ABSTRACT

Chronic early-life stress increases adult susceptibility to numerous health problems linked to chronic inflammation. One way that this may occur is via glucocorticoid-induced developmental programming. To gain insight into such programming we treated zebrafish embryos with cortisol and examined the effects on both larvae and adults. Treated larvae had elevated whole-body cortisol and glucocorticoid signaling, and upregulated genes associated with defense response and immune system processes. In adulthood the treated fish maintained elevated basal cortisol levels in the absence of exogenous cortisol, and constitutively mis-expressed genes involved in defense response and its regulation. Adults derived from cortisol-treated embryos displayed defective tailfin regeneration, heightened basal expression of pro-inflammatory genes, and failure to appropriately regulate those genes following injury or immunological challenge. These results support the hypothesis that chronically elevated glucocorticoid signaling early in life directs development of a pro-inflammatory adult phenotype, at the expense of immunoregulation and somatic regenerative capacity.

KEY WORDS: Chronic stress, Glucocorticoid, Developmental programming, Immune system, RNA-seq, Zebrafish

INTRODUCTION

Chronic psychosocial stress contributes significantly to a wide variety of public health problems endemic to the modern world, many of which are linked to chronic inflammation (Berk et al., 2013; Christian et al., 2006; Cohen et al., 2007; Pawelec et al., 2014; Radek, 2010). A growing body of epidemiological evidence indicates that individuals that experience chronic stress early in life – even prenatally – tend to develop a pro-inflammatory phenotype, and hence increased susceptibility to inflammatory disease and degenerative aging in adulthood (Cohen et al., 2012; Harris and Seckl, 2011; Howerton and Bale, 2012; Miller et al., 2009). The systemic stress response in humans is mediated by the glucocorticoid hormone cortisol, a steroid secreted by the adrenal cortex in response to adrenocorticotropic hormone (ACTH). The latter is released by the anterior pituitary in response to corticotropin-releasing hormone (CRH) emitted by the hypothalamus in response to stress signals from the brain. This system, known as the hypothalamus-pituitary-adrenal (HPA) axis, induces metabolic and other cell physiological changes directed at stress mitigation. Following acute stress, the system turns itself off by way of a negative feedback loop wherein receipt of cortisol in the hypothalamus and pituitary extinguishes emission of CRH and ACTH. Chronic psychosocial stress compromises this mechanism by continuously activating the HPA axis, leading to chronically elevated cortisol levels, and ultimately (at least in some cases) to glucocorticoid resistance (Jung et al., 2015; Maldonado Bouchard and Hook, 2014; Stark et al., 2001).

Cortisol and other corticosteroids exert their biological effects by binding and thereby activating the glucocorticoid and mineralocorticoid receptors, DNA sequence-specific transcription factors in the steroid receptor family (Gomez-Sanchez and Gomez-Sanchez, 2014; Kadmiel and Cidlowski, 2013). Owing to its higher affinity for cortisol, the mineralocorticoid receptor is more or less constitutively active, whereas the glucocorticoid receptor (GR) is activated under stressful conditions, and is largely responsible for inducing the stress-responsive gene expression program (Sorrells and Sapolsky, 2007). The GR regulates gene expression both directly, by binding glucocorticoid responsive DNA sequence elements within the cis-regulatory domains of genes, and indirectly, by binding other transcription factors such as NF-xB. Direct targets include genes required for the metabolic changes entailed by the stress response, whereas indirect targets include immune system genes activated by NF-xB, which is inhibited by the GR, accounting in part for the well-known anti-inflammatory effects of glucocorticoids.

Elevated glucocorticoid signaling has epigenetic effects that are thought to mediate long-term developmental programming in response to early life adversity (Khulan and Drake, 2012). Consistent with this, chronic exposure to elevated corticosteroids during early development has been shown in a number of systems to have lasting phenotypic effects that can not only impact the health of the exposed individual, but also that of the individual’s offspring (Drake et al., 2007). The epigenetic mechanisms for some of these effects have been elucidated, and include methylation of the GR promoter, resulting in lower GR expression and consequently glucocorticoid resistance (Chen et al., 2009; Tsyka et al., 2012; Weaver et al., 2004). However, the stress response system involving glucocorticoid signaling is complex and regulated at many levels, and the systemic effects of glucocorticoid-mediated developmental programming, including the global impact such programming has on somatic gene expression and regenerative capacity in the adult, is largely unknown.

The zebrafish is a premier model organism for studies of regeneration, owing to its remarkable ability to regenerate many of its body parts, including its tailfin, which provides a convenient assay for regenerative capacity. Studies of tailfin regeneration have begun to illuminate the genetic pathways required for regeneration, and it was recently shown that macrophages play an essential role in the process (Petrie et al., 2014), as has also recently been shown to be true for salamander limb regeneration (Godwin et al., 2013). Of interest in light of the results reported here, the GR is known to play a central role in macrophage regulation (Chinenov et al., 2014).
Zebrafish have recently emerged as a model for examining the role of glucocorticoid signaling in developmental programming (Nesan and Vijayan, 2013). In this study we used zebrafish to ask how exposure to chronically elevated cortisol during early development affects gene expression and regenerative capacity in adulthood.

RESULTS AND DISCUSSION
Glucocorticoid signaling is activated in zebrafish embryos exposed to exogenous cortisol

Zebrafish embryos treated continuously with sub- to low-micromolar doses of exogenous cortisol from 0 to 5 days post-fertilization (dpf) were found to produce morphologically normal larvae (Fig. 1A), but with elevated whole body cortisol levels, as indicated by ELISA of larval extracts (Fig. 1B). The treated larvae upregulated known glucocorticoid-responsive genes (Fig. S1A), and larvae of a glucocorticoid-responsive GFP reporter line (SR4G, Krug et al., 2014) displayed elevated fluorescence in response to the treatment (Fig. 1C; Fig. S1B), indicating that the treatment activated the glucocorticoid receptor. Treated larvae had an elevated heart rate and increased levels of reactive oxygen species (Fig. S2), expected effects of stress signaling. We conclude from these results that zebrafish embryos treated with exogenous cortisol activate glucocorticoid receptor signaling and mount a systemic stress-response.

Cortisol-treated larvae upregulate genes involved in immune system processes and the immune response

High-throughput sequencing of RNA (RNA-seq) extracted from larvae at 5 dpf was used to examine the genome-wide transcriptional effects of the cortisol treatment. Of the ~20,000 expressed genes, 411 were found to be significantly (P<0.05) over-expressed and 144 were found to be significantly under-expressed in the cortisol-treated larvae. PANTHER overrepresentation analysis (Mi et al., 2013) of gene ontology (GO) biological process (BP) terms associated with the 555 genes that were significantly differentially expressed indicated a strong effect of the treatment on processes related to immune cell function and inflammation (Fig. 1D; Table S1). Similarly, GO BP term enrichment analysis using the GOrilla algorithm (Eden et al., 2009) to interrogate the full list of 9523 upregulated genes (i.e. all genes with relative expression treated/untreated >1) ranked by significance identified immune system process as the most significantly enriched BP term (Table S2), and visualization of all the significantly enriched BP terms (Supek et al., 2011) identified immune response and cellular lipid catabolism as the two largest parent categories of terms (Fig. S3A). Examples of significantly upregulated genes associated with immune response and/or regulation included irg1l (immuneresponsive gene 1-like; Hall et al., 2014), the macrophage marker mpeg1 (Ellett et al., 2011), and the immunoregulatory gene socs3a (Qin et al., 2012) (Fig. 1E). GOrilla analysis of the 10,547 downregulated genes ranked by significance (Table S2) identified membrane depolarization, cell communication, and long-chain fatty acid metabolism (including arachidonic acid metabolism, known to regulate inflammation; Buckley et al., 2014) as the three largest parent categories of enriched BP terms (Fig. S3B). Examples of significantly downregulated genes associated with regulation and/or resolution of inflammation included cyp2aa8 (Su and Ding, 2004), anxa1c (Vago et al., 2012), and klf2b (Nayak et al., 2013) (Fig. 1F). These data are consistent with known metabolic effects of cortisol-treated larvae upregulate genes involved in immune system processes and the immune response

High-throughput sequencing of RNA (RNA-seq) extracted from larvae at 5 dpf was used to examine the genome-wide transcriptional effects of the cortisol treatment. Of the ~20,000 expressed genes, 411 were found to be significantly (P<0.05) over-expressed and 144 were found to be significantly under-expressed in the cortisol-treated larvae. PANTHER overrepresentation analysis (Mi et al., 2013) of gene ontology (GO) biological process (BP) terms associated with the 555 genes that were significantly differentially expressed indicated a strong effect of the treatment on processes related to immune cell function and inflammation (Fig. 1D; Table S1). Similarly, GO BP term enrichment analysis using the GOrilla algorithm (Eden et al., 2009) to interrogate the full list of 9523 upregulated genes (i.e. all genes with relative expression treated/untreated >1) ranked by significance identified immune system process as the most significantly enriched BP term (Table S2), and visualization of all the significantly enriched BP terms (Supek et al., 2011) identified immune response and cellular lipid catabolism as the two largest parent categories of terms (Fig. S3A). Examples of significantly upregulated genes associated with immune response and/or regulation included irg1l (immuneresponsive gene 1-like; Hall et al., 2014), the macrophage marker mpeg1 (Ellett et al., 2011), and the immunoregulatory gene socs3a (Qin et al., 2012) (Fig. 1E). GOrilla analysis of the 10,547 downregulated genes ranked by significance (Table S2) identified membrane depolarization, cell communication, and long-chain fatty acid metabolism (including arachidonic acid metabolism, known to regulate inflammation; Buckley et al., 2014) as the three largest parent categories of enriched BP terms (Fig. S3B). Examples of significantly downregulated genes associated with regulation and/or resolution of inflammation included cyp2aa8 (Su and Ding, 2004), anxa1c (Vago et al., 2012), and klf2b (Nayak et al., 2013) (Fig. 1F). These data are consistent with known metabolic effects.
elevated GR signaling (Sorrells and Sapolsky, 2007), but also suggest that prolonged activation of glucocorticoid receptor signaling during early development upregulates immune system gene activity. Since many immune system genes are responsive to the transcription factor NF-κB, we reasoned that NF-κB activity may be constitutively higher in cortisol-treated larvae. This was indeed the case, as shown by increased fluorescence of 5 day old cortisol-treated larvae of a transgenic NF-κB-responsive GFP reporter line (Kanther et al., 2011) (Fig. S4).

**Adults developed from cortisol-treated embryos have elevated basal cortisol and aberrant constitutive expression of genes involved in immune system response and regulation**

The cortisol treatment regime did not significantly affect the rate that treated fish grew to adulthood (Fig. S5) and the adults derived from treated embryos displayed no obvious morphological or behavioral defects. However, adult fish derived from the treated embryos were found on average to have constitutively elevated basal cortisol levels (Fig. 2A), indicating that the treatment had a long-term activating effect on the stress system. To assess the long-term effects of the treatment on constitutive gene expression, RNA-seq was performed on RNA extracted from blood and muscle of 4.5-month-old adults derived from cortisol-treated larvae and their untreated (control) siblings. In both tissues more genes were found to be under-expressed than over-expressed in the fish derived from cortisol-treated embryos (Fig. 2B). GOrilla analysis (Tables S3 and S4) identified leukocyte chemotaxis as the most significant parent category for genes under-expressed in muscle, with multiple subcategories in common between the two tissues, including defense response, inflammatory response, response to chemical and response to external stimulus (Fig. S6). Fourteen genes were significantly differentially expressed both in blood and muscle (Fig. 2C, Table 1), twelve of which were under-expressed in fish derived from cortisol-treated embryos. The latter included irg1l and socs3a (Table 1), two genes associated with immune system response and regulation that were among the most highly upregulated genes in the cortisol-treated larvae (see Fig. 1E). Quantitative reverse-transcription and polymerase chain reaction (qRT-PCR) confirmed that the latter two genes were differentially expressed in both blood and muscle of older adults derived from the same batch of treated embryos, although irg1l was found to be over-expressed rather than under-expressed in blood (Fig. 2D). The latter difference may reflect the fact that the fish used for the qRT-PCR analysis, while from the same experimental cohort as those used for RNA-seq, were significantly older (9 months). Indeed, in 5-month-old fish from another experimental cohort irg1l was again found to be significantly under-expressed in both blood and muscle (Fig. 2E). However, in these fish socs3a was not under-expressed, indicating that there is biological variability in the response of specific genes to the treatment. Despite this variability (not unexpected in wild-type fish), these results suggest that short-term treatment of zebrafish embryos with exogenous cortisol has long-term effects on the constitutive (basal) activities of both the stress and immune systems.

**Fig. 2. Effects of early developmental exposure to cortisol on adult basal cortisol levels and gene expression in blood and muscle cells.** (A) Cortisol levels measured by ELISA of whole body extracts from individual 7-month-old adults derived from three experimental cohorts exposed as embryos to the indicated concentrations of cortisol from 0-5 dpf. Error bars in the first two graphs depict standard deviations (s.d.) for technical replicates; for the third graph showing combined data from all three experiments, the error bars represent s.e.m. for biological replicates. For the third experiment a power analysis was performed which determined that a minimum of five individuals was needed for statistically significant results based on the effect sizes obtained in the first two. Statistical significance $^{*}P<0.05, ^{**}P<0.01, ^{***}P<0.001$ by an unpaired t-test, excluding the single outlier (Q-test, $Q_{exp}=0.649>Q_{crit}=0.569$, $P<0.05$) in the control group. (B) Scatter plots of average expression level and fold-change in expression for each gene in blood and muscle tissue from 4.5-month-old adults derived from cortisol treated with 1 μM cortisol and untreated controls. Red dots indicate genes with significantly different expression ($P<0.05$) between the two groups, the numbers of which (over- and under-expressed) are also shown in red. (C) Venn diagram showing numbers of differentially expressed genes in blood and muscle, and the overlap between the two (14 genes, listed in Table 1). (D,E) qRT-PCR measurements of socs3a and irg1l in blood and muscle tissue from cortisol treated embryos (+) and untreated control embryos (−) from (D) 9-month-old zebrafish from the same experimental cohort from which the RNA-seq results shown in B and C were obtained, and (E) 5-month-old zebrafish from a different experimental cohort. Statistical significance $^{*}P<0.05, ^{**}P<0.01, ^{***}P<0.001$ by an unpaired t-test, the error bars represent s.d.
Adul ts d eveloped from cortisol-treated e mbryos have tailfin regeneration defects including aberrant expression of inflammatory genes

In a tailfin regeneration assay, several cohorts of adults in the cortisol-treated groups were found to produce a significantly higher frequency of morphological defects, ranging from mild patterning defects to complete failure of parts of the fin to regenerate (Fig. 3A, B; note caveat in the figure legend). Given the results described above and the recent report that macrophages play an essential role in adult tailfin regeneration (Petrie et al., 2014), we reasoned that the observed regeneration defects might reflect an aberrant inflammatory response to injury. To begin to assess whether this might be so we employed mpx:GFP (Mathias et al., 2006) and mpeg1::YFP (Roca and Ramakrishnan, 2013) transgenic lines to examine the respective dynamics of neutrophil and macrophage infiltration during tailfin regeneration. In fish from the cortisol-treated groups, greater numbers of neutrophils were found to initially enter the wound site, which was ultimately followed by greater numbers of macrophages in the regenerating fin (Fig. 3C), suggesting that these fish mount an exaggerated and possibly prolonged inflammatory response to injury.

RNA-seq analysis of regenerating tailfins from 4.5-month-old fish revealed that at 2 days post-amputation (dpa; corresponding to blastema formation), significantly fewer genes were mobilized in regenerating tissue of fish derived from cortisol-treated embryos (815 vs 2353 in controls; Fig. 3D). Gene ontology BP term enrichment (GOrilla) analysis of the genes differentially expressed between 2 and 0 dpa (Tables S5 and S6) gave strikingly contrasting results for fins from treated and control fish; the upregulated set from the controls was enriched with genes annotated with the BP parent categories cell division, nuclear division, regeneration (including fin regeneration), response to stress, leukocyte migration, and immune system process, whereas genes annotated in those categories were not enriched in the upregulated set of the cortisol-treated group (Fig. S7). Rather, genes annotated in categories associated with the immune system were instead enriched in the downregulated set of the cortisol-treated group (Fig. S8). This result was confirmed by comparing the interaction between treatment and regeneration time-course, which revealed that compared to controls, 274 genes were significantly differentially expressed in the fin tissue from treated fish at 2 dpa vs 0 dpa, with 260 of these being significantly under-expressed (Fig. 3E), many of them associated with the immune system defense response and/or its regulation (Fig. S9, Table S7). Plotting the temporal profiles of several of these genes revealed that they were highly but transiently upregulated at 2 dpa in regenerating fins of control fish, whereas their expression in treated fish was inverted; basally elevated compared to controls in the pre-regenerative tailfin (day 0), then transiently decreased at 2 dpa in the regenerating fin before becoming elevated again with respect to controls at 4 dpa (Fig. 3F). A similar pattern was observed on a global level when RNA-seq data from control and treated groups were compared at each time point and the results subjected to GOrilla analysis (Tables S8-S10): genes associated with immune system process and other categories related to immune system functions were basally over-expressed in the tailfin, but under-expressed in the regenerating tailfin at 2 dpa, then again over-expressed at 4 dpa (Fig. S10). Quantitative RT-PCR measurements of marco, il-1b, infg1-1 and infb from the same cohort of fish six months later confirmed that tailfin expression of these pro-inflammatory genes in the treated fish is basally elevated, then sharply downregulated during regeneration, inverse of their expression in untreated controls (Fig. 3G).

The results described above suggested that adult fish derived from cortisol-treated embryos have both aberrant basal expression of immune system genes and an impeded immune response. To further test this we injected adult fish with lipopolysaccharide (LPS), and used qRT-PCR to assay the expression of the LPS-responsive pro-inflammatory LPS-TLR4-responsive genes il-1b, il6 and infa as well as the il-6-responsive immunoregulatory gene socs3a in peripheral tissues (spleen, skeletal muscle and heart), blood and brain. As expected, in control fish all four genes were strongly and specifically induced by LPS, albeit to different degrees in different tissues (Fig. 4). In contrast, in fish derived from cortisol-treated embryos LPS-induced expression of these genes was much weaker, in many cases being no different (or lower) than the expression induced non-specifically by PBS; in addition, basal expression of il-1b, il-6 and socs3a was elevated in the heart (Fig. 4). Thus, chronic elevation of glucocorticoid signaling during early development has long term effects that increase the basal expression and/or non-specific induction of pro-inflammatory genes in adult tissues, while suppressing their specific induction in response to pathogenic challenge.

Table 1. Genes that are differentially expressed in both blood and muscle of 7-month-old zebrafish derived from cortisol-treated embryos, compared to untreated controls

| ENSEMBL gene ID | Gene symbol | Description | Family | Log2 (fold change) blood | Log2 (fold change) muscle |
|-----------------|-------------|-------------|--------|------------------------|-------------------------|
| ENSDARG00000090382 | BX005305.5 | Protein coding | Peptidase C13 | −2.66 | −5.48 |
| ENSDARG00000086549 | BX005305.1 | Protein coding | Peptidase C13 | −2.30 | −6.31 |
| ENSDARG000000104109 | CUB855711.2 | Protein coding | Peptidase C13 | −1.62 | −4.61 |
| ENSDARG000000100172 | CAB201042671.1 | miRNA | Novel | 5.98 | 6.19 |
| ENSDARG000000100469 | 5S_rRNA | 5S ribosomal RNA | 5.43 | 5.74 |
| ENSDARG00000083378 | SNORA63 | Small nuclear RNA | −2.30 | −6.65 |
| ENSDARG00000088643 | CUB855711.1 | Protein coding | Peptidase C13 | −2.08 | −5.48 |
| ENSDARG00000084532 | SNORA31 | Small nuclear RNA | −1.90 | −2.56 |
| ENSDARG00000025783 | sic:h211-12566.11 | Protein coding | C-type lectin | −2.18 | −3.04 |
| ENSDARG000000103201 | sic:dkcyp-3b12.9 | rRNA | −5.80 | −5.85 |
| ENSDARG00000024528 | socs3a | Suppressor of cytokine signaling 3a | −1.45 | −2.06 |
| ENSDARG00000081082 | SNORA16 | Small nuclear RNA | −4.38 | −3.44 |
| ENSDARG00000062788 | irg1 | Immunoresponsive gene 1-like | −1.95 | −3.15 |
| ENSDARG00000058206 | sic:h211-153b23.5 | Protein coding | Type 1 glutamine amidotransferase | −1.99 | −2.50 |
Collectively these data show that cortisol-treated embryos develop into adults wherein normal regulation of the immune system and inflammatory response is compromised. Our results further suggest that chronically elevated glucocorticoid signaling during early development reduces regenerative capacity of the adult. Although a previous report showed that synthetic glucocorticoids (but not cortisol) impair tailfin regeneration in larvae independently of their anti-inflammatory effects (Mathew et al., 2007), that study did not address the long-term effects of chronically elevated glucocorticoid signaling during early development on tailfin regeneration in adults, which is a very different context not directly comparable to that of the larva. It is thus possible that the regeneration defects that we observed were a consequence of the aberrant immunoregulation in the cortisol-treatment group, consistent with the emerging consensus that regulation of immune system activity is a critical determinant of adult somatic regenerative capacity (Forbes and Rosenthal, 2014). The observed effects on immunoregulation could stem from effects that elevated glucocorticoid signaling had on the developmental specification of immune cell fate, and/or from glucocorticoid-induced epigenetic changes that reduce the sensitivity of immune genes to glucocorticoids or other immunoregulatory signals. While further studies are needed to explore these possibilities, the findings reported here indicate that the zebrafish will provide a useful model to study the chronic early life stress affects adult immunoregulation.

MATERIALS AND METHODS
Zebrafish strains, husbandry and treatments

Unless otherwise noted, all experiments were performed using the AB strain of zebrafish. The SR4G transgenic reporter line (Krug et al., 2014) was kindly provided by Dr Karl Clark of the Mayo Clinic. Thempeg1:YFP (Roca and Ramakrishnan, 2013) and mpx:GFP (Mathias et al., 2006) reporter lines were provided by Dr Voot Yin of the MDI Biological Laboratory. The NF-κB-responsive GFP reporter line (Kanther et al., 2011) was kindly provided by Dr Karl Clark of the Mayo Clinic. The sr4G:GFP transgenic reporter line (Krug et al., 2014) was kindly provided by Dr Karl Clark of the Mayo Clinic. The mpeg1:YFP (Roca and Ramakrishnan, 2013) and mpx:GFP (Mathias et al., 2006) reporter lines were provided by Dr Voot Yin of the MDI Biological Laboratory.
conductivity of 600-700 microsemens, pH of 7.2. Lights are timed on a 14 h light/10 h dark cycle. Matings and embryo/larva culture were carried out using standard procedures (Nasiadka and Clark, 2012). Embryos and larvae were cultured in plastic Petri dishes from 0-5 days post-fertilization (dpf).

For cortisol treatments, stock solutions of cortisol-21-hemisuccinate sodium salt (Sigma H4881) in DMSO were added to the larval medium to achieve the indicated final concentrations; controls consisted of larvae in medium with an equivalent amount of DMSO vehicle. Cortisol and vehicle were replaced daily with media changes. At 5 dpf the fish were transferred to system tanks without water flow. Water flow began at 10 dpf. Fish were kept at a density of 30 fish/liter until 1 month post-fertilization, then were split up to a density of 5 fish/liter maximum. From 5-14 dpf the fish were fed O.S.I. Rotofier Diet three times per day. From 15-21 dpf the fish were fed Golden Pearl Active Spheres (Brine Shrimp Direct) 50-100 μm three times per day, and once a day with Brine Shrimp Direct Spirulina & Kelp Flake. Adults are fed once per day with live artemia, hatched daily and 1× per day with a 50:50 mixture of Golden Pearl Active Spheres 800-1000 μM and Aquaneering Fish Food Zebrafish Select Diet.

Caudal fin amputations and regeneration
Adult zebrafish were anesthetized in tricaine and caudal fins were amputated halfway between the edge of the caudal peduncle and the distal tip of the fin, using a razor blade. Regeneration progress was monitored at post-amputation time points using an Olympus MVX10 fluorescence stereomicroscope. Fin areas were quantified using ImageJ (NIH).

LPS injection
Adult zebrafish were fasted for 24 h, and then anesthetized in tricaine. Fish were injected intraperitoneally with 10 μl 1 mg/ml lipopolysaccharide (Sigma L3012) in PBS or PBS only, using a 10 μl 701-LT Hamilton syringe fitted with a 35-gauge needle. Fish were then transferred immediately back into system water and fasted for an additional 24 h before tissue collection.

Tissue collection
Adult zebrafish were euthanized in tricaine and blood was collected using the method described by Babaei et al. (2013). Other tissues were dissected out and snap frozen in liquid nitrogen. All procedures were approved by the Institutional Animal Care and Use Committee (IACUC).

Cortisol measurements
Larvae were washed of treatment media and pooled (n=30), then snap frozen in liquid nitrogen. Adult zebrafish were euthanized in tricaine and snap frozen in liquid nitrogen and frozen tissues were minced with a razor. Samples were homogenized in PBS using an automatic pestle grinder. A liquid-liquid extraction was performed with ethyl acetate and the organic, cortisol-containing phase was collected. The solvent was evaporated under a stream of nitrogen gas, and the cortisol extract was re-dissolved in Cortisol ELISA Extraction Buffer (Neogen Corporation). Samples were assayed for cortisol content using the Neogen Cortisol ELISA kit and quantified with a standard curve.

ROS measurements
Zebrafish larvae were incubated in 10 μM CM-H2DCFDA (Molecular Probes/Life Technologies C6827) in embryo water at 28°C for 1 h, 15 min
under darkness. Larvae were anesthetized in tricaine and imaged using an Olympus MVX10 fluorescence stereomicroscope. Fluorescence intensity was quantified using ImageJ software (NIH).

**Fluorescence imaging**

Fluorescently labeled zebrafish larvae and adults were imaged using an Olympus MVX10 fluorescence stereomicroscope, following their anesthetization in tricaine. Fluorescence data were quantified using NIH ImageJ software, as integrated pixel intensity within a defined region of interest (ROI), minus the average background pixel intensity times the ROI area.

**RNA-seq**

Total RNA was extracted from 5-day larvae (30 per each of three biological replicates) or dissected tissues from adults (5 adults per each of three biological replicates) using the RNA-EZplus kit from Qiagen, and quantified using a Nanodrop spectrophotometer. Indexed strand-specific polyA+-selected mRNA libraries were prepared and paired-end sequenced on an Illumina HiSeq2500 at the HudsonAlpha Institute for Biotechnology (Birmingham, AL) following manufacturer’s protocols. Following sequence read quality control diagnostic analyses using FastQC version 0.11.2 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/), reads were trimmed using Trimmatic version 0.32 (Bolger et al., 2014). Trimmed reads from the larval libraries were aligned to the zebrafish (Danio rerio) genome assembly version Zv9 using TopHat version 2.0.11 (Trapnell et al., 2009) and read counts per gene were generated using HTSeq version 0.6.1 (Anders et al., 2015). The larval RNA-seq data were deposited in the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database, under accession number GSE80221. The trimmed reads from the 4.5-month-old adult libraries were aligned to the annotated transcriptome from Ensembl version 83 (Flicek et al., 2014), using RSEM version 1.2.5 (Li and Dewey, 2011) and Bowtie 1.2.1 (Langmead et al., 2009). Read counts per gene from either HTSeq or RSEM were analyzed using R statistical computing environment (http://r-project.org) version 3.2.1 and R/edgeR version 3.2.1 (Robinson et al., 2010). The adult RNA-seq data were deposited in the NCBI GEO database, under accession number GSE80260. Both larval and adult data sets can be accessed via the GEO SuperSeries, under accession number GSE80286.

**Gene ontology enrichment analysis**

Gene ontology enrichment analysis was performed using PANTHER (Mi et al., 2013) or GOriilla (Eden et al., 2009), using either a single ranked list or two unranked lists (target and background) of genes, as described in the Results and figure legends for each analysis. Enriched terms found by GOriilla were visualized using REVIGO (Supek et al., 2011).

**Measurement of gene expression by quantitative PCR**

RNA prepared from whole larvae (30 larvae) or dissected adult tissues (from five fish, determined by a power analysis to be sufficient for statistical significance given measured effect sizes) as described above was converted to cDNA using Superscript III (Life Technologies), and transcript levels were quantified by quantitative real-time PCR (qPCR) measurement of SYBR Green fluorescence using Quanta Biosciences PerfeCTa FastMix on a Roche Lightcycler 480II. Primer sequences are shown in Table 2. Relative levels of expression were calculated using the delta-delta Ct method, using the average of Cts obtained for beta-actin and RPL13a for normalization.

**Author contributions**

The authors declare no competing or financial interests.

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**Data availability**

The larval and adult RNA-seq data were deposited in the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database (http://www.ncbi.nlm.nih.gov/geo/), under accession numbers GSE80221 and GSE80260, respectively. Both larval and adult data sets can be accessed via the GEO SuperSeries, under accession number GSE80286.

**Supplementary information**

Supplementary information available online at http://bio.biologists.org/lookup/doi/10.1242/bio.020065.supplemental

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**Table 2. Primer sequences used for qPCR**

| Gene   | Forward primer                      | Reverse primer                      |
|--------|------------------------------------|-------------------------------------|
| b-actin| TGGCCCGTTGATGCCCTCTGTTG            | AGGTCGACGCGGAGATGTC                |
| fkbp5  | GAGAATCCAGAGGGCATGGTTG            | AGGCGCTAGTTGGCTTTAGA               |
| ifrl1-1| GCCCGGCTGATTGGCTCTG                | TTGTTGATCTCTGTTTAAGTAGG            |
| il-1b  | CAGATCGGGAGAGCAAGCAAAGGG         | TGGTTGACAGTGAGCTTTTC               |
| i-6    | TTGCGTACACCTGCTCAGCTCTCA          | GCTGTGGTGGAGCGAAAAGGAC             |
| irg1f1  | AGACCCGTGTTGGGTCAAAGG            | TGGTTGACAGTGAGCTTTTC               |
| marco  | AGCCCAAGGGGATTAAGAGGAC            | AGGACGACATCCAGCAAACCTAAGGAAGG     |
| mmp13  | AGCATCGGACTTCCCAAGAGAAGGAAGGAAGG | AGGACGACATCCAGCAAACCTAAGGAAGG     |
| rpl13a  | TGGTTGACAGTGAGCTTTTC             | AGGACGACATCCAGCAAACCTAAGGAAGG     |
| rplb   | TGGTTGACAGTGAGCTTTTC             | AGGACGACATCCAGCAAACCTAAGGAAGG     |
| rpl13d3| GACAGAGGAAGCTTGCTC               | AGGACGACATCCAGCAAACCTAAGGAAGG     |

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