TL 98 strain at the level of both HPI-axis activity and behaviour (van den Bos et al., 2017a, 2017b, 2019a, 99 2020). We have attributed these differences to the mutation in the connexin 41.8 gene that leads to 100 spots (for discussion: see van den Bos et al., 2020). Measuring the expression of these genes in 101 larvae of the TL strain next to larvae in the AB strain may therefore reveal how robust our findings are.

102 Finally, to assess the role of the GR in more detail, we exposed fertilized eggs 0-6 hpf to

103 dexamethasone, a specific GR agonist (Rupprecht *et al.*, 1993).

104 To functionally assess the activity of the immune system following early life exposure to 105 cortisol or dexamethasone, we used two experimental models for immune activation. First, we used 106 the tail fin amputation assay. This is a well-established model in which amputation of the tail triggers 107 expression of many pro-inflammatory molecules and the recruitment of innate immune cells 108 (neutrophils and macrophages) towards the wounded area (Hall et al., 2014; Renshaw et al., 2006; 109 Roehl, 2018). The tail fin amputation assay was performed using the double transgenic fish line 110 Tg(mpx:GFP/mpeg1:mCherry-F) (Bernut et al., 2014; Renshaw et al., 2006). Recruitment of 111 neutrophils and macrophages was determined following tail fin amputation in larvae at 3 dpf 112 (Chatzopoulou et al., 2016; Xie et al., 2019). Second, we used a sepsis model, which involved a 113 challenge with lipopolysaccharide (LPS), the membrane component of Gram-negative bacteria, in 4 114 dpf larvae. We measured survival, phenotypical changes, and the expression of a series of LPS-115 responsive genes (Dios et al., 2014; Hsu et al., 2018; Novoa et al., 2009; Philip et al., 2017).

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117 2. Materials and Methods

118 **2.1.** Subjects, spawning and care

Housing conditions and breeding procedures were similar as those reported in van den Bos *et al.* (2017a, 2017b, 2019; AB and Tüpfel long-fin (TL) strains) or Xie *et al.* (2019; the double transgenic fish line *Tg(mpx:GFP/mpeg1:mCherry-F)*. They were kept in recirculation systems (~28°C) under a 14h:10h light-dark cycle and fed twice daily.

Breeding started at least one hour after the last feeding of zebrafish (>16:00 h). Males and females of the AB or TL strain were placed in a zebrafish breeding tank, separated by a partitioning wall, with water of ~28°C. After turning on the lights the next morning, the partitioning wall was removed and tanks were placed at a slight angle, such that the fish had the possibility to move into shallow water to spawn.

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129 **2.2.** Cortisol/dexamethasone exposure during early embryonic development (0-6 hpf)

130 Cortisol (hydrocortisone; Sigma-Aldrich, Zwijndrecht, the Netherlands) and dexamethasone (Sigma-

131 Aldrich, Zwijndrecht, the Netherlands) were dissolved in 96% ethanol in the required stock solution

132 concentrations and stored at -20°C. From these stock solutions media with the appropriate 133 concentration were freshly prepared for each experiment (Althuizen, 2018; van den Bos et al., 2019): 134 cortisol-containing medium: 400 µg/l cortisol (1.1 µM), 0.4 ml/l 96% ethanol (0.04% v/v), 5 mM NaCl, 135 0.17 mM KCI, 0.33 mM CaCl₂, 0.33 mM MgSO₄ in dH2O; dexamethasone-containing medium: 430 136 μ g/l dexamethasone (1.1 μ M), 0.4 ml/l 96% ethanol (or 1 ml/l 96% ethanol (0.1% v/v)); depending on 137 the specific experiment), 5 mM NaCl, 0.17 mM KCl, 0.33 mM CaCl₂, 0.33 mM MgSO₄ in dH2O. 138 Control medium consisted of: 0.4 ml/l 96% ethanol (or 1 ml/l 96% ethanol; depending on the specific 139 experiment), 5 mM NaCl, 0.17 mM KCl, 0.33 mM CaCl₂, 0.33 mM MgSO₄ in dH2O. 140 Directly following spawning and fertilization, eggs were collected and randomly assigned to

Petri dishes filled with either cortisol-containing, dexamethasone-containing or control medium. Within 1-1.5 hpf Petri dishes were placed in an incubator set at 28.5 °C (300–350 lux). Eggs were exposed to these solutions for 6 hrs. It has been shown that both cortisol and dexamethasone diffuse inside the eggs in this period (Steenbergen *et al.*, 2017). In addition, we have shown this procedure to be effective in eliciting changes in physiology and behaviour at 5 dpf (Althuizen, 2018; van den Bos *et al.*, 2019).

Following this, cortisol-containing, dexamethasone-containing and control media were replaced by E3 medium (5 mM NaCl, 0.17 mM KCl, 0.33 mM CaCl₂, 0.33 mM MgSO₄, 3 ml/l 0.01% (w/v) methylene blue in dH2O). Embryos were rinsed twice in E3 medium to ensure that the original treatment media were completely removed. Petri dishes were returned to the incubator allowing the embryos to develop further (28.5°C; 14h:10h light-dark period (lights on: 09.00h – 23.00h); light phase: 300–350 lux; dark phase; 0 lux). At 1 and 4 dpf E3 medium was refreshed and unfertilized eggs, dead eggs/embryos/larvae and chorions were removed from the dishes.

154 All experiments were carried out in accordance with the Dutch Experiments on Animals Act 155 (http://wetten.overheid.nl/BWBR0003081/2014-12-18), the European guidelines for animal 156 experiments (Directive 2010/63/EU; http://eur-lex.europa.eu/legal-content/NL/TXT/HTML/?uri= 157 CELEX:32010L0063) and institutional regulations (Radboud University or Leiden University). Larvae 158 were euthanized by placing them in ice slurry for at least 20 minutes followed by adding bleach to the 159 slurry. In case of anaesthesia 0.01% 2-phenoxyethanol (Sigma Aldrich, Zwijndrecht, the Netherlands; 160 experiments Radboud University) or 0.02% buffered aminobenzoic acid ethyl ester (tricaine; Sigma 161 Aldrich, Zwijndrecht, the Netherlands; experiments Leiden University) was used.

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163 **2.3 Baseline gene expression analysis.**

164 Following 0-6 hpf exposure to the different media, larvae of the AB and TL strain were sampled for 165 gene expression at 5 dpf between 16:00 hrs and 19:00 hrs (van den Bos et al., 2019; experimental 166 time-line: figure 1). Genes of interest were: socs3a, mpeg1.1, mpeg1.2 and irg1l (for primer 167 sequences of these genes: see van den Bos et al., 2017a). Gene expression was determined by 168 qPCR analysis as described below. 169 170 2.4 Tail fin amputation assay 171 The tail fin amputation assay was performed using the double transgenic fish line 172 Tg(mpx:GFP/mpeg1:mCherry-F; Bernut et al., 2014; Renshaw et al., 2006). Three-day-old zebrafish 173 larvae were anaesthetized in E3 medium containing 0.02% buffered tricaine and loaded onto 2% 174 agarose-coated Petri dishes (experimental time-line: figure 1). Amputation was performed with a 1 mm 175 sapphire blade (World Precision Instruments, Friedberg, Germany) using a Leica M165C 176 stereomicroscope and a micromanipulator (Leica Microsystems BV, Amsterdam, the Netherlands; see 177 below, figure 3A). Larvae were fixed in 4% paraformaldehyde in PBS at 4 hours post amputation (hpa) 178 and stored overnight at 4°C. The following day larvae were washed twice for one minute and then 179 twice for five minutes in PBS containing 0.01% Tween 20 (Sigma Aldrich, Zwijndrecht, the 180 Netherlands). 181 A Leica M205FA fluorescence stereomicroscope supported by LAS software (version 4.12.0; 182 Leica Microsystems BV, Amsterdam, the Netherlands) was utilised to visualise the leukocytes. 183 Detection of neutrophils and macrophages was based on their fluorescent GFP and mCherry signals 184 respectively. To quantify cell migration towards the wounded area, cells within a distance of 200µm 185 from the amputation site were counted manually, as previously described (Xie et al., 2019). Data 186 (numbers of migrated neutrophils and macrophages per individual) were pooled from three individual 187 experiments (n>10 per experiment), and the presented data are means (± SEM). 188

2.5. LPS exposure

First, we conducted a pilot study to assess the optimal LPS dose, exposure duration and parameters
to be measured (protocols adapted from: Dios *et al.*, 2014; Hsu *et al.*, 2018; Novoa *et al.*, 2009; Philip

192 *et al.*, 2017). Incubation for 30 minutes in 150 μ g/ml LPS (B11.04; Sigma Aldrich, Zwijndrecht, the 193 Netherlands) was effective in eliciting a robust increase in *il1* β expression (assessed using qPCR 194 analysis), changes in tail fin morphology (swollen or damaged tails) and increased levels of reactive 195 oxygen species (ROS; measured by a fluorescent labelling method according to Philip *et al.*, 2017). 196 Hence, we used this dose in subsequent experiments.

197 Subsequently, two experiments were conducted. In both experiments 4 dpf larvae were 198 exposed to 150 µg/ml LPS (B11.04; Sigma Aldrich, Zwijndrecht, the Netherlands) in E2 medium or 199 control E2 medium for 30 minutes in Petri dishes (n=50 in 25 ml; experimental time-line: figure 1). 200 Following this exposure. LPS-containing medium or control E2 medium was replaced by fresh E2 201 medium (larvae were rinsed two times to ensure that the original media were removed). Larvae either 202 remained in the Petri dishes for sampling for gene expression at later time points or were transferred 203 individually to 24 wells plates (Greiner Bio-One BV, Alphen a/d Rijn, the Netherlands) for assessing 204 phenotypical changes and survival (volume per well: 1-1.5 ml). Six treatment groups were thus 205 created: 0-6 hpf control, cortisol or dexamethasone treatment, combined with either 4 dpf LPS or 206 control treatment (for 30 min).

In the first experiment the level of gene expression at 0 hr, 0.5 hr (i.e. directly following
exposure), 1 hr (i.e. 30 minutes after ending exposure) and 3 hrs (i.e. 2.5 hrs after ending exposure;
Novoa *et al.*, 2009) was determined by qPCR analysis as described below. Genes of interest (see
papers by: Hsu *et al.*, 2018; Kanwal *et al.*, 2013; Novoa *et al.*, 2009; Philip *et al.*, 2017) were genes
encoding proteins involved in barrier function of the vascular endothelium (*cldn5a*, *cldn2*, *oclnb*), Tolllike receptors (*tlr2*, *tlr4ba*, *tlr4bb*, *tlr5a*, *tlr5b*), and regulators of the immune response (*il1β*, *il10*, *myd88*, *cxcr4a*, *cxcr4b*, *ptpn6*). Primer sequences are listed in Table 1.

214 In addition, in the first experiment survival and phenotypical changes were determined at 0, 4 215 (i.e. 3.5 hrs after ending exposure) and 28 hrs (i.e. 27.5 hrs after ending exposure). Phenotypical 216 changes included (see Philip et al., 2017): changes in tail fin morphology (normal, swollen (oedema) 217 or damaged), presence of heart oedema and changes in shape (straight or curved). In the second 218 experiment (that also served as replicate for the first experiment) survival and phenotypical changes 219 were measured at 0, 0.5, 1, 3, 6 and 24 hrs (i.e. before exposure, directly after exposure, 0.5 hrs after 220 ending exposure, 2.5 hrs after ending exposure, 5.5 hrs after ending exposure and 23.5 hrs after 221 ending exposure).

223 **2.6 Gene expression analysis**

For the assessment of gene expression levels by qPCR analysis, 3-5 larvae were transferred to a 2-ml Eppendorf tube; thus, one sample contained material from 3-5 larvae. Residual medium was removed with a pipette, tubes were snap frozen in liquid nitrogen, kept on ice during the sampling procedure, and subsequently stored at -80 °C until total RNA extraction.

228 RNA isolation, RNA preparation, removal of genomic DNA from the samples and synthesis of 229 cDNA was performed according to the protocol described in van den Bos et al. (2017a). Total RNA 230 content of each sample was isolated. This was done by homogenising the tissue with 400 µl Trizol 231 reagent (Invitrogen, Carlsbad, USA) in a Grinding Mill (Retsch GmbH, Germany) for 20 s at 20 Hz. 232 After homogenisation, samples were kept at room temperature for 5 min. Next, 80 µl chloroform was 233 added and the solution was mixed by shaking for 15 s. Afterwards, samples were kept at room 234 temperature for 2 min. The samples were centrifuged at 14,000 rpm for 10 min in a cooled centrifuge 235 (4°C) and the aqueous phase of the samples was transferred to a new tube. To this phase, 200 µl 236 isopropanol was added and this solution was mixed well by inversion of the tube. The solution was 237 then stored at -20°C for 2 h and centrifuged afterwards for 15 min at 14,000 rpm in a cooled centrifuge 238 (4°C). The supernatant was decanted and the pellet washed with 500 µl 75% ethanol and centrifuged 239 10 min at 14,000 rpm in a cooled centrifuge (4°C). The supernatant was decanted, after which the 240 pellet was centrifuged for 5 s to remove all the remaining supernatant using a pipette. The pellet 241 containing the RNA was air-dried for 10 min at room temperature and afterwards dissolved in 100 µl 242 ice cold DEPC-treated dH2O. To this RNA solution, 10 µl 3M NaAc (pH 5.4) and 250 µl 100% ethanol 243 were added. The solution was mixed by inverting the tube and samples were stored for 2 h at -20°C. 244 Subsequently, the samples were centrifuged for 15 min at 14,000 rpm in a cooled centrifuge (4°C), 245 and the supernatant was decanted and the pellet washed washed as described earlier. Finally, the 246 RNA pellet was dissolved in 15 µl DEPC-treated dH2O. The concentration and quality of RNA in each 247 sample were measured using a nanodrop spectrophotometer at 260 nm wavelength (Nanodrop, 248 Wilmington, DE, USA). 249 Isolated RNA was treated with DNase to remove any (genomic) DNA from the sample; 400 ng

250 RNA was transferred into a PCR strip, and DEPC-treated dH2O was added to a volume of

251 8 μl. To this, 2 μl of DNase mix was added, containing 1 μl 10x DNase I reaction buffer and

1 μl (1 U/μl) amplification grade DNase I (both from Invitrogen, Carlsbad, USA). The resulting mix was
 incubated for 15 min at room temperature. Afterwards, 1 μl 25 mM EDTA was added to stop the
 DNase reaction and the reaction mix was incubated for 10 min at 65°C and returned on ice.

After the DNase treatment, samples were used to synthesize cDNA by the addition of 1 µl random primers (250 ng/µl), 1 µl 10 mM dNTP mix, 4 µl 5 x 1st strand buffer, 1 µl 0.1M DTT, 1 µl RNase inhibitor (10 U/µl), 0.5 µl Superscript II (reverse transcriptase) (200 U/µl) (all from Invitrogen, Carlsbad, USA) and 0.5 µl DEPC-treated dH2O. The resulting mix was incubated for 10 min at 25°C for annealing of the primers and then 50 min at 42°C for reverse transcription. Enzymes were hereafter inactivated by incubating samples at 70°C for 15 min. Finally, 80 µl dH2O was added to dilute the samples five times for the qPCR reaction.

262To measure the relative gene expression in each sample, real-time qPCR was carried out for263each gene of interest. For each qPCR reaction, 16 μl PCR mix (containing 10 μl SYBR green mix (2x)264(BioRad, Hercules, USA), 0.6 μl forward and reverse gene-specific primer (10 μM) and 4.8 μl H2O)265was added to 4 μl of cDNA. The qPCR reaction (3 min 95°C, 40 cycles of 15 s 95°C and 1 min 60°C)266was carried out using a CFX 96 (BioRad, Hercules, USA) qPCR machine. Analysis of the data was267carried out using a normalisation index of two reference genes (viz. elongation factor alpha (elf1a) and268ribosomal protein L13 (rpl13)) (Vandesompele et al., 2002).

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270 **2.7. Statistics**

271 For gene expression analyses, outliers were removed following Grubb's outlier test (p≤0.01). We 272 explored the interrelationships of transcript abundance levels using Principal Component Analysis 273 (PCA) with orthogonal rotation (Varimax rotation with Kaiser normalization; see van den Bos et al., 274 2017b). In case of missing samples, data were excluded list-wise. The number of retained 275 components was based on eigenvalues (>1) and visual inspection of the scree plot. The Kaiser-276 Meyer-Olkin (KMO) measure of sampling adequacy and Bartlett's test of sphericity were done to 277 ensure that data obeyed analysis criteria; both are measures to assess whether the correlation matrix 278 is suited for factor analysis (Budaev, 2010). Component scores were saved and used for further 279 statistical analysis. The following component loading cut-off points were considered: ≤-0.600 or 280 ≥0.600 (Ferguson, 1989; Budaev, 2010).

281	For the basal gene expression values a two-way or three-way Analysis of Variance (ANOVA)
282	was run with treatment, strain or batch (where applicable) as independent factors. In the tail fin
283	amputation assay a Student's t-test was run on the number of neutrophils or macrophages at 4 hpa
284	comparing 0-6 hpf treatment groups (cortisol versus control; dexamethasone versus control).
285	In the LPS exposure experiment, for gene expression a multivariate analysis of variance (MANOVA)
286	was run (to account for multiple comparisons) followed by univariate analysis of variance (0-6 hpf
287	treatment and time as independent factors followed by post-hoc testing (Tukey HSD)). In addition per
288	time point a one-way ANOVA was run with 0-6 hpf treatment as factor followed by post-hoc testing
289	(Tukey HSD).
290	In the LPS exposure experiment differences in survival rate were assessed using the Kaplan-
291	Meijer procedure (Log Rank Mantel-Cox). Differences in phenotypical changes were compared using
292	Chi-square tests.
293	Significance was set at p≤0.05 and trends are indicated (p≤0.10) where appropriate; ns: not
294	significant: p>0.10. Unless otherwise stated all p-values are two-tailed. All statistical analyses were

done using IBM SPSS version 23 (IBM, Armonk, NY, USA).

296

297 3 Results

298 **3.1 Baseline gene expression analysis**

Rather than analysing transcript abundance of different genes following glucocorticoid treatment
 (cortisol or dexamethasone, 0-6 hpf) separately, we explored the effects of exposure through the
 interrelationships of transcript abundance of genes using PCA.

302 A PCA for the 0-6 hpf cortisol treatment experiment revealed two components (figure 2A; 303 supplementary figure 1A shows the transcript abundance of individual genes). The KMO value was 304 sufficiently high (0.522) and Bartlett's test of sphericity was highly significant (Chi-square=17.707, 305 df=6, p≤0.007), indicating that the data were adequate for a PCA. The first component (explaining 306 44.0% of the variance) was comprised of irg11 (loading: 0.854) and socs3a (loading: 0.890); the 307 second component (explaining 28.9% of the variance) of mpeg1.1 (loading: 0.890) and mpeg1.2 308 (loading: 0.890). Each component was analysed separately using ANOVA. Treatment (0-6 hpf) with 309 cortisol enhanced transcript abundance of genes in component 1 (irg1l and socs3a) independent of 310 strain (three-way ANOVA (strain, treatment, batch): treatment: F(1,23)=6.477, p≤0.018). Transcript

abundance was overall higher in TL than AB larvae (strain: F(1,23)=7.023, p≤0.014). Treatment (0-6

hpf) with cortisol enhanced transcript abundance of genes in component 2 (*mpeg1.1* and *mpeg1.2*)

313 independent of strain (three-way ANOVA (strain, treatment, batch): treatment: F(1,23)=7.660,

314 p≤0.011).

A PCA for the 0-6 hpf dexamethasone treatment experiment revealed only one component (figure 2B; supplementary figure 1B shows the transcript abundance of individual genes). The KMO was sufficiently high (0.626) and Bartlett's test of sphericity was highly significant (Chisquare=102.945, df=6, p<0.001), indicating that the data were adequate for a PCA. This component explained 76.6% of the variance. Loadings onto this component were: *irg11* (0.867). *socs3a* (0.835),

320 *mpeg1.1* (0.928) and *mpeg1.2* (0.868). Treatment with dexamethasone enhanced transcript

321 abundance of genes independent of strain (two-way ANOVA (strain, treatment): treatment:

- 322 F(1,33)=6.745, p≤0.014).
- These data show that glucocorticoid treatment was effective in eliciting changes in baseline
 expression of (a selected set of) immune-related genes.
- 325

326 **3.2 Tail fin amputation assay**

327 Figure 3A shows the site of the tail fin amputation performed at 3 dpf. Neutrophil recruitment at 4 hpa

328 was enhanced in the 0-6 hpf dexamethasone treatment group (Student's t-test: t=2.917, df=106,

p≤0.004; figure 3C), but not in the 0-6 hpf cortisol treatment group (Student's t-test: t=0.621, df=89, ns;

figure 3B), compared to the group treated with control medium at 0-6 hpf. No effects were found for

331 macrophage recruitment for either 0-6 hpf cortisol treatment (Student's t-test: t=0.923, df=89, ns;

figure 3B) or 0-6 hpf dexamethasone treatment (Student's t-test: t=-1.095, df=106, ns; figure 3C).

333

334 **3.3 LPS exposure**

335 Survival

In the LPS exposure experiments, we never observed any morphological changes or mortality in the 4 dpf control groups. Hence, we only present the data of the 4 dpf LPS-treated groups. Figure 3 shows the survival data of both LPS exposure experiments. Only 27.8% of the 0-6 hpf control-treated larvae survived in the first LPS exposure experiment (figure 4A), while 79.2% survived in the second LPS

340 exposure experiment (figure 4B). In both LPS exposure experiments the number of larvae that

341 survived following glucocorticoid treatment 0-6 hpf appeared to be higher (Log Rank Mantel Cox; first 342 LPS exposure experiment: overall Chi-square=20.863, df=2, p<0.001; second LPS exposure 343 experiment: overall Chi-square=5.824, df=2, $p \le 0.054$). Pair-wise comparison revealed that (1) in both 344 LPS exposure experiments survival was significantly higher in 0-6 hpf cortisol-treated than 0-6 hpf 345 control-treated larvae ((Log Rank Mantel Cox; Chi-square=14.385, df=1, p<0.001; Chi-square=5.466, 346 df=1, p \leq 0.019), while (2) survival was significantly higher in the first LPS exposure experiment (Log 347 Rank Mantel Cox; Chi-square=12.436, df=1, p<0.001), but not the second LPS exposure experiment 348 (Log Rank Mantel Cox; Chi-square=1.370, df=1, ns) in 0-6 hpf dexamethasone-treated larvae 349 compared to 0-6 hpf control-treated larvae. 350 351 Phenotypical effects and gene expression endothelium-related genes 352 In the first LPS exposure experiment phenotypical changes were measured at 4 and 28 hrs, i.e. 3.5 353 hrs and 27.5 hrs after ending exposure. Table 2 shows the numbers of dead larvae and of larvae that 354 were alive and displayed phenotypical changes. We observed LPS-induced changes in the shape of 355 the larvae (curved larvae), tail fin morphology (swollen or damaged tail fins) and heart cavity 356 (oedema). Larvae were scored as either affected (at least one of these changes present) or not (no 357 changes in any of the parameters). Supplementary table 1 shows the scores of the individual

358 parameters at 4 hrs (the number of larvae of the control treated group alive at 28 hrs was too low for a

359 meaningful statistical analysis between treatments).

While the scores at 4 hrs suggested that the LPS-induced effects were less strong in glucocorticoid-treated larvae than in control-treated larvae, this was not (as yet) significant (Chisquare=5.81, df=2, ns). LPS-induced effects were less strong in glucocorticoid-treated larvae than in control-treated larvae at 28 hrs (overall Chi-square=25.33, df=2, p<0.001). Pair-wise comparison showed that both cortisol-treated (Chi-square=18.05, df=2, p<0.001) and dexamethasone-treated (Chi-square=18.34, df=2, p<0.001) groups showed fewer dead and fewer malformed larvae than control-treated subjects following LPS exposure.

In addition to the phenotypical changes that we studied, we measured expression levels of genes related to endothelial barrier function (*clnd5a*, *clnd2* and *oclnb*). Transcript abundance of *clnd5a* was higher in cortisol-treated (Tukey HSD: p<0.001) and dexamethasone-treated larvae (Tukey HSD: $p\leq 0.067$) compared to control-treated larvae (two-way ANOVA; treatment and time as independent

371 factor; treatment: F(2,36)=9.815, p<0.001; figure 5A). Transcript abundance of *clnd2* was lower in 372 cortisol-treated (Tukey HSD: p≤0.034) and dexamethasone-treated larvae (Tukey HSD: p≤0.093) 373 compared to control-treated larvae (two-way ANOVA; treatment and time as independent factor; 374 treatment: F(2,36)=9.815, p<0.001; figure 5B). At 3 hrs transcript abundance of *clnd2* was higher than 375 at other time points in all treatment groups (Tukey HSD; p<0.001; time: F(2,36)=13.425, p<0.001). 376 Transcript abundance of oclnb was lower in cortisol-treated and dexamethasone-treated larvae than in 377 control-treated larvae at baseline (0 hr), but higher at 3 hrs following LPS exposure (two-way ANOVA; 378 treatment and time as independent factor; treatment *time: F(3,36)=6.404, p<0.001; figure 5C).

379 In the second LPS exposure experiment we measured changes in tail fin morphology and 380 shape as we hardly observed any oedema in the heart cavity in the first LPS exposure experiment. 381 Figure 6 shows the changes in tail fin morphology following LPS exposure. In control-treated larvae 382 there was a clear and rapid loss of normal tail fin structure and a shift towards swollen or damaged tail 383 fins, while this was not the case in the glucocorticoid-treated larvae. Statistical analysis showed that at 384 3 hrs (Chi-square=10.25, df=4, p≤0.036), 6 hrs (Chi-square=10.27, df=4, p≤0.036) and 24 hrs (Chi-385 square=10.25, df=4, p≤0.042) treatment groups differed significantly from one another. More in 386 particular, both cortisol-treated larvae (3 hrs: Chi-square=5.16, df=2, p≤0.076; 6 hrs: Chi-square=6.94, 387 df=2, p≤0.03; 24 hrs Chi-square=5.77, df=2, p≤0.056) and dexamethasone-treated larvae (6 hrs: Chi-388 square=6.98, df=2, p≤0.03; 24 hrs Chi-square=6.53, df=2, p≤0.038) showed less severe changes in 389 tail fin morphology than control-treated larvae. We did not observe a strong effect of LPS exposure on 390 the shape of the larvae. After 24 hrs only small percentages of each treatment group showed a curved 391 shape: control-treated larvae (see supplementary table 2 for all time points): 15.8% (n=19); cortisol-392 treated larvae: 8.3% (n=24); dexamethasone-treated larvae: 9.1% (n=22); these differences were not 393 significant (Chi-square=1.71, df=2, ns).

394

395 Gene expression analysis

First, we explored the effects of glucocorticoid treatment through the interrelationships of transcript abundance of genes using PCA. Then in each component, we selected genes of interest of which the transcript abundance of the different treatments matched the differences in phenotypical changes and survival that we observed between treatment groups.

- A PCA revealed three components explaining in total 72.9% of variance (supplementary table
 3). The KMO was sufficiently high (0.619) and Bartlett's test of sphericity was highly significant (Chisquare=303.010, df=55, p<0.001) indicating that the data were adequate for a PCA.
- 403 The first component, explaining 32.9% of the variance, consisted of the genes of different Toll-404 like receptors: *tlr2* (factor loading: 0.677), *tlr4ba* (0.669), *tlr4bb* (0.764), *tlr5a* (0.779) and *tlr5b* (0.838). 405 The overall pattern of the factor regression scores across time was an inverted U-shape: a two-way 406 ANOVA (independent factors: time and treatment) for this component revealed a highly significant 407 effect of time (F(3,36)=15.502, p<0.001) with time points 0 hr and 3 hrs having significantly higher 408 factor regression scores than time points 0.5 hr and 1 hr (Tukey HSD; supplementary Table 3). Only 409 weak effects were found between the different treatments (F(2,36)=3.012, $p\leq 0.062$; F(6,36)=2.038, 410 $p \le 0.086$). The only gene of this component of which the transcript abundance of the different 411 treatments seemed to match the differences in phenotypical changes and survival was tlr4bb (see 412 supplementary table 4 and figure 7A): transcript abundance increased less strongly over time in 413 glucocorticoid-medium treated subjects than in control-medium treated subjects.
- 414 The second component, explaining 23.1% of the variance, consisted of $il1\beta$ (factor loading: 415 0.940), myd88 (0.844) and il10 (0.758). As time progressed factor regression scores increased (two-416 way ANOVA (independent factors: treatment and time); time: F(3,36)=68.309, p<0.001) with a post-417 hoc Tukey HSD revealing that the factor regression scores differed significantly from one another at all 418 time points (supplementary table 3). The 0-6 hpf control group had increasingly higher factor 419 regression scores than the 0-6 hpf cortisol treated group and the 0-6 hpf dexamethasone treated 420 group as time progressed (supplementary table 3; Tukey HSD; treatment: F(2,63)=22.664, p<0.001; 421 treatment*time: F(6,36)=13.491, p<0.001). Of all genes of this component, transcript abundance of the 422 different treatments seemed to match the differences in phenotypical changes and survival 423 (supplementary table 4 and figure 7 B-D): transcript abundance increased less strongly over time in 424 glucocorticoid-medium treated subjects than in control-medium treated subjects. 425 The third component, explaining 16.9% of the variance, consisted of regulators of the immune 426 response: cxcr4a (factor loading: 0.794), cxcr4b (0.809) and ptpn6 (0.799). Overall the factor 427 regression scores decreased over time: a two-way ANOVA (independent factors: treatment and time) 428 revealed a highly significant effect of time (F(3,36)=4.464, $p \le 0.009$) with 3 hrs and 0 hr being 429 significantly different from one another (Tukey HSD). The 0-6 hpf dexamethasone group had lower

- 430 factor regression scores than the 0-6 hpf control treated group and the 0-6 hpf cortisol treated group
- 431 (treatment: F(2,36)=11.056, p<0.001; treatment*time: F(6,36)=5.571, p<0.001). The only gene of this
- 432 component of which the transcript abundance of the different treatments seemed to match the
- 433 differences in phenotypical changes and survival was *cxcr4a* (see supplementary table 4 and figure
- 434 7E): transcript abundance increased in control-medium treated subjects but decreased in
- 435 glucocorticoid-medium treated subjects.
- 436

437 **4. Discussion**

The data of this study showed that treatment of zebrafish embryos with cortisol or dexamethasone during the first six hours after fertilization modulated the function of the immune system and thereby enhanced survival after an immune challenge. This suggests that in zebrafish maternal stress through enhancing oocyte cortisol levels and thereby increased GR stimulation leads to an adaptive response to immune challenges.

443

444 **4.1. Baseline expression of immune-related genes**

445 Hartig and colleagues (2016) have shown that 5-day exposure to (1 µM) cortisol in zebrafish 446 embryos/larvae (0-5 dpf) enhanced baseline expression of immune-related genes such as of socs3a, 447 mpeg1 and irg1l at 5 dpf. Here, we show that exposure at the first six hours of life (0-6 hpf) is already 448 sufficient to induce enhanced baseline expression of these genes at 5 dpf. As GR is the only 449 corticosteroid receptor present in these early life stages (Alsop and Vijayan, 2008; Pikulkaew et al., 450 2010, 2011), this suggests that activation of GR in these early stages is responsible for mediating 451 these effects. Indeed, we show that 0-6 hpf exposure to the specific GR agonist dexamethasone 452 enhanced baseline expression of these immune-related genes in 5 dpf larvae as well. This is in 453 general agreement with data from studies showing that exposure of embryos/larvae to other GR 454 agonists (for variable time-periods from fertilisation) increased baseline expression of (some of) these 455 genes (Willi et al., 2018, 2019; Zhao et al., 2016).

We have previously observed that 0-6 hpf exposure to cortisol enhanced baseline levels of cortisol at 5 dpf in larvae of the AB strain but not in larvae of the TL strain (van den Bos *et al.*, 2019). In addition, we have observed that 0-6 hpf exposure to dexamethasone had no effect on baseline levels of cortisol at 5 dpf in larvae of the AB or TL strain (Althuizen, 2018). Overall these data suggest that the effects on the expression of immune-related genes in AB and TL larvae are independent ofbaseline levels of cortisol.

While we have previously observed substantial differences between larvae of the AB and TL strains in gene expression, physiology and behaviour (van den Bos *et al.*, 2017a, 2017b, 2019, 2020), the effects of 0-6 hpf exposure to cortisol and dexamethasone on baseline expression of immunerelated genes were similar in both strains. This suggests a robust strain-independent effect. Future studies should elucidate whether this effect also holds in other strains revealing thereby fundamental aspects of how early-life levels of cortisol may affect offspring functioning and survival.

468

469 **4.2 Tail fin amputation assay**

It has been shown that tail fin amputation in 3 dpf larvae leads to a rapid recruitment of macrophages (within 2 hpa) remaining at a plateau for at least 24 hours thereafter, while the number of neutrophils reaches a peak 4 hpa declining thereafter (Chatzopoulou *et al.*, 2016). Here, we observed that dexamethasone increased the number of neutrophils at 4 hpa, while it had no effect on the number of macrophages at 4 hpa. These data suggest that pre-exposed subjects mount a stronger response to injury, possibly enhancing protection against invading micro-organisms (Hall *et al.*, 2014).

476 Dexamethasone, but not cortisol, exposure was effective in increasing the recruitment of 477 neutrophils. Two explanations may be forwarded for this. First, this may be related to 478 dexamethasone's higher affinity for GR (Rupprecht et al., 1993). In line with this we observed 479 enhanced expression of FK506 binding protein 5 (*fkbp5*), a marker for levels of GR stimulation (e.g. 480 Willi et al., 2018, 2019) in dexamethasone-exposed embryos, but not cortisol-exposed embryos 481 (Althuizen, 2018; van den Bos et al., 2019). Second, it has been shown neutrophil recruitment, but not 482 macrophage recruitment, is sensitive to acute treatment with GR agonists, including cortisol, which 483 decreases the number of neutrophils at 4 hpa (Chatzopoulou et al., 2016; Hall et al., 2014; Xie et al., 484 2019). We have observed that baseline levels of cortisol are enhanced following cortisol but not 485 dexamethasone treatment (Althuizen, 2018; van den Bos et al., 2019). Hence, in the case of cortisol 486 pre-exposure enhancement of neutrophil recruitment through GR stimulation (as indicated by the 487 effect of dexamethasone) may potentially be offset at 3 dpf by the higher baseline levels of cortisol. If 488 so, this suggests a fine-tuning of the response. Future studies should address these alternative 489 explanations.

491 **4.3 LPS challenge**

492 In line with results from other studies (Dios et al., 2014; Hsu et al., 2018; Novoa et al., 2009; Philip et 493 al., 2017) LPS exposure produced phenotypical changes (tail fin oedema, swollen or damaged tail 494 fins, and curved animals), mortality and increased expression levels of immune-related genes. To 495 assess the molecular mechanisms underlying the effects of LPS exposure, we measured the 496 expression of *cldn5a*, *cldn2* and *oclnb*; genes of which the expression has been shown to be changed 497 following LPS exposure (Hsu et al., 2018; Philip et al., 2017) and which are involved in endothelial 498 barrier function (Kása et al., 2015; Odenwald and Turner, 2013; Shen et al., 2011; Yoseph et al., 499 2016). The strong increase in the expression of *cldn2* at 3 hours in control-treated subjects is in line 500 with data from other studies (Hsu et al., 2018; Philip et al., 2017). Increased expression of cldn2 is 501 associated with endothelial hyper-permeability due to increased pore-pathway activity possibly 502 mediated by increased expression of IL-13 (Kása et al., 2015; Odenwald and Turner, 2013; Shen et 503 al., 2011; Yoseph et al., 2016). At variance with other studies (Hsu et al., 2018; Philip et al., 2017) we 504 did not observe strongly decreased expressions of cldn5a and oclnb. Decreased expression levels of 505 cldn5a and oclnb are related to endothelial hyper-permeability due a lower sealing function of the 506 pore-pathway and a less functional leaky pathway respectively (Kása et al., 2015; Odenwald and 507 Turner, 2013; Shen et al., 2011; Yoseph et al., 2016). One reason for this may be that we measured 508 gene expressions at 3 hours after the start of the LPS exposure, while in other studies this was 509 measured at substantially later time points, i.e. 6 and 8 hours (Hsu et al., 2018; Philip et al., 2017). 510 Both cortisol and dexamethasone exposure at 0-6 hpf were associated with milder effects to 511 LPS exposure (in both experiments), as indicated by milder phenotypical changes (lower number of 512 larvae expressing tail fin oedema, damaged tail fins or curved animals), lower mortality and lower 513 gene expression of immune-related genes, such as $ill\beta$. As to the expression of endothelium-related 514 genes we noted that the expression of cldn5a was higher and expression of cldn2 lower in cortisol-515 and dexamethasone-treated subjects compared to control-treated subjects. This suggests a lower

permeability of the endothelium due to a lower pore-pathway activity (Kása *et al.*, 2015; Odenwald and

517 Turner, 2013; Shen *et al.*, 2011; Yoseph *et al.*, 2016) supporting the milder phenotypical effects

518 following LPS exposure. The expression of *oclnb* was strongly increased after three hours in 0-6 hpf

519 cortisol-treated and dexamethasone-treated subjects compared to control-treated subjects. This

520 suggests lower permeability due to more protective leaky pathway activity (Kása *et al.*, 2015;

521 Odenwald and Turner, 2013; Shen *et al.*, 2011; Yoseph *et al.*, 2016). It has been shown that 24 hours

522 following LPS challenge in zebrafish larvae the expression of *oclnb* is strongly up-regulated facilitating

523 tissue-repair (Hsu *et al.*, 2018). Again, the data support the milder phenotypical effects that we see

524 following LPS exposure in cortisol-treated and dexamethasone-treated subjects compared to control-

525 treated subjects.

526 To explore the underlying mechanisms of glucocorticoid treatment we measured the 527 expression of a series of genes of interest and related their expression to the outcome of the 528 phenotypical changes and mortality. Five genes stood out in this respect, reflecting two receptors 529 (*tlr4bb* and *cxcr4a*), a factor involved in the transduction pathway of the expression of cytokines 530 (*myd88*) and two cytokines (*il1β* and *il10*).

531 While LPS exerts its effects through transduction mechanisms following binding to TLR4 in 532 mammals (Goulopoulou et al., 2015; Kása et al., 2015), in zebrafish this is not clear as yet: tlr4ba and 533 tlr4bb have been suggested to be paralogues rather than homologues and TLR4BA and TLR4BB 534 have thus far not been shown to be activated by LPS possibly by lack of a binding site for LPS 535 (Sepulcre et al., 2009; Sullivan et al., 2009). However TLR4BB has been shown to be involved in 536 inflammatory processes as *tlr4bb* transcript abundance is increased following tail fin amputation 537 (Chatzopoulou et al., 2016). Here, we observed an increase in tlr4bb expression in 0-6 hpf control-538 treated subjects over time, accompanied by a strong inflammatory response, which was less strong in 539 0-6 hpf cortisol-treated or dexamethasone-treated subjects, accompanied by a milder inflammatory 540 response. This suggests a role for TLR4BB in the LPS-induced response.

541 In humans CXCR4 has been implicated in recognition of LPS or being part of a 'LPS sensing 542 apparatus' in addition to TLR4 (Triantafilou et al., 2001, 2008). Furthermore, LPS increases the 543 expression of *cxcr4* through an NF-KB signalling pathway associated with increased micro-vascular 544 leakage in the lungs (Konrad et al., 2017) or increased colorectal tumor metastasis (Liu et al., 2017). 545 Here, we observed an increase in *cxcr4a* expression in 0-6 hpf control-treated subjects over time, 546 which was absent in 0-6 hpf cortisol-treated or dexamethasone-treated subjects. This difference in 547 gene expression may be associated with differences in the extent of vascular leakage as suggested 548 by the differences in the expression of genes involved in the endothelial barrier and differences in tail 549 fin oedema between treatments as discussed above. In zebrafish cxcr4a is found in endothelial cells

550 (blood vessels), while cxcr4b is not (Wei Chong et al., 2001), which may explain that we only observed 551 a phenotype-related effect for the expression profile of cxcr4a. Interestingly, CXCR4 has been 552 implicated in the development of tolerance to lethal doses of LPS in zebrafish larvae (Dios et al., 2014; 553 Novoa et al., 2009). Thus, this suggests that CXCR4 is involved in modulating the response to LPS. 554 Overall therefore our data warrant further studies into the role of TLR4BB and CXCR4 in LPS-555 induced sepsis in zebrafish as well as into the effects of early life glucocorticoid stimulation hereon. 556 Earlier studies have shown that MYD88 knockout larvae show enhanced survival to LPS 557 challenge (Hsu et al., 2018) and no increase in *il1* persistion (van der Vaart et al., 2013). Lethal, but 558 not sub-lethal, doses of LPS have been found to be associated with high expression levels of $il1\beta$ and 559 il10 in zebrafish larvae, indicative of a hyper-inflammatory response (Dios et al., 2014). In line with 560 these findings we observed that the expression levels of *myd88*, *il1* β and *il10* were strongly increased 561 3 hours after LPS challenge associated with low survival in 0-6 hpf control-treated subjects, but lower 562 levels of expression of all three genes with higher survival in 0-6 hpf cortisol-treated and 563 dexamethasone-treated subjects. Myd88 is an adaptor protein critical to toll-like receptor signalling 564 (except for TLR3; Goulopoulou et al., 2016) and IL1β receptor signalling (see Kanwal et al., 2013; van 565 der Vaart et al., 2013) and thereby cytokine expression. It should be noted that myd88 expression was 566 already low in 0-6 hpf cortisol-treated and dexamethasone-treated subjects, suggesting lower 567 transduction pathway activity, possibly leading to a lower stimulation of inflammatory pathways. It is 568 clear that this deserves further study.

As indicated above following cortisol or dexamethasone treatment at 0-6 hpf we noted increased base-line expression levels of *irg1l*, *socs3a*, *mpeg1.1* and *mpeg1.2* compared to control treatment at 0-6 hpf. These increased base-line levels may aid in increased clearance of bacteria and preventing excessive inflammation and hence aid in increasing survival (Benard *et al.*, 2015; Hall *et al.*, 2014; Jo *et al.*, 2005).

574 The data of LPS exposure in larvae observed here seem to match the data of LPS exposure 575 in adult zebrafish following 0-5 dpf exposure to cortisol: LPS exposure did not increase *il1* β expression 576 in different tissues measured (Hartig *et al.*, 2016). Unfortunately no survival was measured in the latter 577 study.

578 Overall these data suggest that the response to LPS of subjects pre-exposed to cortisol or 579 dexamethasone is less strong than the response of subjects of pre-exposed to control medium. How 580 differences in survival, phenotype and gene-expression levels are causally related remains to be

581 studied. As we have observed that 0-6 hpf exposure to dexamethasone had no effect on baseline

582 levels of cortisol at 5 dpf in larvae of the AB strain (Althuizen, 2018), the data suggest that these

- 583 effects are independent of baseline levels of cortisol.
- 584

585 **4.4 Limitations**

A clear limitation is that we only used one dose of cortisol and dexamethasone. For convenience we used equimolar doses of cortisol and dexamethasone, which may have led to different levels of stimulation of GR 0-6 hpf. Thus, in future studies different dose-ranges may be warranted, e.g. to study whether higher concentrations of cortisol in the tail fin amputation assay have an effect on neutrophil recruitment.

591 Regarding the tail fin amputation assay it has been shown that glucocorticoids may play a role 592 in the differentiation of macrophages into a pro-inflammatory (M1) phenotype (Xie et al., 2019). So, 593 future studies should study in more detail the effects of early life exposure of glucocorticoids on the 594 inflammatory response and wound healing. Similarly, we used LPS to induce a hyper-inflammatory 595 response, i.e. sepsis (Hsu et al., 2018; Philip et al., 2017), as a model to study the effectiveness of our 596 early life treatments. To assess the ecological relevance of our findings and their more general nature 597 the effects of early-life exposure on larval exposure to different pathogens, such as of bacterial, viral or 598 fungal origin, should be studied (see e.g. Meijer and Spaink, 2011; van der Vaart et al., 2013).

599 We observed a variable response to the LPS challenge in the two exposure series. This is not 600 uncommon as we also observed variable responses in other immune-related paradigms such as the 601 response to a dextran sodium sulphate (DSS) challenge (van den Bos et al., unpublished 602 observations). While as yet speculative differences in baseline levels of expression of $il1\beta$ may be one 603 associated factor as we observed that higher baseline levels seem associated with milder responses 604 to LPS (van den Bos et al., unpublished data). This is not unprecedented as this has also been 605 observed in mice: enhanced levels of IL1 are associated with a milder response to LPS (Alves-Rosa et 606 al., 2002). It has been shown that following hatching $il1\beta$ expression increases due to exposure to 607 microbes in the medium (Galindo-Villegas et al., 2012). This tunes the activity of the innate immune 608 system of the zebrafish larvae and determines thereby their disease resistance. In the laboratory 609 however this may lead to variation between experimental series as the microbial load of the medium

610 may vary from experimental series to series. It is clear that this warrants further studies. It has been

611 suggested that tuning the innate immune system occurs through chromatin modifications (see e.g.

612 Foster *et al.*, 2007; Galindo-Vargas *et al.*, 2012; Netea *et al.*, 2016, 2017). Our future studies are

613 directed at understanding the variable responses in this context.

614 The immune system affects HPI-axis activity and vice versa (Wendelaar Bonga, 1997). 615 However, we did not address changes in HPI-axis activity as a consequence of our procedures (tail fin 616 amputation or LPS exposure) in this study as the primary aim was to study whether our pre-treatments 617 would affect immune function. Still changes in HPI-axis activity may be anticipated. For example, a 618 recent study showed in European sea bass (Dicentrarchus labrax) larvae 5 day post-hatching that at 619 120 hours following infection with the bacterium Vibrio anguillarum, when mortality was already high, 620 HPI-axis activity increased (Reyes-López et al., 2018). This increased activity coincided with 621 increased expression of pro-inflammatory and anti-inflammatory genes. While the underlying 622 mechanism was not clear as yet, the data show that infections may impact HPI-axis activity. It has 623 been shown that early-life exposure to cortisol dampens the response of the HPI-axis to stressors in 624 zebrafish larvae (Nesan and Vijayan, 2016). Hence, future studies should address how early-life 625 exposure to cortisol affects the relationship between immune function and HPI-axis activity, and how 626 this relates to the increased survival that we have observed here.

627 We have used 0-6 hpf cortisol exposure by the medium as a model or proxy of increased 628 oocyte levels of cortisol due to chronic stress in mothers (van den Bos et al., 2019), while others have 629 used micro-injection of cortisol in the yolk of single-cell embryos (Best et al., 2017; Nesan and Vijayan, 630 2012, 2016). Chronic stress, whether due to excessive predation, food shortage, crowding or out-of-631 range-temperatures, is associated with increased levels of cortisol (Wendelaar Bonga, 1997) and at 632 this level these procedures may be a valid approach of mimicking maternal stress. Still, these different 633 stressors may have additional neuro-endocrine and/or metabolic signatures that affect oocyte yolk-sac contents and hence thereby development of embryos and larvae. Thus, future experiments should 634 635 compare the current results to data from chronically stressed mothers using different types of 636 stressors.

637

638 **4.5 Conclusion**

639	These data show that early	v-life exposure to cortisol.	as a model or prox	v of maternal stress. indu	lces
00,					

- 640 an adaptive response to immune challenges, which seems mediated via the glucocorticoid receptor.
- 641 These data are of relevance for both ecological research (Sopinka *et al.*, 2017) and biomedical
- research (Stewart *et al.*, 2014) in understanding the effects of stressful conditions and exposure to
- 643 endocrine disruptors on disease susceptibility and survival of offspring.
- 644

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- 649

650 6. Conflict of interest

- 651 The authors have no conflict of interest to report.
- 652

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- 655

656 8. Contributions

- 657 RvdB and MS conceived the project. SC, KT, JA and RW performed experiments. RvdB, SC, KT, JA
- and RW analysed data. JZ and SC performed qPCR analysis. RvdB, GF and MS wrote the
- 659 manuscript.
- 660
- 661

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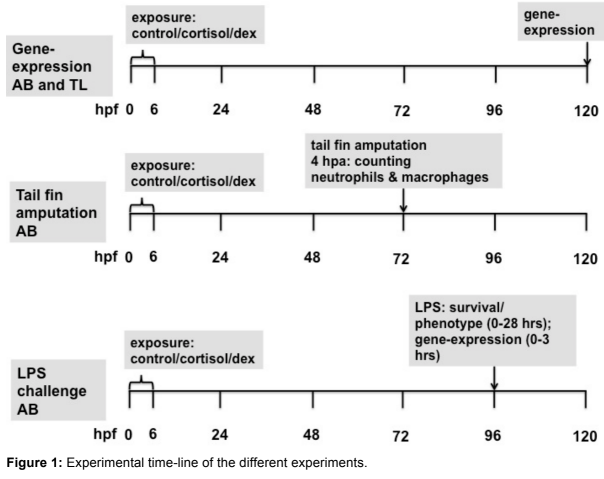
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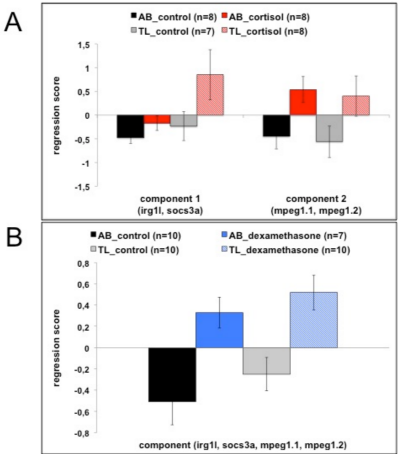
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903 **10. Legends figures**



907 dex=dexamethasone

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910 911 **Figure 2, panel A**: Regression scores (mean± SEM) of components 1 and 2 from the PCA for the

912 different treatments (cortisol or control) and strains (AB or TL). Genes that contributed to the

913 components are indicated in the figure. One subject was removed from the statistical analyses (TL

914 $\,$ $\,$ control) as it was a consistent outlier following Grubb's outlier test.

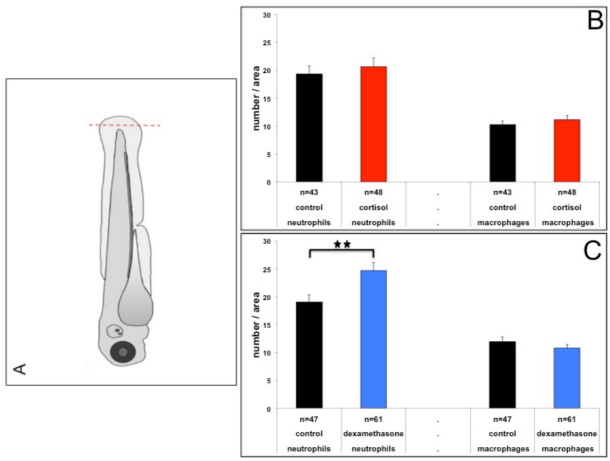
915 **Figure 2, panel B**: Regression scores (mean± SEM) of the only component of the PCA for the

916 different treatments (dexamethasone or control) and strains (AB or TL). Genes that contributed to the

917 component are indicated in the figure.

918

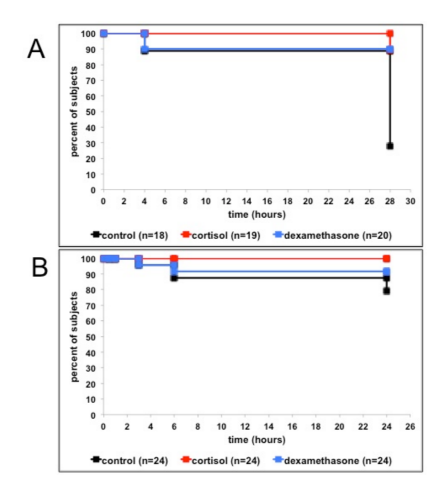
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- 920
 neutrophils
 neutrophils
 macrophages
 macrop

 921
 Figure 3, panel A Picture that illustrates the position of the site of the tail fin amputation in 3 dpf

 922
 larvae.
- 923 **Figure 3, panel B**: Number of neutrophils and macrophages (mean± SEM) at 4 hpa for the different
- 924 treatments (cortisol or control).
- 925 Figure 3, panel C: Number of neutrophils and macrophages (mean± SEM) at 4 hpa for the different
- 926 treatments (dexamethasone or control).
- 927
- 928



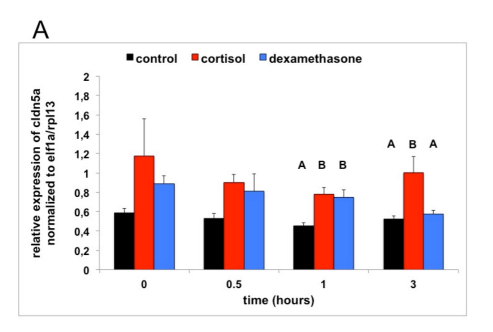
930 Figure 4, panel A: Per cent of surviving larvae following LPS treatment (from 0-0.5 hrs) and 0-6 hpf

control treatment, cortisol treatment or dexamethasone treatment in the first exposure series (time-points: 0 hr, 4 hrs and 28 hrs).

933 **Figure 4, panel B**: Per cent of surviving larvae following LPS treatment (from 0-0.5 hrs) and 0-6 hpf

control treatment, cortisol treatment and dexamethasone treatment in second exposure series (time-points: 0hr, 0.5 hr, 1 hr, 3 hrs, 6 hrs and 24 hrs).

936





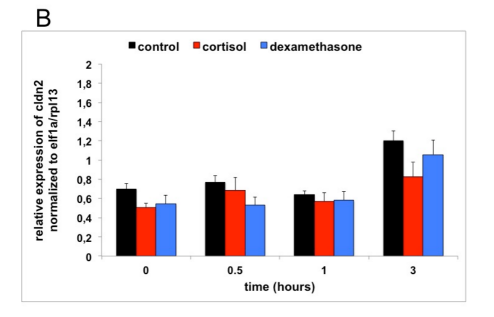
939 **Figure 5, panel A:** Transcript abundance (relative normalized expression; mean+SEM) of *cldn5a*

940 following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

941 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

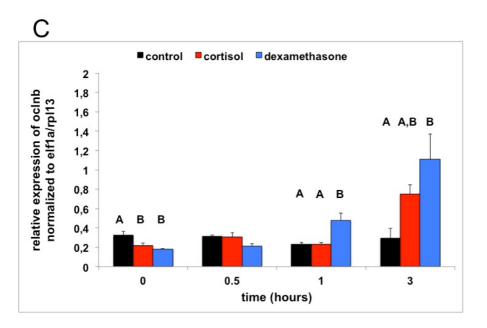
942 significantly differ from one another (Tukey HSD following a significant treatment effect for this time-

943 point). Note: control-treated subjects showed no significant change over time: F(3,12)=2.009, ns.



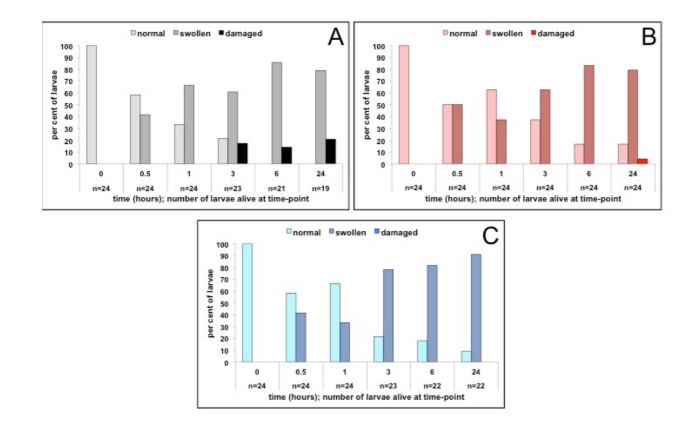
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Figure 5, panel B: Transcript abundance (relative normalized expression; mean+SEM) of *cldn2*following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or
dexamethasone treatment (n=4 samples per time-point). Note: control-treated subjects showed an
increased expression at 3 hrs: F(3,12)=13.253, p<0.001; Tukey HSD.



952	Figure 5, panel C: Transcript abundance (relative normalized expression; mean+SEM) of ocInb
953	following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or
954	dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not
955	significantly differ from one another (Tukey HSD following a significant treatment effect for this time-
956	point). Note: control-treated subjects showed no significant change over time: F(3,12)=0.562, ns.
957	
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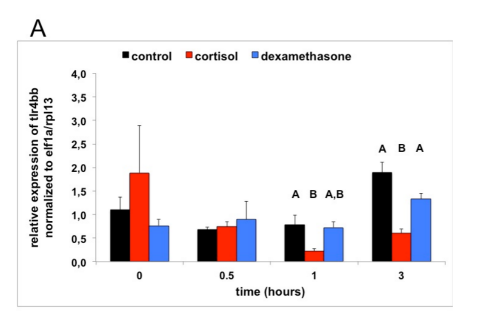
Figure 6, panel A: Per cent of larvae following LPS treatment (from 0-0.5 hr) in the 0-6 hpf control

treatment group showing normal, swollen or damaged tail fins (time-points: 0hr, 0.5hr, 1 hr, 3 hrs, 6hrs and 24 hrs).

963 Figure 6, panel B: Per cent of larvae following LPS treatment (from 0-0.5 hr) in the 0-6 hpf cortisol

treatment group showing normal, swollen or damaged tail fins (time-points: 0 hr, 0.5 hr, 1 hr, 3 hrs, 6

- 965 hrs and 24 hrs).
- 966 Figure 6, panel C: Per cent of larvae following LPS treatment (from 0-0.5 hr) in the 0-6 hpf
- 967 dexamethasone treatment group showing normal, swollen or damaged tail fins (time-points: 0 hr, 0.5
- 968 hr, 1 hr, 3 hrs, 6 hrs and 24 hrs).
- 969
- 970



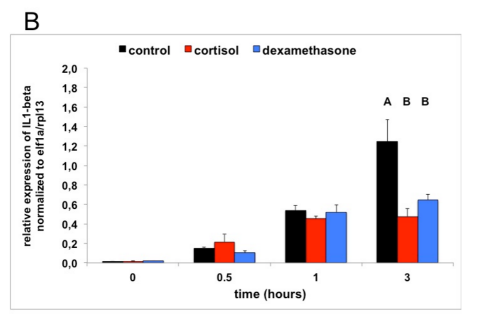
972 Figure 7, panel A: Transcript abundance (relative normalized expression; mean+SEM) of *tlr44bb*

973 following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

974 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

975 significantly differ from one another (Tukey HSD following a significant treatment effect for this time-





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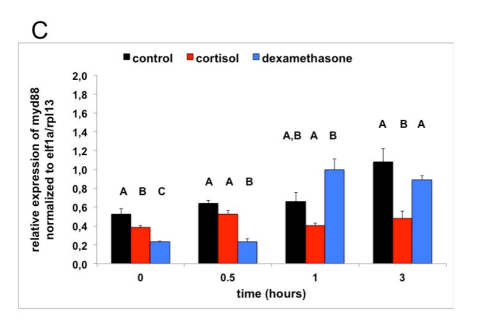
978 **Figure 7, panel B**: Transcript abundance (relative normalized expression; mean+SEM) of *IL1-beta*

following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

980 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

981 significantly differ from one another (Tukey HSD following a significant treatment effect for this time-

982 point).

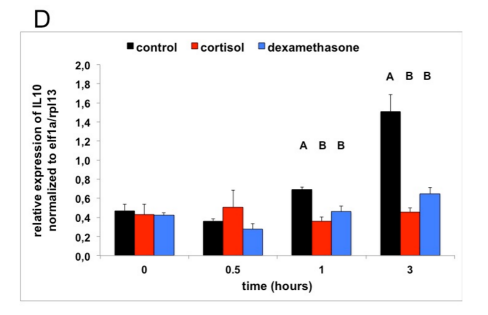


984 Figure 7, panel C: Transcript abundance (relative normalized expression; mean+SEM) of myd88

985 following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

986 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

987 significantly differ from one another (Tukey HSD following a significant treatment effect for this time-988 point).



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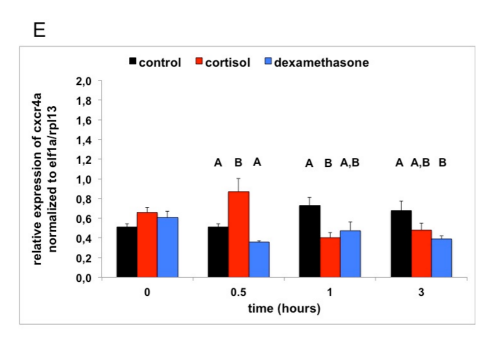
990 **Figure 7, panel D**: Transcript abundance (relative normalized expression; mean+SEM) of *IL10*

991 following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

992 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

993 significantly differ from one another (Tukey HSD following a significant treatment effect for this time-

994 point).



996 **Figure 7, panel E**: Transcript abundance (relative normalized expression; mean+SEM) of *cxcr4a*

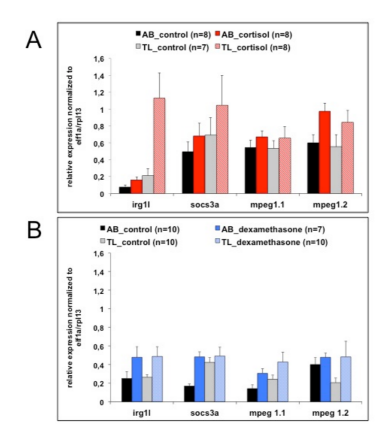
following LPS treatment (from 0-0.5 hr) and 0-6 hpf control treatment, cortisol treatment or

998 dexamethasone treatment (n=4 samples per time-point). Groups with the same capitals do not

significantly differ from one another (Tukey HSD following a significant treatment effect for this time-

- 1000 point).
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1003

Supplementary figure 1, panel A: Transcript abundance (relative normalized expression;

1005 mean+SEM) of the different immune related genes for the different treatments (cortisol or control) and

1006 strains (AB or TL). One subject was removed from the statistical analyses (TL control) as it was a

1007 consistent outlier following Grubb's outlier test. Treatment effects were found for *irg11*

1008 (F(1,23)=11.789, p≤0.01; treatment * strain: F(1,23)=8.446, p≤0.01; AB: p≤0.05; TL: p≤0.01) and

1009 mpeg1.1 (F(1,23)=8.614, p \leq 0.01).

- 1010 **Supplementary figure 1, panel B**: Transcript abundance (relative normalized expression;
- 1011 mean+SEM) of the different immune related genes for the different treatments (dexamethasone or
- 1012 control) and strains (AB or TL). Treatment effects were found for *irg1l* (F(1,33)=6.484, p≤0.05),
- 1013 socs3a (F(1,33)=7.655, p≤0.01) and mpeg1.1 (F(1,33)=5.487, p≤0.05).
- 1014

Gene	Forward primer sequence (5'-3')	Reverse primer sequence (5'-3')	Ampli con	Source	
			length (bp)		
elf1a	CTGGAGGCCAGCTCA AACAT	TCAAGAAGAGTAGTACC GCTAGCATTAC	85	NM_131263.1	
rpl13	TCTGGAGGACTGTA AGAGGTATGC	AGACGCACAATCTTGAG AGCAG	147	NM_212784	
cldn5a	CTCGGGGGGAAGCAT ACTCTG	TATTCCCCGTTGGATGT GGC	101	NM_213274.1	
cldn2	CCCCTACTGGGCAAC ATCTG	CTTGCAGGTCGACGGTT AGT	155	XM_005170272.4	
oclnb	TGACAAAGTCAGTGC TGCTCA	GGATATTCATCTGTGTC AAGCTCAT	162	NM_001008618.1	
tlr2	TGCTGTCGGTCGATT ACCTG	ACACAGGGAAAACGAAG GCT	130	NM_212812.1	
tlr4ba	ACTTTCAAGATTTCC GTTGAGGTAT	CCACAAGAACAAGCCTT TGCAG	209	NM_001131051.1	
tlr4bb	ATCACACTGCTGTGT TGCGA	AGGCCAGTGAGAGCTTT GTG	178	NM_212813.2	
tlr5a	TGGTTCCGGTTGTGA GTTCC	TCTTGAAGATGACACCG CGT	162	XM_001919017.6	
tlr5b	GGTCTTGAAGCGAG TGGTGT	CATTTTGCGCCAAGGTC AGT	118	NM_001130595.2	
il1β	GATGGCATGCGGGCA ATATG	AGCTCATTGCAAGCGGA TCT	116	NM_212844.2	
il10	GACCATTCTGCCAAC AGCTC	ACCATATCCCGCTTGAG TTCC	102	NM_001020785.2	
myd88	GTTTGCGCTCAGTCT TTGCC	GGTCAGAAAGCGCAGAA TGC	103	NM_212814.2	
cxcr4a	ACTTGTACAGCAGCG TCCTC	CTTTGGCGAACACCAAG TCG	174	NM_131882.3	
cxcr4b	GCGCCTTTTTGAGCA CACTT	ATTGCTGACTGAGAGGT CGC	127	NM_131834.1	
ptpn6	ACAGGAAGAACTGC GCTCAT	TGCAGCACAGCCAAGTA GAT	225	NM_199960.1	

Table 1: Nucleotide sequences of forward and reverse primers used for qPCR.

1017 Reference genes: elongation factor 1α (*elf1* α); ribosomal protein L13 (*rpl13*) 1018

4 hrs	treatment	%dead	%(alive+affected)	%(alive+intact)
			(tail fin, shape, heart)	
	control (n=18)	11.1	83.3	5.6
	cortisol (n=19)	0.0	73.7	26.3
	dexamethasone (n=20)	10.0	60.0	30.0
28 hrs	treatment	%dead	%(alive+affected)	%(alive+intact)
			(tail fin, shape, heart)	
	control (n=15)	86.7	13.3	0.0
	cortisol (n=16)	12.5	37.5	50.0
	dexamethasone (n=16)	12.5	31.3	56.3

1021**Table 2**: Per cent of larvae dead, alive and affected (regardless of phenotypical1022changes in the tail fin, shape or cardiac area), and alive and intact at 4hrs and 28hrs1023after LPS treatment (0-0.5hr) in the different treatment groups: 0-6 hpf treatment with1024control, cortisol-containing or dexamethasone-containing medium. Note that the1025number of larvae at 28hrs is lower than at 4hrs; per treatment n=3-4 larvae (alive)

1026 were randomly selected for staining for reactive oxygen species in the tail fin (not

1027 reported here); they were not scored for phenotypical changes.