

Figure S1.
The stress predominantly affects the first phase of heart regeneration.

(A-K) Heart sections at 60 dpci after AFOG staining were analyzed in the same manner as described in Fig. 2. After cryoinjury, zebrafish were subjected to daily stress either during the whole period of 60 days (crowding: C-E) or only during the first 30 days (crowding: F-H; heat shock: I-L). (L-P) Heart sections at 30 dpci after AFOG staining. In this case, zebrafish were exposed to daily crowding either during the first 2 weeks (L-N) or during the final 2 weeks (O, P). (L-N) The cryoinjured hearts of fish that were stressed for the first 2 weeks followed by 2 weeks of recovery at normal conditions display impaired regeneration. 78% of the animals displayed a partial or complete blockage of heart regeneration. (O, P) The cryoinjured hearts of fish that were kept at normal conditions for the first 2 weeks and then exposed to stress for the final 2 weeks, regenerated the hearts. The analysis of the regenerative process in the different groups revealed that daily exposure to stress has a stronger impact on the initial phase of heart regeneration. (Q) Analysis of the body weights before cryoinjuries and at 30 dpci revealed no significant weight changes during this period in control and stressed animals. (R) Glucose measurements revealed higher levels of blood glucose in the control animals as compared to stressed fish at 30 dpci. N≥8. No difference was observed at 7 dpci. Data are represented as mean ± SEM. \*\*P < 0.01; N=5. Scale bar (A) = 100 μm.

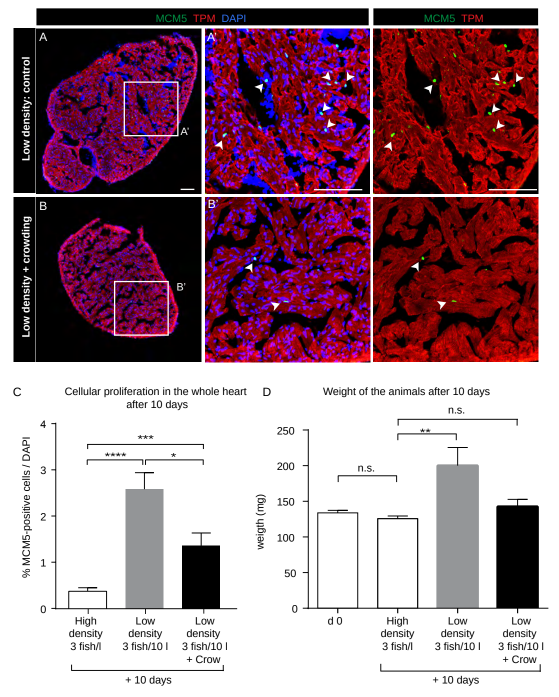


Figure S2.

Rapid cardiac growth is affected by daily acute stress exposure.

(A, B) Sections of hearts of control and stressed zebrafish 10 days after the transfer to low density conditions (3 fish / 10 liters) stained with antibodies against Tropomyosin (TPM, red), MCM5, a G1/S-phase marker (green) and with DAPI (blue). Juvenile zebrafish (2 months of age) were exposed 2 x per day to crowding (10 fish/ 250ml during 1h) during 10 days to investigate the effect of stress on cardiac homeostatic growth. (A', B') Higher magnifications of the framed area shown in (A, B). A higher number of proliferating cardiac cells (CMs and non CMs) were identified in the control fish when compared to the stressed animals (arrows). (C) Bar graphs show the percentage of proliferating cells (MCM5-positive DAPI-positive / DAPI-positive in the tropomyosin-labeled myocardium) in the animals maintained at high density (5 fish per liter) and in control and stressed animals transferred to low density conditions (3 fish in 10 liters). Ventricles exposed to daily stress (10 fish/ 250 ml, 2 x 1h/day) showed a significantly lower proportion of proliferating cardiac cells than control hearts. (D) Bar graphs show the average weight of the animals at day 0, after 10 days in high density conditions (3 fish in 1 liter), after 10 days in low density conditions (3 fish in 10 liters) and after 10 days in low density conditions with daily exposure to crowding (10 fish/ 250 ml, 2 x 1h/day). Data are represented as mean ± SEM. \*P < 0.05, \*\*P < 0.01 \*\*\*P < 0.001, \*\*\*\*P < 0.0001; N≥4.

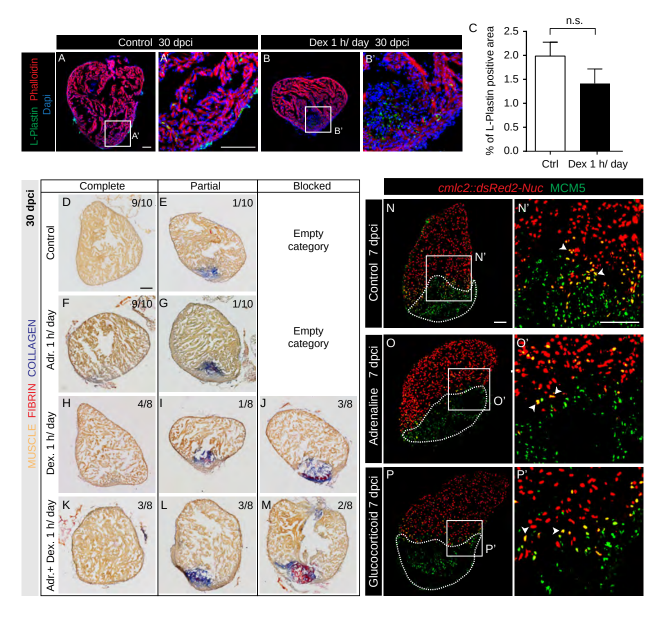


Figure S3.

The concomitant administration of dexamethasone and adrenaline after cryoinjury mimics the effect of stress on heart regeneration.

(A, B) Representative images of cryoinjured hearts at 30 dpci labeled with phalloidin (red, F-actin), dapi (blue) and antibody against L-Plastin (green, leukocytes). (A', B') Higher magnification of the frame areas shown in (A, B). (C) Quantification of L-Plastin-positive area normalized to the total area of the ventricles revealed no significant effect of daily acute dexamethasone (2 mg/l) treatment on cardiac inflammation. N > 4. (D-M) Sections of hearts at 30 days post cryoinjury (dpci) after Aninlin blue acid Fuchsin Orange-G (AFOG). (F-M) The acute administration (1 hour per day) of dexamethasone (Dex, 2 mg/l) alone (H-J) or concomitantly with adrenaline (adr, 1 mg/l) (K-M) resulted in cardiac regenerative impairment in 50% and 62.5% of the fish, respectively. In contrast, animals treated with adrenaline alone during 1 hour per day displayed similar regenerative scores as the control (D-G). (N-P) Heart sections of cmlc2::dsRed2-Nuc transgenic zebrafish at 7 dpci immunostained against MCM5 (green). (N'-P') Higher magnification of the framed areas shown in (N-P). Proliferating CMs could be identified in all groups by the overlap between MCM5 and DsRed (arrowheads). In contrast to adrenaline (1 mg/l, 1h /day), the treatment with glucocorticoid (hydrocortisone, 1 mg/l) lead to a reduction in CM proliferation at 7 dpci. Cryoinjured parts are encircled with a dashed line. Scale bar (A, D, N) = 100 um.

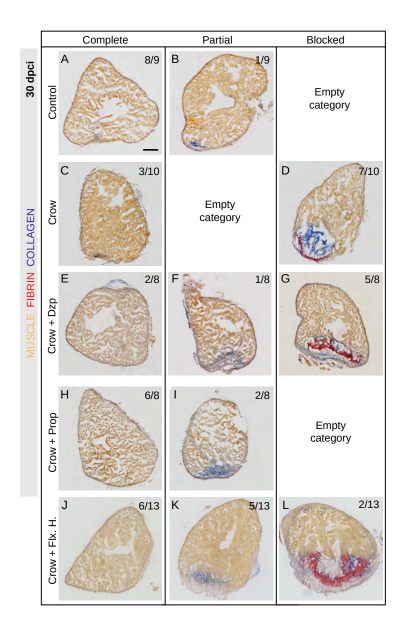


Figure S4.

Propranolol and fluoxetine hydrochloride administration have a beneficial effect on cardiac regeneration in the stressed animals.

(A-L) Heart sections at 30 dpci after AFOG staining. (E-L) The treatment of the stressed animals with propranolol (1 mg/l, 1 h/day, 3 days pretreatment, H, I) and fluoxetine hydrochloride (100  $\mu$ g /l, continuous, 2 weeks pretreatment, J-L) had a significant rescue effect on the stress-induced regenerative impairment whereas the administration of diazepam (1 mg/l, continuous, 3 days pretreatment, E-G) did not rescue the negative effect of stress on heart regeneration. Scale bar (A) = 100  $\mu$ m.

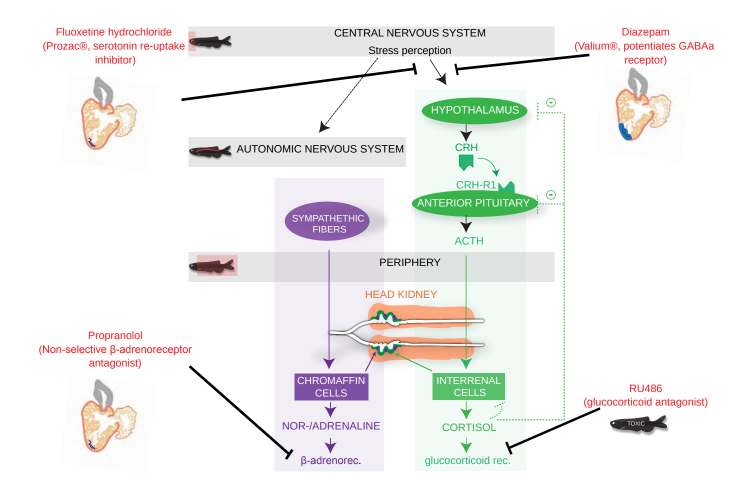


Figure S5.

Modulation of the stress response: effect on cardiac regeneration.

In zebrafish, exposure to different stressors activates two main axes: (1) the hypothalamus-pituitary interrenal (HPI) axis (green) and (2) the sympathetic-chromaffin cell axis (purple).

The treatment of the stressed animals after cryoinjury with fluoxetine hydrochloride (serotonin re-uptake inhibitor) and propranolol (non-selective  $\beta$ -adrenoreceptor antagonist) had a significant rescue effect on the stress-induced regenerative impairment. The administration of diazepam (GABAa receptor enhancer) did not show any beneficial effect on scar resorption. The intermittent administration of RU486 (glucocorticoid antagonist) did not rescue cardiomyocyte proliferation in the stressed animals and the prolonged treatment with this drug was toxic and resulted in a high lethality.

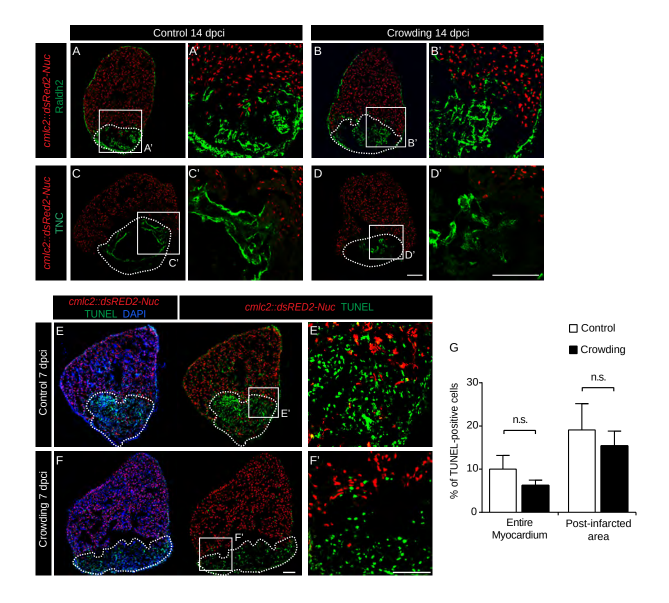


Figure S6.

Epi/endocardium activation, Tenascin C expression and cellular apoptosis are not affected by daily exposure to stress.

(A-D) Sections of *cmlc2::dsRed2-Nuc* hearts of control and stressed zebrafish at 14 dpci immunostained against the retinoic acid synthetizing enzyme (Raldh2, green), which is a marker of the activated epi-endocardium (A, B) or against Tenascin C (TNC, green), which is an ECM protein involved in tissue remodeling (C, D). (A'-D') Higher magnifications of the framed areas shown in (A-D). Expression of the Raldh2 enzyme and Tenascin C could be detected at the injured site and in the epicardium both in control (A', C') and stressed (B', D') fish. This indicates no major impact of daily stress on epicardial/endocardial activation or on Tenascin C regulation. Scale bar (D, D') = 100 um.

(**E**, **F**) Sections of *cmlc2::dsRed2-Nuc* hearts at 7 dpci after TUNEL assay and DAPI staining. (**E**', **F**') Apoptotic cells were detected by signal overlap between TUNEL and DAPI in the cryoinjured parts but also in the rest of the ventricle. (**G**) Quantification of TUNEL and DAPI positive cells allowed to estimate the level of cellular apoptosis in the ventricle of the control and crowded zebrafish. No significant difference in apoptosis was observed between control and stressed fish at 7 dpci. Cryoinjured parts are encircled with a dashed line. Data are represented as mean  $\pm$  SEM. Scale bar (F, F ') = 100  $\mu$ m.

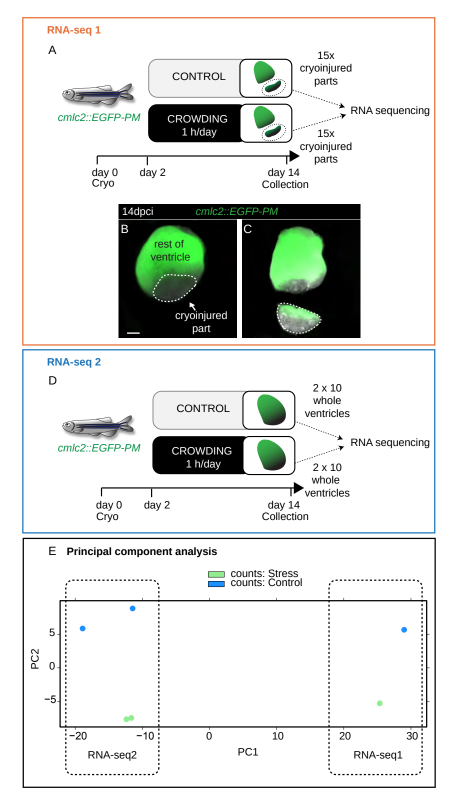


Figure S7.

Experimental setup for the RNA-seq1 & 2 experiments and principal components analysis of both experiments.

(A) Experimental setup for the RNA sequencing 1 performed at 14 dpci with the RNA extracted from the cryoinjured parts of the stressed and control hearts. (B, C) The use of *cmlc2::EGFP-PM* transgenic zebrafish, expressing GFP in the plasma membrane of CMs, enabled to identify and isolate the cryoinjured parts of the ventricle used for the RNA-sequencing experiment (15 cryoinjured parts were pulled together for each group in order to get sufficient amount of RNA). (D) Experimental setup for the RNA sequencing 2 performed at 14 dpci with the RNA extracted from the wholes ventricles of the stressed and control hearts. (E) Principal component analysis plot performed with RNA-seq1 and RNA-seq2. The Principal Compenent Analysis (PCA) is a classical method of statistical analysis. By transforming the set of observations into linearly uncorrelated variables called "principal component". By definition the PCs are ordered by the variance they can explain in the data. PC1 explains the largest variance, then PC2, then PC3 etc. In our experiment, PC1 allows discriminating the two independant RNAseq experiments, and PC2 discriminates the stress and the control samples in both RNAseq experiments. Scale bar (B)=  $100 \mu m$ .

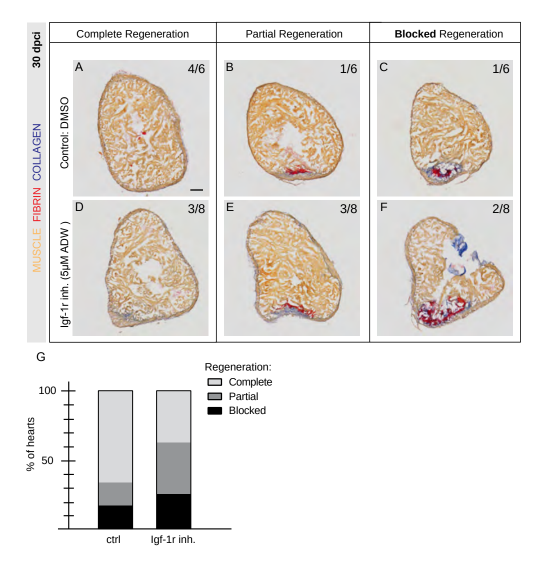


Figure S8. Inhibition of IGF-1 receptor impairs heart regeneration.

(A-F) Heart sections at 30 dpci after AFOG staining. (A-C) In the control group treated with 0.05% DMSO, 66% of the zebrafish completely regenerated the heart at 30 dpci. (D-F) In the group treated with NVP-ADW742 (5  $\mu$ M), a specific inhibitor of the lgf1r kinase activity, 62.5% of the fish showed impaired cardiac regeneration at 30 dpci. (G) Histograms represent the percentage of zebrafish hearts with complete (white), partial (gray) or blocked (black) regeneration at distinct experimental settings (regenerative scores). Scale bar (A) = 100  $\mu$ m.

Name	# of Entities	Expanded # of Entities	# of Measured Entities	Median change		
Proteins Involved in Pathogenesis of Melanoma	245	262	180	1.121233899	5.5849E-06 Disease Collections	
Built Pathway_IGFBP1b_downstream cell processes	178	763	238	1.086427761	2.7121E-05 Private pathways	
Proteins Involved in Pathogenesis of Glioma	304	340	207	1.083054452	2.8003E-05 Disease Collections	
Defective Clearance of Apoptotic Keratinocytes in Systemic Lupus Erythematos	79	245	89	1.071944531	2.9092E-05 Disease Collections	
BMP7-ACVR2 Expression Targets	27	29	21	1.610822919	0.00026135 Expression Targets Pathways	
TGFB1-ACVRL1 Expression Targets	221	233	138	1.175402723	0.00028927 Expression Targets Pathways	
EGF/CTNN Expression Targets	143	156	96	1.279815128	0.00032076 Expression Targets Pathways	
B Cell Activation	62	841	506	1.181790244	0.00032227 Cell Signaling	
TLR4/NF-kB/IRF Expression Targets	70	76	37		0.00034869 Expression Targets Pathways	
Pathway_Genes_Imp	954	954	628		0.00037041 Private pathways	
B-cell Chronic Lymphocytic Leukemia Overview	122	430	273			
Extracellular Matrix Turnover	36	166	76		0.00039486 Cell Process Pathways	
Melanoma Overview	176	627	429		0.00043896 Disease Collections	
EphrinR -> actin signaling	15	216	141		0.00051062 Receptor Signaling	
Dystrophin Glycoprotein Complex Signaling in Duchenne Muscular Dystrophy	67	792	490		0.00054743 Disease Collections	
OXIDATIVE(ROS)Dystrophin Glycoprotein Complex Signaling in Duchenne Mus		792	490		0.00054743 Private pathways	
Proteins Involved in Pathogenesis of Glioblastoma	188	210	138		0.00055643 Disease Collections	
Hodgkin Lymphoma Overview	146	614	285		0.00062251 Disease Collections	
TLR4 -> IRF signaling	14	14	10		0.0007392 Receptor Signaling	
TGFB1-TGFBR1 Expression Targets	89	91	47		0.00076762 Expression Targets Pathways	
Proteins Involved in Pathogenesis of Cataract	95	95	53		0.00099064 Disease Collections	
Role of Hexosamine Pathway in Diabetic Microangiopathy	26	145	71	1.1450458		
T Cell Activation	80	948	471		0.00101336 Cell Signaling	
PDGF/STAT Expression Targets	80	89	57		0.00102698 Expression Targets Pathways	
Proposed Mechanisms of Antiepileptic Effects of a Ketogenic Diet	56	365	219		0.00108017 Disease Collections	
Atlas of Signaling	381	6035	3424		0.00113812 Cell Signaling	
Insulin/CEBPA/CTNNB/FOXA/FOXO Expression Targets	145	171	109		0.00173612 Cell Signaling 0.00129622 Private pathways	
Insulin/CEBPA/CTNNB/FOXA/FOXO Expression Targets	145	171	109		0.00129622 Expression Targets Pathways	
Notch_Insulin/CEBPA/CTNNB/FOXA/FOXO Expression Targets	145	171	109		0.00129622 Expression rargets Fatriways 0.00129622 Private pathways	
Cell Cycle Regulation	135	2176	1436			
Common Non-genomic Effects of Thyroid Hormones	60	338	1430		0.00135125 Cell Signaling	
,					0.00140719 Disease Collections	
BMP2 Activates WNT Signaling in Pulmonary Artery Smooth Muscle Cells	26	253	163 77		0.00154104 Disease Collections	
TGFB1-TGFBR2 Expression Targets	116	125			0.00157163 Expression Targets Pathways	
Actionyosin-Based Movement	25	85	31		0.00160695 Cell Process Pathways	
Actin Cytoskeleton Regulation	51	546	369		0.00179848 Cell Signaling	
Ca2+ Overload in Duchenne Muscular Dystrophy	59	256	150		0.00184601 Disease Collections	
Role of HMGB1 and IL1B in Neuroinflammation in Epilepsy	25	29	19		0.00186536 Disease Collections	
TNF/NF-kB Expression Targets	127	135	68		0.00192162 Expression Targets Pathways	
INHBA-ACVR2/ACVR1 Expression Targets	25	27	20		0.00193913 Expression Targets Pathways	
T-cell Receptor Signaling	71	371	203		0.00194978 Immunological Pathways	
Onset of Atopic Dermatitis	72		203		0.00194978 Disease Collections	
IL1B Expression Targets	169	195	109		0.00196439 Expression Targets Pathways	
Vascularization in Hepatocellular Carcinoma	25	101	52		0.00203218 Disease Collections	
Secondary Glioblastoma	66	343	237		0.00204329 Disease Collections	
ROS metabolism	45	108	16		0.00217801 Metabolic Pathways	
EGF/MEF/MYOD/NFATC Expression Targets	145	231	138		0.00222991 Expression Targets Pathways	
Leptin/STAT Expression Targets	96	107	63		0.00239132 Expression Targets Pathways	
AGT/CREB Expression Targets	116	193	110		0.00244381 Expression Targets Pathways	
Vitamin A (retinol) metabolism and visual cycle	93	165	19		0.00261586 Metabolic Pathways	
Sister Chromatid Cohesion	27	174	113	1.317461315	0.00276692 Cell Process Pathways	

Table S1.

Gene set enrichment analysis for the the fused RNA seq experiment (RNA-seq1&2) to highlight changes in gene expression between control and stressed zebrafish hearts. The table shows the 50 most significant pathway gene sets for the fused RNA-seq experiment: RNA-seq1 & RNA-seq2. The analysis was performed with Pathway Studio.

Name	# of Entities	Expanded # of Entities	# of Measured Entities	Median chan	p-value	Hit type
SRP-dependent cotranslational protein targeting to membrane	108	108	74	1.77234506	3.0145E-10	biological_process
viral transcription	82	82	56	1.84158913	4.4373E-10	biological_process
translational termination	89	89	62	1.7790759	1.112E-08	biological_process
viral life cycle	93	93	63	1.77234506	4.2407E-08	biological_process
translational elongation	149	149	74	1.76277199	3.1223E-07	biological_process
cell adhesion	658	658	342	1.03973094	1.4525E-06	biological_process
response to virus	145	145	68	1.37681504	1.67E-06	biological_process
cytosolic large ribosomal subunit	68	68	33	1.7790759	1.7438E-06	cellular_component
actin binding	395	395	223	-1.0781643	1.789E-06	molecular_function
Z disc	124	124	71	1.0368129	4.4501E-06	cellular_component
extracellular region	2319	2319	793	1.10562778	1.0497E-05	cellular_component
cytosol	2752	2752	1886	1.1016447	1.3491E-05	cellular_component
homophilic cell adhesion	160	160	47			biological_process
nuclear-transcribed mRNA catabolic process, nonsense-medi	122	122	86			biological_process
extracellular matrix	250	250	154			cellular_component
cytosolic small ribosomal subunit	59	59	28			cellular_component
proteinaceous extracellular matrix	318	318	167			cellular_component
DNA strand elongation involved in DNA replication	31	31	21			biological_process
cytokine binding	22	22	11			molecular function
structural constituent of muscle	52	52				molecular function
innate immune response	669	669	341			biological process
structural constituent of ribosome	416	416	112			molecular function
cellular response to exogenous dsRNA	14	14	6			biological_process
myosin complex	59	59				cellular_component
heparin binding	166	166	75			molecular_function
collagen binding	61	61	42			molecular_function
inositol phosphate-mediated signaling	12	12	.2			biological_process
transmembrane signaling receptor activity	178	178	71			molecular_function
muscle filament sliding	39	39	24			biological_process
positive regulation of neutrophil chemotaxis	19	19	11			biological_process
translational initiation	149	149	100			biological_process
actin filament binding	97	97				molecular function
response to vitamin D	27	27				biological_process
protein homodimerization activity	765	765				molecular function
RNA metabolic process	250	250	195			biological_process
cellular response to mechanical stimulus	86	86				biological process
mRNA metabolic process	227	227	180			biological process
mitotic cell cycle	394	394	300			biological_process
cardiac muscle cell action potential involved in contraction		10				biological process
positive regulation of potassium ion transport	16	16	9			biological_process
myoblast migration	5	5	5			biological_process
superoxide anion generation	15	15	8			biological_process
immune response	449	450	119			biological_process
regulation of stress fiber assembly	7	7				biological_process
cellular amino acid metabolic process	50	50				biological_process
· ·	320	320	32 243			-
viral process	320 48	48	243			biological_process
cell periphery	48 174	48 174				cellular_component
regulation of small GTPase mediated signal transduction	48	174				biological_process
small ribosomal subunit			18			cellular_component
energy reserve metabolic process	103	103	68	-1.11351284	0.00129081	biological_process

**Table S2.** Gene set enrichment analysis for the fused RNA seq experiment (RNA-seq1&2) to highlight changes in gene expression between control and stressed zebrafish hearts. The table shows the 50 most significant **G0-term** gene sets for the fused RNA-seq experiment: RNA-seq1 & RNA-seq2. The analysis was performed with Pathway Studio.

ganalD	log2EC fused	padj_fused_type of RNA
geneID IFITM1	3.742286001	
ENSDARG00000084533	3.881034115	
ENSDARG00000081938	8.047716049	
SLC25A4	-2.760180063	1.18E-21
ENSDARG00000088865	3.443302383	1.18E-21 miRNA
LGALS3BP	3.113514297	
CORO2A	-5.652379587	
ENSDARG00000088976 ENSDARG00000086686	3.582835203	
FHL2	3.47104141 2.929914201	
ENSDARG00000089384	4.183491967	
ENSDARG00000070212	3.256322883	
ENSDARG00000090280	3.435020079	
ENSDARG00000091738	3.373336585	6.97E-18 miRNA
PARP6	-2.569819142	1.76E-17
MYBPH	-2.572453077	
ENSDARG00000093902	7.612314957	. 0
PCOLCE	2.707511492	
GRIN3A	-4.873753905	
ENSDARG00000087783	3.665121865 3.429343718	
ENSDARG00000088564 IFI27	2.641037412	
ADCY6	-2.924964698	
CPAMD8	3.645294968	
ENSDARG00000089068	7.09582739	
MIH7B	-2.872828154	
ENSDARG00000090619	2.873602128	3.78E-13 miRNA
ENSDARG00000086606	7.03980247	3.78E-13 pseudogene
ENSDARG00000086256	-2.733709962	4.41E-13 protein_coding
CRIP1	2.06748148	
GNB3	-5.560717107	
ENSDARG00000086396	2.928302686	
SEC61G ENSDARG00000086085	2.213266624 3.124629975	
IFI27L1	2.450376839	
ENSDARG00000088436	2.770051212	
ENSDARG00000078859	4.727957055	
MYRFL	-2.53518149	. – •
ENSDARG00000087953	3.056798055	1.97E-12 miRNA
ENSDARG00000089382	2.864534502	. – 3
PDZRN3	-2.361487556	
SLC16A1	-2.31344781	
BMP10	-2.566374785	
ENSDARG00000096403 ENSDARG00000070873	2.7441639 2.558192575	
ENSDARG00000070073	2.774640053	. – •
ENSDARG00000087337	2.801632112	
HEPACAM2	6.641683192	
ENSDARG00000017246	-2.401867587	
ENSDARG00000090146	2.984329301	4.74E-11 miRNA
RBP4	2.002717065	
ENSDARG00000090175	2.882947206	
ENSDARG00000082789	2.100222947	
ENSDARG00000086192 KCNMB2	3.098469469 -2.581176089	
ENSDARG00000015815	-1.809270076	
NR4A1	-2.146381259	
EPHB3	-2.249748285	3.19E-10
SULT2B1	2.547289475	
ENSDARG00000058348	2.976849528	
ENSDARG00000088311	2.795463059	
ETV4	2.433975283	
ENSDARG00000088838	2.703044112	
ENSDARG00000085168	6.285124238	
ENSDARG00000097100 ENSDARG00000087747	-2.044045148 6.227650516	. – •
ENSDARG00000087747	6.225175014	
LGR4	-2.076095077	
	=::::3000011	

**Table S3.** Table showing the 50 most significant DE **non-coding RNAs**(miRNAs are highlighted in red) for the fused RNA-seq experiment: RNAseq1 & RNA-seq2. The analysis was performed with Pathway Studio.