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1	Sub-lethal UV radiation during early life stages alters the
2	behaviour, heart rate and oxidative stress parameters in
3	zebrafish (<i>Danio rerio</i>)
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18 Abstract

19 Environmental UV radiation in sufficient doses, as a possible consequence of climate change, is 20 potent enough to affect living organisms with different outcomes, depending on the exposure life 21 stage. The aim of this project was to evaluate the potentially toxic effects of exposure to sub-lethal 22 and environmentally relevant doses of UVA (9.4, 18. 7, 37.7 J/cm²) and UVB radiation (0.013, 0.025, 0.076 J/cm²) on the development and behaviour in early life stages (4.5 – 5.5 hours post 23 fertilization, hpf) of the zebrafish (Danio rerio). The used doses were all below the median lethal 24 dose (LD₅₀) and caused no significant difference in survival, deformities, or hatching between 25 exposed and control groups. Compared to controls, there were transient UVA and UVB exposure 26 27 effects on heart rate, with dose dependent reductions at 50 hpf, and at 60 hpf for UVA only. The UVB exposure caused an increasing trend in reactive oxygen species (ROS) formation at the two 28 highest doses, even though only significant at 120 hpf for the second highest dose. Both UVA and 29 30 UVB caused an increasing trend in lipid peroxidation (LPO) at the highest doses tested at 72 hpf. 31 Furthermore, UVA exposure led to significant reductions in larval movement following exposure 32 to the two highest doses of UVA, i.e., reduction in the time spent active and the total distance 33 moved compared to control at 100 hpf, while no effect on the swimming speed was observed. The 34 lowest dose of UVA had no effect on behaviour. In contrast, the highest dose of UVB led to a possible increase in the time spent active and a slower average swimming speed although these 35 effects were not significant (p = 0.07). The obtained results show that UV doses below LD₅₀ levels 36 are able to cause changes in the behaviour and physiological parameters of zebrafish larvae, as 37 38 well as oxidative stress in the form of ROS formation and LPO. Further testing is necessary to assess how this type of radiation and the effects observed could affect fish population dynamics. 39

40 Key words: zebrafish; UV; locomotor; heart rate; ROS; lipid peroxidation

41 **1 Introduction**

42 Ultraviolet light is ubiquitously present in the environment and classified into three categories: 43 UVA (400–315 nm), UVB (315–280 nm), and UVC (280–100 nm), which is absorbed by the ozone layer and does not occur as part of the solar spectrum reaching the troposphere. The 44 depletion of the ozone layer and climate change together are increasing the exposure of aquatic 45 organisms to increasing levels of UVB and UVA radiation (Bais et al., 2018). It has been proposed 46 that exposure to an altered UV regime can potentially cause differences in behavioural responses 47 and possibly influence the level of biodiversity and food web dynamics in aquatic ecosystems 48 (Bais et al., 2018). 49

Most studied aquatic organisms, particularly those inhabiting shallow aquatic environments, show 50 51 susceptibility to the detrimental effects of UV radiation exposure (Häder et al., 2007). In general, 52 it has been reported that fish spawning in shallow waters are most susceptible to the biologically damaging effects of UV radiation due to exposure of the vulnerable early larval stages, at a time 53 when extensive DNA replication and organogenesis is taking place (Béland et al. 1999, Hunter et 54 55 al., 1979). In sufficient doses (i.e. a longer exposure time), UV radiation can impair embryonic development in fish (Andrade et al., 2016; Fujimoto et al., 2007), and additionally it was found 56 57 that zebrafish embryos at the gastrulation stage (starting from 5.25 hours post fertilization (hpf)), 58 were more tolerant to UV radiation compared to later developmental stages (Dong et al., 2007). Further, it was shown that even UVC radiation, at a wavelength outside the solar spectrum could 59 inflict severe biological damage, whereby hindering the embryonic development in zebrafish 60 (Danio rerio) via impairment of epiboly in the earliest post-zygotic stages (Strähle and Jesuthasan, 61 1993). 62

Moreover, UV radiation in sufficient doses can initiate a series of redox reactions to generate reactive oxygen species (ROS), which cause oxidative stress to cells and tissues during irradiation, but also as a result of disturbed cellular metabolic processes (Stańczyk et al., 2005). Although the exposure effects on oxidative stress parameters in larval fish have been reported after chronic UV irradiation during several days (Lesser et al., 2000; Mekkawy et al., 2010), it is less known whether these effects are persisting at later developmental stages.

In addition to the potential of UV radiation to induce oxidative stress, previous studies have shown 69 70 that UVB exposure also caused differences in physiological and behavioural responses in fish 71 larvae (Icoglu Aksakal and Ciltas, 2018), which are key life fitness traits essential for the growth and survival. Alterations in these responses would have severe consequences for the survival of 72 these vulnerable early life-history stages. For example, an impairment of avoidance behaviour was 73 demonstrated after exposure to environmentally relevant doses of UVB in cod (Gadus morhua) 74 larvae (Fukunishi et al. 2012). In an earlier study, Alemanni et al., (2003) investigated the 75 76 neurobehavioural effects of UVB exposure in juvenile rainbow trout (Oncorhynchus mykiss). These authors observed that irradiation with UVB from fluorescent tubes irreversibly increased 77 trout O₂-consumption by individual fish. Further, rapid tail and fin movement as well as rapid and 78 79 erratic displacements were observed at doses that caused changes in the O_2 consumption. In another study, Häkkinen et al. (2004) reported that exposure of newly fertilized pike (*Esox Lucius*) 80 81 eggs to UVB-doses similar to one daily erythema weighted ambient dose in Finland in May (0.27 82 J/cm², solar radiant exposure weighted by an action spectrum), resulted in neurobehavioural 83 disorders such as inability to swim straight, circular movement and eventual mortality. However, to date insufficient data is available on the potential persistence of deleterious effects of UV 84 irradiation during early life stages prior to hatching in fish. 85

The objective of this study was to investigate whether zebrafish sub-lethal UVA and UVB 86 exposures during a vulnerable early life stage can cause persisting changes in physiological, 87 88 oxidative stress parameters and lead to locomotor behavioural changes later in life. For this purpose, the zebrafish was selected as a model organism as it is a well-known model for 89 developmental and behavioural toxicity assessment following environmental toxicant exposures 90 91 (Ton et al., 2006; Parng et al., 2007; Selderschlaghs et al 2010; Colwill and Creton, 2011; Tierney et al., 2011). The doses used in this study correspond to a typical mid-summer, midday and clear 92 93 sky average outdoor exposure in Oslo (60 °N) of 10 and 150 min of UVB and UVA, respectively. 94 Zebrafish from the late blastula to early gastrula stages (4.5-5.5 hpf), when the cell fate specification onset takes place (Kimmel et al., 1995; Montero et al., 2005) were used for the 95 exposure studies. In addition to changes in larval behaviour, changes in heart rate as well as 96 changes in oxidative stress were assessed. 97

98

99 2 Materials and methods

100 **2.1 Fish husbandry**

The study was performed at The Norwegian Zebrafish Platform of the Norwegian University of 101 Life Sciences, Oslo, Norway. The unit is licensed by the Norwegian Animal Research Authority 102 103 (NARA) (www.mattilsynet.no) and accredited by the Association for Assessment and Accreditation of Laboratory Animal Care (www.aaalac.org). The study was carried out under the 104 regulations approved by the unit's animal ethics committee (Institutional Animal Care and Use 105 106 Committee/IACUC) following Norwegian laws and regulations controlling experiments and procedures on live animals in Norway. AB wild-type zebrafish were maintained at 28°C under a 107 108 14:10 light/dark photoperiod. Adult care and breeding was in accordance with the local protocols

previously described in Hurem et al. (2017). To generate embryos, adults were placed in spawning tanks in the afternoon, and the fish were spawned following the cessation of light (08:00) the next day, and the embryos collected (09:00) and maintained in sterile embryo media (60 μ g/mL Instant Ocean® sea salts) until the time of exposure.

113 **2.2. Ethical statement**

All animal experiments in this study were performed in accordance with the Norwegian Animal Protection Act (implemented EU Directive 2010/63/EU) and larvae were euthanized at 120 hpf using an overdose of Tricaine (MS-222, Sigma Aldrich), followed by rapid freezing at (-70°C).

117 2

2.3 UV exposure and embryo toxicity

Embryos between the late blastula (4.5 hpf) and early gastrula (5.5 hpf) stage of development were 118 119 used for the UVA and UVB exposures (Table 1). All exposures were performed in polystyrene 50 x 9 mm Petri dishes (VWR, Radnor, PA, U.S) without the lid with 10 embryos in a 1 mL volume. 120 Radiation exposure was performed using a modified exposure unit (Polylux PT, Dreve-Dentamid, 121 Unna, Germany) consisting of three 9 W PL 12 UVB lamps (Philips, Eindhoven, The Netherlands) 122 or three UVA-lamps, Osram GmbH DULUX S BL UVA 9 W/78. In order to remove UV with 123 124 shorter wavelengths than 280 nm a filter material consisting of 5 mm Poly-Methyl-Methacrylate (Atoglas, Altuglas International) was placed in front of the exposure unit. The transmission of the 125 filter was 100 % for wavelengths above 300 nm. During irradiance measurement of the UVB-126 127 lamps, the filter was placed between the lamp and the detector to account for any absorption or light scatter in the material. The samples to be irradiated with UVB were placed 10 cm from the 128 129 exposure unit. The UVA irradiation was performed with two exposure units placed on top of each 130 other in a "sandwich" configuration with Petri dishes placed on a plate made of Atoglas in the gap between the exposure units. Thereby the dishes transparent to UVA were irradiated from both 131

sides. The irradiance at the level of the dishes was estimated by adding the upward and downward
fluxes. The spectrum and irradiance were determined by a scanning spectral radiometer (Bentham,
UK, DTM 300 with a fibre optic light guide and cosine adapted diffuser D7). Constancy of the
irradiance values was routinely performed with a Solar Light Co, PMA2100 (Philadelphia, USA)
radiometer with appropriate detectors. The irradiance levels were 10.4 mW/cm² and 0.42 mW/cm²
in UVA and UVB, respectively. The controls for UVB and UVA embryos were kept at room
temperature (22°C) during irradiation.

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Table 1. Doses for zebrafish UVA and UVB exposure experiments, group denotations andcomparison to LD₅₀.

UVA exp	posure, 10.4 m	nW/cm ²	UVB exposure, 0.42 mW/cm ²					
Group	Exposure	Dose (J/cm ²),	Group	Exposure	Dose (J/cm^2) ,			
	time (s)	(approx. % of LD ₅₀)		time (s)	(approx. % of LD ₅₀)			
Control	0	0	Control	0	0, 0			
UVA 1	900	9.4, (17 %)	UVB 1	30	0.013, (13 %)			
UVA 2	1800	18.7, (34 %)	UVB 2	60	0.025, (25 %)			
UVA 3	3600	37.4, (68 %)	UVB 3	180	0.076, (76 %)			

In order to determine the LD₅₀, 40 embryos distributed in 4 wells (10 embryos/ well) of a 12-well plate (NuncTM, Thermo-Fischer Scientific) were irradiated at approximately 5 hpf over the whole dose range. The number surviving a certain dose was scored at 48 hpf and expressed as surviving fraction relative to an unexposed control. The LD₅₀ was found by linear extrapolation of data from 4-5 independent experiments (Table 1A). The subsequent lower UVA and UVB doses for the

behaviour studies were chosen from the LD₅₀ estimation. In order to determine the toxic effects of 148 the used lower doses of UVA and UVB radiation exposure on the survival and development of the 149 150 embryos and larva, including the LD_{50} , the zebrafish embryo toxicity test (OECD, 2013) was applied. Following exposure, embryos were incubated in 96 well plates (Nunc[™], Thermo-Fischer 151 Scientific) until 96 hpf. The survival, occurrence of deformities, and the median hatching time 152 153 (HT_{50}) were assessed in embryonic and larval zebrafish exposed to doses lower than the LD₅₀ presented in Table 1. In addition, body length was assessed at 72 hpf using a stereomicroscope in 154 155 20-30 replicate larvae without deformities per each exposure dose.

156 2.4 Heart rate

In order to determine the effects of UVA and UVB radiation exposure on the metabolism, the heart rate was assessed at 50 and 60 hpf using a light microscope and counted as the number of heart beats in a 15 sec period. Eight to ten larvae/group were scored for each biological replicate (n = 38-53/group). For UVB, one biological replicate was missing, therefore an additional 8-10 larvae were analyzed within the subsequent biological replicate.

162 **2.5 Oxidative stress**

163 **2.5.1 ROS formation**

Intracellular ROS production was determined in zebrafish after UV irradiation using the fluorescent probe 2',7'-dichlorodihydrofluorescein diacetate (H₂DCFDA, Invitrogen, Molecular Probes Inc., Eugene, OR, USA) and according to the method described in Hurem et al., (2017). Briefly, embryos were individually collected and incubated in a 96-well black microplate (Corning Costar, Cambridge, MA, USA) for 1.5 hours with H₂DCFDA, with 20-24 replicate embryos per exposure group at 70 hpf. Fluorescence was recorded at approximately 72 and 120 hpf in mean relative fluorescence units (RFU) using the Cytation 3 Cell Imaging Multi-Mode Reader (Biotek, Winooski Vermont, USA) and analyzed using Gen5 Microplate Reader and Imager Software (Biotek, Winooski Vermont, USA). Natural fluorescence of irradiated egg water in combination with the probes (without presence of embryos) for each dose rate and the resulting fluorescence subtracted, including a positive control (1% H202) were also analysed. The relative fluorescence obtained for each exposure group was expressed as fold induction comparative to the control.

176 2.5.2 Lipid peroxidation

The lipid peroxidation was assessed by two methods. First, the probe C11-BODIPY^{581/591} was used 177 for measuring LPO in zebrafish larvae in a time-dependent manner. This probe is a fatty acid 178 179 analogue with specific fluorescence properties, which can easily enter the lipid bilayer and be subject to oxidation by oxyl-radicals together with the endogenous fatty acids, once inside the 180 cellular membrane (Drummen et al., 2002). Similarly to ROS formation, exposed embryos and 181 controls were individually collected and incubated in a 96-well black microplate (Corning Costar, 182 Cambridge, MA, USA) for 2 hours with C11-BODIPY^{581/591} (final concentration 10 µM), with 20-183 23 replicate embryos per exposure group at 70 hpf. Fluorescence was recorded by use of the same 184 system as for ROS at 72, 96 and prior to 120 hpf and the results expressed as fold induction 185 comparative to the control. 186

Lipid peroxidation was also determined in 72 hpf larvae by measurement of malondialdehyde (MDA) and 4-hydroxyalkenals (4-HNE) concentrations upon decomposition by polyunsaturated fatty acid peroxides, following the method by Erdelmeier et al. (1998), previously described in zebrafish larvae (Hurem et al., 2017). Here, 15 zebrafish larvae were pooled per sample in triplicate biological samples per dose, with exception of duplicates in UVA2 and UVB3 (where an additional technical replicate was used).

193 **2.6 Behavioural testing**

For the behavioural analyses, immediately following exposure, individual embryos were placed in 194 the wells of square 96 well plates (#7701-1651, Whatman, USA) with 500µL of media and placed 195 inside an incubator set to 28°C with a 14:10 day/night cycle. UVA and UVB treated embryos were 196 transferred to separate plates. For the first three biological replicates, the control and all 3 UVA 197 198 and UVB doses were equally represented across two 96 well plates (n = 24/dose/plate), whereas 199 for the final three biological replicates the control and all 3 UVA and UVB doses were equally represented across only one 96 well plate (n = 24/dose). The locomotor activity (LMR) of the 200 201 larvae (total 139-143 larvae from 6 experiments) was visualized over a set time interval. Behavioural tests were conducted using a ViewPoint® Zebrabox system and the accompanying 202 video tracking software (ViewPoint Life Sciences, Lyon, France), which is a high-throughput 203 204 image analysis system that can visualize and quantify the zebrafish behavioural response. Behavioural screening was undertaken at 100 hpf. This corresponds to tests beginning 330 minutes 205 206 (13:00) and 390 (14:00) minutes after the cessation of light (07:30) in the incubator for UVA and UVB, respectively. Larval behaviour, including the cumulative distance travelled and the time 207 spent active per minute, were simultaneously measured for all larvae on a plate during a 50 minute 208 209 simulated light-dark-light cycle, consisting of 20 minutes of light, 20 minutes of darkness, and final 10 minutes of light. The average swimming speed was calculated by dividing the cumulated 210 211 distance travelled with the total time spent active. The light level was set to 100 % on the 212 ViewPoint software. The larval activity was tracked during the dark period. After the behavioural 213 test, the larvae were inspected with a stereo microscope to identify dead or deformed larvae.

214 **2.7 Statistical analysis**

After evaluating and arranging the data in Excel, the differences in general toxicity, heart rate and 215 216 LPO between exposure groups were analysed using a one-way ANOVA and Tukey's multiple comparison tests (GraphPad Prism 7 Software Inc., La Jolla, CA, USA). Differences between dose 217 and time were compared for ROS production using a Two-way ANOVA followed by the post-hoc 218 219 Tukey test (GraphPad Prism 7 Software Inc., La Jolla, CA, USA). For behavioural analyses, data 220 were transferred to R version 2.15.0 (R Development Core Team, http://www.r-project.org). Dead 221 and deformed larvae were excluded from behavioural analyses. Only the cumulative data from the 222 20 minutes dark period of the test were used, as movement was minimal during the lighted periods. Linear mixed effect (LME) models were used within the "nlme" package of R to assess behaviour. 223 The dependent variable was either the cumulative time spent active, the cumulative distance 224 travelled, or average speed (calculated as the cumulated distance travelled/cumulated time spent 225 226 active), with dose as a categorical independent variable, and replicate as a random effect. For all 227 models, examination of the residual plots verified that no systematic patterns occurred in the errors (e.g. q-q plots). To assess individual doses to the controls, we used the contrast results provided 228 within R. Significance in all tests was assigned at $p \le 0.05$. 229

230

231 **3 Results**

3.1 Developmental effects and heart rate

Analyses of mortality, deformities or hatching at 48, 72 and 96 hpf between controls and the exposed groups using the doses below LD_{50} showed no significant differences compared to controls and were generally below 10% (Table 2A and 3A). There was no difference in mortality between 48 hpf and later time points. Additionally, there was no difference in body length at 72hpf between exposed and control larvae (Table 2A).

UVA exposure significantly decreased the heart rate at 50 hpf in all exposed groups compared to controls, while at 60 hpf, the decrease remained significant only in the 18.7 and 37.4 J/cm² UVA doses (p = 0.04 and p = 0.003, respectively) (Fig 1A). The results of UVB exposure showed a significant decrease in mean heart rate at the highest UVB dose compared to the controls (p < 0.01) at 50 hpf , while no significant differences were observed in the two lower UVB doses compared to controls (Fig 1B). By 60 hpf, no differences in heart rate were found between the UVB exposed and control groups.



Fig 1. Heart rate measured at 50 and 60 hpf in zebrafish exposed to sub-lethal UV radiation. Data presented as mean \pm SEM. (One way ANOVA, p < 0.006). Significant difference between groups denoted with asterisks: (*) p < 0.05, (**) p < 0.001, (***) p < 0.0001 according to Tukey's test. (A) UVA exposure. (B) UVB exposure.

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251 **3.2 Oxidative stress**

To assess the potential of UV radiation to generate ROS in zebrafish, the time dependent formation 252 of ROS using a fluorescent probe was measured in all exposure groups. The results showed that 253 for UVA both time and dose were significant for the differences seen in exposed larvae (p < 0.0001254 and p = 0.0025, respectively). No clear pattern of increasing ROS formation was observed in the 255 256 larvae after UVA exposure (Fig 2A), while a trend of increasing ROS formation at the two highest UVB-doses was observed (Fig 2B), although significantly increased only in the highest dose at 257 120 hpf. Two-way ANOVA also showed that both time and dose affected ROS formation 258 259 significantly for the UVB exposed groups (p = 0.0006 and p < 0.0001, respectively), and that their interaction was also significant (p < 0.0001) (Table 4A). 260



Fig 2. ROS fold induction in zebrafish larvae from 72 hpf to 120 hpf exposed to UV radiation. Results presented as mean \pm SEM. Significance in comparison to control denoted with (*) (Twoway ANOVA, p < 0.05; Tukey's test, p < 0.05). (A) UVA exposure. (B) UVB exposure.

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The formation of oxyl-radicals (HO, ROO, RO and peroxynitrite) responsible for lipid oxidation was assessed by the fluorescent dye C11-BODIPY^{581/591} in zebrafish larvae exposed to UVA and UVB. Results showed a small decrease in LPO after 72 hpf in the two highest UVA-doses when compared to the control (Fig 3A), while no LPO was detected after 96 hpf. However, no formation of oxyl-radicals was detected in larvae exposed to UVB (Fig 3B).



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Fig 3. Formation of oxyl-radicals in 72 hpf to 96 hpf zebrafish larvae exposed to UV radiation. Results presented as mean \pm SEM. Significance in comparison to control denoted with (*). (One way ANOVA, p < 0.05; Tukey's test, p < 0.05). (A) UVA exposure. (B) UVB exposure.

On the other hand, the end-products of LPO, MDA and 4-HNE were determined at 72 hpf, where an increase (1.9-fold) in the highest dose was detected after the UVA exposure (Fig 4A). In the

UVB exposure doses, the highest UVB dose demonstrated a non-significant increase in lipid peroxidation compared to control, while the lowest UVB dose caused a significantly decreased lipid peroxidation compared to control (Fig 4B). Therefore, in both wavelength regions, a dose dependent increasing trend in LPO was observed.



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Fig 4. Lipid peroxidation in 72 hpf zebrafish larvae after exposure to UV radiation. Results presented as mean \pm SD. Significant difference between groups denoted with different letters, whereby letters shared by groups represent no significant difference. (One way ANOVA, p < 0.05; Tukey's test, p < 0.05). (A) UVA exposure. (B) UVB exposure.

287 3.3 Behaviour

Analysis of the locomotor activity (LMR) assay data showed that exposure to the highest and second highest dose of UVA significantly reduced the time spent active (p = 0.02 and 0.04, respectively) (Fig 5A), while the highest dose also decreased the total distance moved compared to controls (p = 0.03), but had no effect on swimming speed. The lowest dose of UVA had no effect on behaviour. Exposure to the highest dose of UVB led to an increase in the time spent active (Fig 5B), but a slower average swimming speed although these effects were not significant (p = 0.07). Neither of these tendencies were observed at lower UVB doses.



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Fig 5. Time spent in activity during the dark cycles of the locomotor assay measured in 100 hpf larval zebrafish after exposure to UV radiation. Data presented as means \pm SEM. Significance between groups denoted with different letters, whereby letters shared by groups represent no significant difference (linear mixed effect models $p \le 0.05$, n = 139-143 larvae). (A) UVA exposure. (B) UVB exposure.

302

4 Discussion

This study examined the biological effects in zebrafish larvae following a short and low dose exposure to UVA and UVB radiation during a very sensitive life stage. The results demonstrated that the heart rate, oxidative stress parameters and the behaviour in fish aged 72 to 120 hpf may be persistently altered even at very low doses and that these alterations are wavelength and dosedependent.

309 4.1 General toxicity and heart rate

UV radiation in high levels is able to induce acute toxicity in fish embryos and larvae. The LD₅₀ 310 311 determined following exposure to the mentioned UVA and UVB regimes confirmed that the doses used in this study are below the acute toxic levels. Comparable to the present results, Icoglu 312 Aksakal and Ciltas (2018) reported effects on different parameters in zebrafish, whereby a 313 mortality of 20 % was observed at 24 hpf in embryos exposed to 0.1 J/cm² during a 3h period in 314 the blastula stage of development. Dong et al. (2007) used doses and dose rates higher than in the 315 316 present study, but with different spectra, with LD_{50} about 20x higher in UVB and about 10x higher 317 in UVA, whereby the segmentation stage (12-24 hpf) was more sensitive than the mid-blastula stage. Banerjee and Leptin (2014) exposed zebrafish embryos to lamps with wavelength around 318 320 nm, i.e., between UVA and UVB. The dose inducing close to 50 % embryo mortality at 24 319 hpf was 0.024 J/cm², which is lower than here, but in accordance with the data of Dong et al. 320 (2007), who found that mortality after UVB was higher at 24 hpf than at 3 hpf. 321

322 In order to study the behavioural effects and other effects not induced by acute toxicity, the used doses for late blastula to early gastrula (4.5-5.5 hpf) embryo exposure to UVA and UVB radiation 323 corresponded to maximum 68 and 76% of the LD₅₀, respectively. Correspondingly, no differences 324 325 were observed in development, hatching and body length at later stages of development from these exposures. Deformities included spinal aberrations, yolk sac or cardiac oedema, aberrations in 326 327 pigmentation, and loss of equilibrium, but they were not statistically significant (Table 2A). However, the heart rate was significantly decreased compared to control after exposure to both 328 329 UVA and UVB radiation at 50 hpf. This difference persisted until 60 hpf in the two highest doses of UVA compared to control groups, while in UVB no differences were observed at this stage. 330 Together these results indicate that heart rate is a very sensitive endpoint susceptible to change 331

after exposure to sub-lethal UV radiation in fish larvae, but that it also might be a temporary effect in the lower doses. In addition, the lowered heart rate may be connected to changes in metabolism and/or other physiological parameters, such as oxidative stress, as reported by Icoglu Aksakal and Ciltas (2018). Embryos of Atlantic cod (*Gadus Moruha*) exposed to UVB from the sun seemed to be less sensitive by a factor >10 than zebrafish (Béland et al., 1999). It should be emphasized that the exposure conditions and the shape of the spectra varied widely in the studies cited above and direct comparisons of lethal doses are therefore impractical.

339 4.2 Oxidative stress

The use of fast and direct assays using fluorescent dyes has proven effective for the detection of 340 oxidative stress caused by radiation at a whole-organism level, as previously seen in zebrafish 341 larvae exposed to gamma radiation (Hurem et al., 2017). Results obtained using the H₂DCFDA 342 probe showed that no significant formation of ROS was generated in zebrafish larvae exposed to 343 344 UVA, contrary to what was expected. On the other hand, a more clear time and dose dependent 345 increase was seen at the two highest UVB-doses, even though only significant at 120 hpf for the second highest dose. The reason for not observing an increase in ROS might be related to the time 346 point of the ROS assessment. Zebrafish larvae exposed to both UV wavelengths could have 347 348 undergone different chain reaction processes involved in the oxidative stress mechanism during exposure that could have accounted for a continuous formation and recycling of reactive species, 349 350 which after 72 hpf were not present or not detected by H_2DCFDA . Another possibility for this lack of ROS formation in exposed larvae is the combined action of the antioxidant defence system that 351 might have mediated the ROS formed during and after exposure. A study of chronic exposure of 352 Atlantic cod (*Gadus morhua*) larvae to UVA and UVB radiation (0.001 and 0.006 J/cm² weighted 353

dose), reported a significant increase in antioxidant enzyme activity after constant irradiation for
12 days, which is consistent with the previous notion.

LPO is a marker of oxidative damage and can potentially lead to cell death (Ayala et al., 2014; 356 Livingstone, 2001; Halliwell and Gutteridge, 2007). In this study, no significant increase in LPO 357 levels were detected using the C11-BODIPY^{581/591} probe in zebrafish larvae exposed to both UVA 358 and UVB (Fig A1), even at the UVB doses where an increase in ROS levels was seen. This non-359 360 linearity between the formation of ROS and LPO levels detected in zebrafish larvae can be explained by the specificity of the two fluorescent probes towards different reactive species. The 361 fluorescent probe H_2DCFDA has been shown to be reactive to a variety of ROS, particularly H_2O_2 , 362 HO, NO, ROO, O2- and peroxide-derived oxidants, while the C11-BODIPY^{581/591} probe is 363 triggered only in the presence of oxyl-radicals such as HO, ROO, RO and ONOO⁻ (Drummen et 364 al., 2002). The absence of oxyl-radicals formation in exposed zebrafish can also be related to the 365 formation of other end products of LPO, which are not detectable by the this probe. In fact, the 366 results obtained for LPO when measured as MDA and 4-HNE were more consistent to those 367 obtained for ROS formation and showed significant damage at the highest UVA and UVB doses 368 at 72 hpf. Although the ROS and LPO levels determined were not significantly different between 369 the highest UVB dose and controls, the lowest UVB dose (0.013 J/cm²) demonstrated a significant 370 371 decrease in both parameters at 72 and 120 hpf, respectively. Together, these findings indicate that UVA at doses \geq 37.4 J/cm² is potent enough to cause lipid peroxidation and consequently oxidative 372 damage. It could be speculated that adaptive responses to the highest level of UVB exposure could 373 374 lead to decreased LPO. This study has shown that oxidative stress parameters such as time 375 dependent ROS formation and LPO can demonstrate changes persisting a longer time after early life UV exposure. 376

377 **4.3 Behaviour**

UV radiation levels in aquatic environments are strongly influenced by UV absorption in the water 378 379 and sediments, and current levels have the potency to affect aquatic organisms and induce behavioural changes (Bais et al., 2017). Changes in behaviour may represent either compensatory 380 381 and reversible adaptive responses in order to mitigate potential overt effects after perception of stress), such as reported in Atlantic cod after sea temperature changes (Alemanni et al., 2003; 382 383 Freitas et al., 2015). They also may be irreversible effects of a toxicant on a behavioural mechanism or expression after toxicokinetic and toxicodynamic processes have started (Nellore, 384 2015) and are found to be an indicator of overall welfare (Martins et al., 2012). Some claim that 385 386 behavioural changes might be pointing to neurodevelopmental toxicity of studied agents (Levin and Cerutti, 2009; Rihel and Schier, 2012). 387

Zebrafish larval behaviour was previously shown to be affected after exposure to various toxicants 388 389 at the early embryonic stages (Nellore, 2015; Fraser et al., 2017). Here, behavioural changes 390 resulting from a short duration early life exposure to UV were assessed 5 days post fertilization, 391 and results showed that exposure to the two highest UVA doses resulted in a significant decrease 392 in larval activity compared to the controls. As an example, a decrease in total movement can be an indicator of differences in anti-predator behaviour, concurring with earlier reports showing 393 394 impaired escape behaviour in fish larvae after UV exposure (Fukunishi et al., 2012). The same 395 exposure groups demonstrated a decrease in heart rate, which together with the decreased 396 locomotor activity may be indicative of an overall lower metabolic activity as a consequence of UVA exposure. UVB exposure had no significant effect on larval activity. This result contradicts 397 398 results obtained in studies of behaviour after exposure to environmentally relevant doses of UVB,

whereby decreases in total movement in cod (*Gadus morhua*) larvae (Fukunishi et al. 2012) as well as behavioural differences in juvenile rainbow trout (*Oncorhynchus mykiss*) (Alemanni et al., 2003), were observed. Additionally, in pike eggs (*Esox Lucius*), mortality occurred after exposure to UVB doses similar to one daily erythema weighted ambient dose in Finland in May, in addition to swimming disorders (about 0.27 J/cm²) (Häkkinen et al., 2004), indicating that influence of UVB irradiation effects on the behaviour could have been a factor contributing to increased mortality.

Even though studies demonstrating an interaction of ROS production and behaviour in zebrafish larvae are lacking in the literature, at later developmental stages in zebrafish, an interaction of ROS production and movement was observed after chronic 3-hour daily exposure to UVB radiation (Seebacher et al., 2016). In this study, the ROS formation was significantly decreased in the 37.4 J/cm² UVA dose at 72 hpf. The LPO in this group on the other hand, was increased at the same time point. In addition to the increased LPO in the highest UVA dose, the displayed decreased locomotor activity in these larvae might indicate that oxidative damage is affecting the behaviour.

413

414 **5** Conclusion

Taken in account that climate change may increase exposure of aquatic organisms to increased UV radiation levels, it is important to assess how subtle changes in the UV regime might affect the physiological parameters and the behavior as a key life fitness trait in aquatic populations. From the present findings, it can be concluded that an early life stage exposure to UVA and UVB radiation to sub-lethal and non-detrimental doses to zebrafish development can lower the metabolic activity in later stage embryos and fish larvae. However, depending on the exposure duration and wavelength, this effect persists only temporarily in the shortly exposed UVB groups. On the other hand, in the longer exposed UVA group (68% of the LD₅₀); lipid peroxidation persists for a longer time after the exposure, including the change in resting heart rate, while the total activity of fish larvae is reduced. The findings in this study show that even a very small change in the UV regime during a sensitive developmental stage can induce behavioral changes. Considering that these changes persist long after exposure to low doses of UV radiation during early life, they might have further implications for the fish population dynamics and warrant further studies.

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