

1 **Toxicity Effects of the Environmental Hormone 4-tert-octylphenol in Zebrafish**
2 **(*Danio rerio*)**

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19 **Abstract** 4-tert-octylphenol (4-t-OP), an environmental exogenous estrogen is produced
20 by microbial degradation of alkylphenol polyethoxylates (APEOs). Although it is well
21 known that 4-t-OP can cause the feminization of male, sterility and deficiency of gonad
22 development of aquatic animals by disrupting the endocrine reproductive signaling, less
23 is known about the effects of 4-t-OP on embryonic development. Moreover, the presence
24 of 4-t-OP were detected in umbilical cord blood samples of newborns suggesting infants
25 during development may expose to the risk of 4-t-OP contaminant, hence to investigate
26 the effect of 4-t-OP on physiological function during embryonic development is necessary.

27 In the present study, zebrafish embryos exposed to 4-t-OP were used to evaluate the
28 toxicity of 4-t-OP. The 50% lethal dose (LD50) for wild type zebrafish embryos exposure
29 to 4-t-OP for 3 days is approximately 1.0 M, and a high ratio of cardiovascular defects
30 were showed in survival embryos. To observe the cardiovascular defects more efficiently,
31 Tg(fil-1:EGFP) zebrafish embryos was used in 4-t-OP exposure treatment. Following
32 exposure Tg(fil-1:EGFP) zebrafish embryos to 4-t-OP at 1.0 M for 4 days, a highly
33 proportion of defects revealed in cardiovascular system, including pericardical edema,
34 irregular shape or incomplete looping of ventricle and atrium, the absence of
35 intersegmental vessel in the tail of notochord, unformed parachordal vessel and kinks in
36 the caudal vein. The phenotype of cardiovascular defects was accompanied by reduced
37 heart rate and impaired blood circulation. The mRNA expression levels of transcription
38 factors, which are critical for zebrafish heart chamber formation and blood vessel
39 development, were analyzed by RT-PCR. The results showed that the presence of 4-t-OP
40 significantly induce expression level of *ER α* and *ER β 2*, and caused cardiovascular defects
41 by suppressing transcription factor *Nkx2.7*, *Hand2*, *Tbx2a*, *Tbx2b*, *Tbx5a*, *FGF1a*, *GATA-*
42 *4*, *-5* and *-6* in zebrafish. The present study suggests that 4-t-OP affects the cardiovascular
43 development in zebrafish and elucidated that early life exposure to 4-t-OP potentially may
44 take a risk of impaired cardiovascular function.

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46 **Keywords:** 4-tert-octylphenol; Toxicity; Zebrafish; Cardiovascular development

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48 **1 Introduction**

49 Accompanying with the progress of human activity, loads of industrial or agricultural
50 chemicals introduced into the aquatic environment and have been found to elicit adverse
51 health effects in human and wildlife. One kind of these chemicals is called endocrine

52 disrupting chemicals (EDCs) due to its interfered effects on a physiological function by
53 mimicking or antagonizing the action of the natural hormone. The presence of
54 environmental EDCs in animals body may alter reproduction, secretion, transport, binding
55 and action of natural hormones that are responsible for maintenance of homeostasis (Frye
56 et al., 2012; Kirkley, and Sargis, 2014; Meeker, 2012; Orlando, and Ellestad, 2014).
57 Alkylphenol polyethoxylates (APEOs) are considered as one sort of EDCs, which belong
58 to the group of nonionic surfactants and are widely used in the manufacturing of detergent,
59 plastics, cosmetics, paint, and agrochemicals (Kovarova et al., 2013; Ying et al., 2002).
60 Alkylphenolic contaminants, 4-tert-octylphenol (4-t-OP), is one of microbial degradation
61 products of APEOs and predominant existed in various mediums of water environment,
62 such as sewage sludge, sediments and waste water treatment plants. Several investigations
63 have reported that 4-t-OP contamination occurred in rivers and estuaries of Asian,
64 European, Australia, Africa and South American rivers (Jiang et al., 2012; Watanabe et
65 al., 2007; Brix et al., 2010; Ying et al., 2009; Salgueiro-Gonzalez et al., 2015; Oketola,
66 and Fagbemigun, 2013; Mayer et al., 2007). However, although several reports showed
67 that limited level of 4-t-OP were detected in the worldwide river, the presence of 4-t-OP
68 in aquatic environment was potentially suggesting the release of 4-t-OP from industrial
69 activities. The releasing of 4-t-OP into aquatic environment from manufacturing
70 industries increase the probability of living organism exposure to 4-t-OP and led to the
71 bioaccumulation of 4-t-OP in living organism through direct or indirect uptake process.
72 In recent year global concern regarding 4-t-OP contamination in the environment
73 potentially resulted in toxicity and damage to health due to its xenoestrogen role to disrupt
74 endocrine function through competitive binding to the nature estrogen receptors,
75 consequently, investigators using diverse animal model to evaluate the effects of 4-t-OP
76 on live organism.

77 Studies have shown that the harmful effects of 4-t-OP on reproductive function and
78 endocrine action of diverse fish species, for examples, the adult female of zebrafish
79 exposed to 4-t-OP higher than 25 g/L for 3 weeks resulted in declined ovary somatic
80 index (OSI) in zebrafish (Van den Belt et al., 2001); a regress in testicular growth and
81 vitellogenin (VTG) level induction was observed respectively in male and juvenile
82 rainbow trout after a 3-week exposure to 30 g/L 4-t-OP (Jobling et al., 1996; Van den
83 Belt et al., 2003); exposure to 4-t-OP induced VTG synthesis and disrupts testis
84 morphology in South American freshwater fish (*Cichlasoma dimerus*) (Rey Vazquez et
85 al. 2009); diet supplement of 4-t-OP in *Sparus aurata* induced alteration of liver
86 morphology and degeneration and mediated induction of heat shock protein 70 (Hsp 70)
87 and cathepsin genes, which are bioindicators of endocrine disruption (Traversi et al.,
88 2014). In addition to fish model, 4-t-OP also has been proved to alter cyp19a1 expression
89 profiles involving in gonadal differentiation of male American bullfrog (Wolff et al.,
90 2015), and susceptible to vascular function and led to the reduction of vascular contractile
91 in rats (Hsieh et al., 2009). Due to the toxicity of 4-t-OP and harmful effects over diverse
92 species, many countries including European Union members have legislated to restrict the
93 use of APEOs in domestic application. However, in spite of that, human still have many
94 other pathways exposure to 4-t-OP. Recently, clinical reports showed that 4-t-OP was
95 detected in urine samples from a 57.4% population of the 2517 subjects, and the
96 concentration range of 4-t-OP in subjects is between 0.2 ng/mL and 20.6 ng/mL (Calafat
97 et al., 2008). This result potentially indicated that human already have exhibited high risk
98 exposure to 4-t-OP from living environment including drinking water or food. This view
99 point also can be supported by reports which showed that 4-t-OP was detected in human
100 milk samples which is the main nourishment for newborn, and correlated finding with
101 dietary factors (Ademollo et al., 2008; Chen et al., 2010). The presence of 4-t-OP

102 contaminant in human milk may increase health risks in newborn or infant. Moreover,
103 report also showed that 4-t-OP was detected in 31 samples in concentrations from < 0.05
104 to 1.15 ng/ml from 180 umbilical cord blood samples of newborns, suggesting that
105 expectant mothers exposed to 4-t-OP and leading to contamination of fetus through blood
106 delivery (Tan and Ali Mohd, 2003). It is widely believed that embryos and infants during
107 development are highly sensitive to chemicals that cause serious damage to development
108 and growth, however the effect of 4-t-OP on embryonic development and physiological
109 function of fetus so far is still unclear.

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111 Zebrafish possess discrete organs and tissue that are similar to their human counterparts
112 at the anatomical, physiological, and molecular level. It has become a common
113 experimental model for studying developmental toxicity due to the advantage of rapid
114 development, transparent body for observation, most genes have been characterized from
115 genome databases and a larger number of offspring for providing sufficient experimental
116 material. In the present study, the influences of 4-t-OP on embryonic development and
117 physiological function of a fetus were investigated by using zebrafish embryos exposure
118 to 4-t-OP.

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120 **2 Materials and Method**

121 **2.1 Experimental animals and Compound**

122 Adult AB-strain zebrafish and transgenic zebrafish Tg(fil-1:EGFP) were acquired from
123 the Taiwan Zebrafish Core Facility at Academia Sinica (Taipei, Taiwan). The fish were
124 acclimatized in the laboratory culture condition and observed for clinical health for at least
125 one week prior to experiments. The fish were raised in 10-L tanks and maintained at 28
126 °C in recirculating freshwater with a controlled light cycle (14 h light/10 h dark), and fed

127 daily with commercial pellet. A pair-wise breeding instead of group-breeding was used
128 for breeding of zebrafish in this study to have a better interpretation of the effects.
129 Fertilized embryos generated by pair-wise breeding was used for immersion treatment of
130 4-tert-octylphenol (4-t-OP). All zebrafish were handled in compliance with the local
131 animal welfare regulations. The alkylphenol 4-t-OP with 97% purity (CAS No. 140-66-9)
132 was purchased from Sigma-Aldrich. The 4-t-OP was dissolved in absolute ethanol as 6
133 mM stock solution and then diluted in embryos medium for immersion treatment of
134 zebrafish embryos.

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136 **2.2 Immersion experimental design**

137 Gastrulation is a key event during embryonic morphogenesis and therefore zebrafish
138 embryos with gastrulation stage (5 hour post-fertilization) were used for our exposure
139 studies. Wild type zebrafish embryos at 5 hours post-fertilization (hpf) were collected and
140 put in 12-well microplate for immersion treatment of 4-t-OP. One hundred embryos in
141 each well were immersed with 3 ml of embryos medium (14 mM NaCl, 0.54 mM KCl,
142 0.026 mM Na₂HPO₄, 0.3 mM K₂HPO₄, 0.1 mM CaCl₂ and 0.1 mM MgSO₄ ·7H₂O in
143 deionized water) containing 0.2 mM 1-phenyl-2-thiourea (PTU) and a various
144 concentration of 4-t-OP, and then incubated at 28 °C for 67 h. PTU added in embryos
145 medium was used to prevent pigmentation. Embryos immersed with embryos medium
146 containing 0.2 mM PTU was used as control group. The embryo medium was renewed
147 daily to maintain the water quality and 4-t-OP concentration. Survival rate, hatching rate
148 and malformation were evaluated at 3 days post-fertilization (dpf). The experiment was
149 performed in triplicate for each condition and repeated by three times.

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151 **2.3 Heart rate determination and morphological analysis**

152 Tg(fil-1:EGFP) zebrafish embryos, which enhanced green fluorescent protein (EGFP)
153 was specifically expressed in heart and blood vessel, were used to evaluate the phenotypes
154 of cardiovascular defects resulting from 4-t-OP treatment. Immersion treatment of Tg(fil-
155 1:EGFP) zebrafish embryos with 4-t-OP were carried out as follows: One hundred
156 embryos in each well of 12-well microplate was exposed to 0.5 M or 1.0 M of 4-t-
157 OP from 5 hpf until to the end of embryogenesis (96 hpf). The experiment was performed
158 in triplicate. Twenty Tg(fil-1:EGFP) zebrafish embryos were picked into a petri dish
159 containing 15 ml of embryos medium at 48, 72 and 96 hpf, and heart rate of each zebrafish
160 embryos were calculated under microscopy (Leica Z16 APO). Ten embryos were
161 collected at 48 and 72 hpf for real-time PCR. To observe the morphological defects of
162 heart and blood vessel, live control and 4-t-OP treated embryos were anesthetized with
163 tricaine methanesulfonate (MS222) before mounting in 3% methyl-cellulose (Sigma M-
164 0387) and examined under a Leica stereomicroscope. Digital images or video was
165 acquired using a Leica camera (Leica DFC310 FX).

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167 **2.4 Gene expression detected by real-time PCR**

168 The total RNA was isolated from the Tg (fil-1:EGFP) zebrafish embryos with or without
169 4-t-OP treatment (control group). The expression levels of estrogen receptor (*ER*, *ER 1*,
170 *ER 2*, Nk2 homeobox 5 (*Nkx2.5*), *Nkx2.7*, heart and neural crest derivatives expressed 2
171 (*Hand2*), GATA-binding protein 4 (GATA-4), GATA-5, GATA-6, fibroblast growth
172 factor 1a (*FGF1a*), T-box 2a (*Tbx2a*), *Tbx2b*, *Tbx5a* and elongation factor 1- α (*efl- α*)
173 were determined using quantitative PCR. The *efl- α* was used as an internal control. The
174 specific PCR primers used in this study are listed in Table 1. Real-time PCR was
175 performed using SYBR Green PCR reagents and an Applied Biosystems StepOnePlus
176 Real-Time PCR system. The cycling profile was as follows: 60 °C for 2 min, 95 °C for 10

177 min followed by 40 cycles of denaturing at 95 °C for 15 s, and annealing and primer
178 extension at 60 °C for 1 min. Equal quantities of total RNA were examined in triplicate
179 for each condition. The relative expression levels of each group were normalized to *efl-α*
180 and expressed as the mean ± S.E. Student's t-test was used to statistically analyze and
181 compare two groups. Multiple-group comparisons were analyzed for significant
182 differences between group using one-way ANOVA with a Tukey test (Statistica version
183 5.1; StatSoft. Inc., USA). The differences were defined as significant at $p < 0.05$.

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Table 1 Primer sequences and gene names.

Gene name	Primer sequence (5'→3')	PCR size (bp)	Accession number
Estrogen receptor (<i>ER</i>)	F:CCGGCCCTACACAGAGATCA R:AGCCAAGAGCTCTCCAACAAC	150 bp	NM_152959
Estrogen receptor 1 (<i>ER1</i>)	F: CTGTGCCGTCTGCAGTGATT R: CGGCGGTTCTTGTCGATAGT	150 bp	AF516874
Estrogen receptor 2 (<i>ER 2</i>)	F: TCCGACACCTCAGCAACAAA R: TTTCTGGGCTCTGTTGTCTGCT	150 bp	AF349413
NK2 homeobox 5 (<i>Nkx2.5</i>)	F: CGGGATGGTAAACCGTGTCT R: GCTCGACGGATAGTTGCATGA	150 bp	NM_131421
NK2 homeobox 7 (<i>Nkx2.7</i>)	F: AGCTCACATCCACACAGGTCAA R: GAGCTCCGTGACAGGGTTTG	150 bp	NM_131419
Heart and neural crest derivatives expressed 2 (<i>Hand2</i>)	F: TGTCATGAAGAACCCCTAT R: CCCCGTACTCCTCCGTAGT	150 bp	NM_131626
GATA-binding protein 4 (<i>GATA-4</i>)	F: CCAGTCTGCAACGCATGTG R: GATCGCCGACTGACCTTCAG	150 bp	NM_131236
GATA-binding protein 5 (<i>GATA-5</i>)	F: GGGACGCCAGGGAACCTTA R: CACGCGTTGCACAGGTAGTG	150 bp	NM_131235
GATA-binding protein 6 (<i>GATA-6</i>)	F:AGTCGCGACCCAGTACCTTCAA R: CCTTCGGGATTGCAGTGAGT	150 bp	NM_131557
Fibroblast growth factor 1a (<i>FGF1a</i>)	F: ATGGCAAGCTGTACGCTTCA R: GGCCCGTTTCATTTC	150 bp	NM_200760
T-box 2a (<i>Tbx2a</i>)	F:ACGTTTTCCCTGAGACCGATT R:ATGGAAGGGTCAGCTGTTTCC	150 bp	AF179405
T-box 2b (<i>Tbx2b</i>)	F: ACGTTTTCCCTGAGACCGATT R:ATGGAAGGGTCAGCTGTTTCC	150 bp	NM_131051
T-box 5a (<i>Tbx5a</i>)	F: CGGATGTTTCCCAGCTTCAA R: CATCGCAGGCTCAGCTTTC	150 bp	NM_130915
Elongation factor 1 (<i>ef-1</i>)	F: TGGTGGTGTCCGTGAGTTTG R: AAACGAGCCTGGCTGTAAGG	150 bp	AY422992

204 **3 Results**205 **3.1 Developmental toxicity of 4-tert-octylphenol**

206 To evaluate the toxic effects of 4-t-OP on zebrafish embryogenesis, embryos were
207 exposed to 0.1 M, 0.5 M, 1 M, 2.5 M and 5 M 4-t-OP and compared with their
208 corresponding control group (embryo medium contain PTU only). The survival rate,
209 hatching rate and malformation rate were used as criteria to evaluate the toxicity of 4-t-
210 OP to zebrafish embryos. As a result showed in Table 2, the survival rate, hatching rate
211 and malformation rate exhibit dose effects to 4-t-OP concentration. Embryos treated with
212 0.1 M 4-t-OP as well as control group developed normally, and the survival rate and
213 hatching rate at 3 dpf were more than 95%. However, the survival rate and hatching rate

214 at 3 dpf were declined accompanied by the increasing of 4-t-OP concentration. A 12% of
 215 survival rate and 23% of hatching rate at 3 dpf were showed in the presence of 5 M 4-t-
 216 OP, and all the embryos were seen to be deformed. Concentration higher than 5 M
 217 resulted in 100% mortality at 2 dpf and 3 dpf. Around 50% of survival rate and 60% of
 218 malformation rate at 3 dpf were shown in the embryos treated with 1 M of 4-t-OP, and a
 219 high proportion of cardiovascular defect was revealed in malformation samples. Thus, 1
 220 M of 4-t-OP concentration was used to characterize the cardiovascular phenotype by
 221 exposing them continuously from 5 hpf to 96 hpf.

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225 Table 2 survival rate, hatching rate and malformation rate of zebrafish embryos exposed to different concentration of 4-t-OP at 3 dpf.

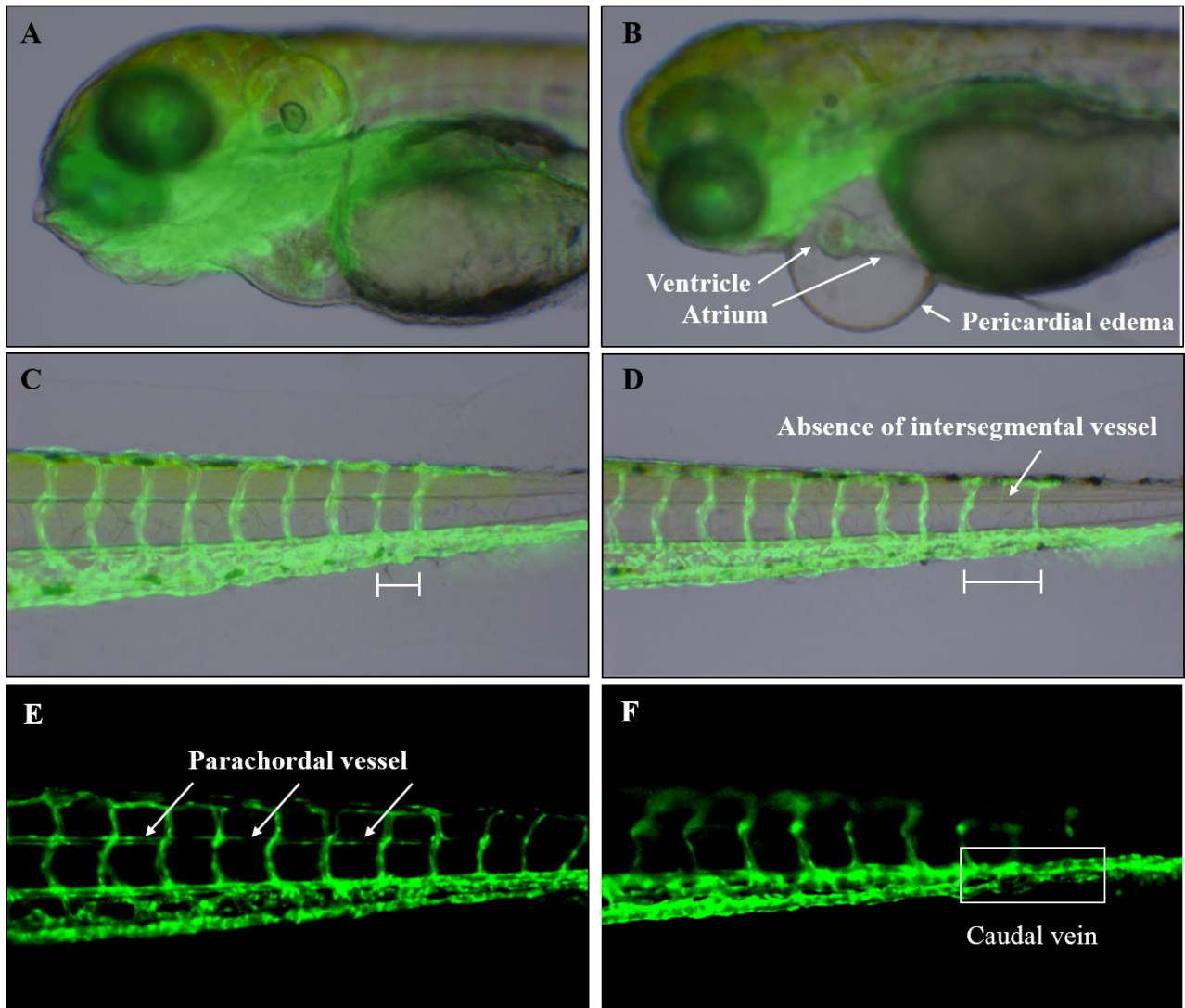
226 Note: ^a Data was evaluated from three times experiment and each experiment was triplicated (n=100). ^b Control indicated the embryo
 227 medium only containing 0.2% PTU.

4t-OP conc. (M)	Survival rate (%) ^a	Hatching rate (%) ^a	Malformation rate (%) ^a
Control (0 M) ^b	96±2.1%	96±2.9%	0 %
0.1 M	95±1.8%	95±1.1%	0%
0.5 M	79±1.5%	83±1.7%	28±1.6%
1.0 M	52±1.8%	71±2.8%	60±1.3%
2.5 M	26±2.8%	45±2.3%	87±4.4%
5.0 M	12±0.4%	23±1.8%	100%

228 3.2 4-tert-octylphenol induced cardiovascular defects

229 To easily observe the cardiovascular defect, Tg (fil-1:EGFP) zebrafish embryos were
 230 used for immersion administration. As result shown in Table 3, exposure of Tg (fil-

231 1:EGFP) zebrafish embryos to 4-t-OP resulted in visible cardiovascular defects from 24
232 hpf. The cardiovascular development was normal in the control group (Fig. 1A, C and D),
233 however the severity of the cardiovascular defects were not consistent among 4-t-OP
234 treated embryos. The majority of embryos exhibited pericardial edema (56% at 2 dpf) and
235 irregular shape or incomplete looping of ventricle and atrium (28% at 2 dpf) (Fig. 1B).
236 The proportion of these phenotypes was increased following 4-t-OP exposure for 3 days,
237 and then declined at 4 days due to the increased mortality. At certain region in the
238 notochord tail, the absence of intersegmental vessel caused the change of the distance
239 between intersegmental vessels and it was also exhibited in 4-t-OP treated embryos (Fig.
240 1C). Furthermore, unformed parachordal vessel and kinks in the caudal vein resulted in
241 blockage of blood flow were exhibited in the 4-t-OP treated embryos (Fig. 1F). These
242 phenotypes were categorized as abnormal blood vessel development, and the proportion
243 was increased accompanying with the time of 4-t-OP treatment (Table 2).



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Figure 1 Cardiovascular defects induced in Tg (fil-1:EGFP) zebrafish at 2 dpf following exposure to 1 M of 4-t-OP. Lateral view of (A, C and E) control Tg (fil-1:EGFP) zebrafish and (B, D and F) 1 M 4-t-OP treated Tg (fil-1:EGFP) embryos. (A, B, C and D) Photos were acquired by merging bright field and dark field using FITC filter. (E and F) Photos were acquired form dark field using FITC filter.

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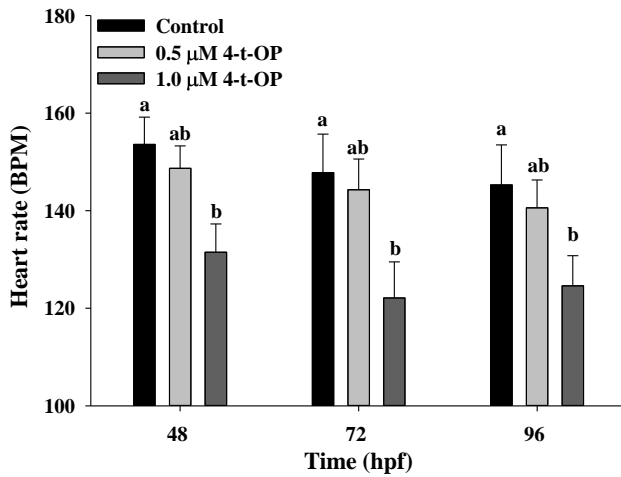
Table 3 Cardiovascular defects induced by 4t-OP

Developmental stages	Treatment	Number of embryos examined	Embryos with cardiovascular defects		
			Pericardial edema	Irregular shape of atrium and ventricle	Abnormal blood vessel development
1 dpf	Control	312	None	None	None
	1 M 4t-OP	287	123 (43%)	49 (17%)	26 (9%)
2 dpf	Control	304	None	None	None
	1 M 4t-OP	273	153 (56%)	76 (28%)	44 (16%)
3 dpf	Control	301	None	None	None
	1 M 4t-OP	178	110 (62%)	83 (47%)	66 (37%)
4 dpf	Control	297	None	None	None
	1 M 4t-OP	143	61 (43%)	31 (21%)	67 (47%)

259 Zebrafish embryos were exposed to control (embryo medium contain PTU only) or 1 M
260 4t-OP and the cardiovascular defects was examined on days 1, 2, 3 and 4. Data shown are
261 the pooled results of triplicated experiments.

262 3.3 4-t-octylphenol damage cardiovascular function

263 The phenotypes of cardiovascular defect induced by 4-t-OP urge us to investigate the
264 effect of 4-t-OP on cardiovascular function. Heart rate variability is a representative index
265 for evaluating the function of cardiovascular function. The development of the cardiac
266 circulation in zebrafish is completed by 48 hpf, so the heart rate was examined at 48,
267 72 and 96 hpf to evaluate the effects of 4-t-OP on cardiac contraction. Although the
268 heart rate of zebrafish at 48, 72 and 96 hpf were decreased than control group following
269 exposure to 4-t-OP at 0.5 M, there was no significant statistical difference between control
270 and 4-t-OP treated group; however zebrafish embryos exposed to 1 M of 4-t-OP
271 significantly reduced heart rate at 48, 72 and 96 hpf compared to that in control group.
272 The heart rate in 4-t-OP treated zebrafish at 48, 72 and 96 hpf were decreased around
273 14.4%, 17.1% and 14.5% respectively compared to control group (Fig. 2). Furthermore,
274 zebrafish exposed to 1 M of 4-t-OP resulted in a slower blood flow rate or a blockage
275 of blood flow was also observed. These results elucidated that exposure to 4-t-OP had
276 potentially damage to cardiovascular function.



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Figure 2 The effect of 4-t-OP on Tg (fil-1:EGFP) zebrafish heart rate following the 96 hours exposure. Data are presented as mean \pm SD of three independent experiments. The asterisk (*) denotes statistically significant differences ($p < 0.05$) between control and 4-t-OP treated groups as measured by Student's t test.

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3.4 Effects of 4-t-OP on the expression of cardiovascular system-related genes in zebrafish

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The significant harmful effect on cardiovascular development and function in zebrafish motivated us to investigate the effect of 4-t-OP on the change of molecular level in cardiovascular system. First, we analyzed the effect of 4-t-OP on the expression of estrogen receptor. Expression levels of ER α and ER 2 genes are significantly increased in zebrafish following exposure to 4-t-OP at 48 and 72 hpf. The result showed that 1 M of 4-t-OP treated zebrafish produced 3.15- and 8.9-fold significantly higher mRNA expression level of ER and ER 2 at 72 hpf compared to control, respectively; however although 1 M of 4-t-OP also induced 1.5- and 2.0-fold significantly increasing in mRNA expression level of ER 1 at 48 and 72 hpf, the increasing level was not strong compared to that in ER and ER 2 (Fig. 3). Moreover the expression level of genes, which was associated with cardiovascular development and function including *Nkx2.5*, *Nkx2.7*, *Hand2*, *Tbx2a*, *Tbx2b*, *Tbx5a*, *FGF1a* and *GATAs* families, in 4-t-OP exposed zebrafish were analyzed by real-time PCR. Compare to the expression of *Nkx2.5*, the expression of

296 *Nkx2.7* were significantly declined in 4-t-OP exposed zebrafish embryos at 48 and 72 hpf.
 297 The mRNA expression level of *Hand2*, *Tbx5a*, *FGF1a* and *GATAs* families including
 298 GATA-4, -5 and -6 were significantly suppressed in 4-t-OP exposed zebrafish at 48 and
 299 72 hpf. The expression level of *Tbx2a* and *Tbx2b* are significantly suppressed only in the
 300 presence of 1 M 4-t-OP at 48 hpf, and significantly suppressed in the presence of 0.5 M
 301 and 1 M 4-t-OP at 72 hpf (Fig. 4). These results suggested that 4-t-OP suppresses the
 302 expression level of cardiovascular development-related genes during zebrafish
 303 embryogenesis.

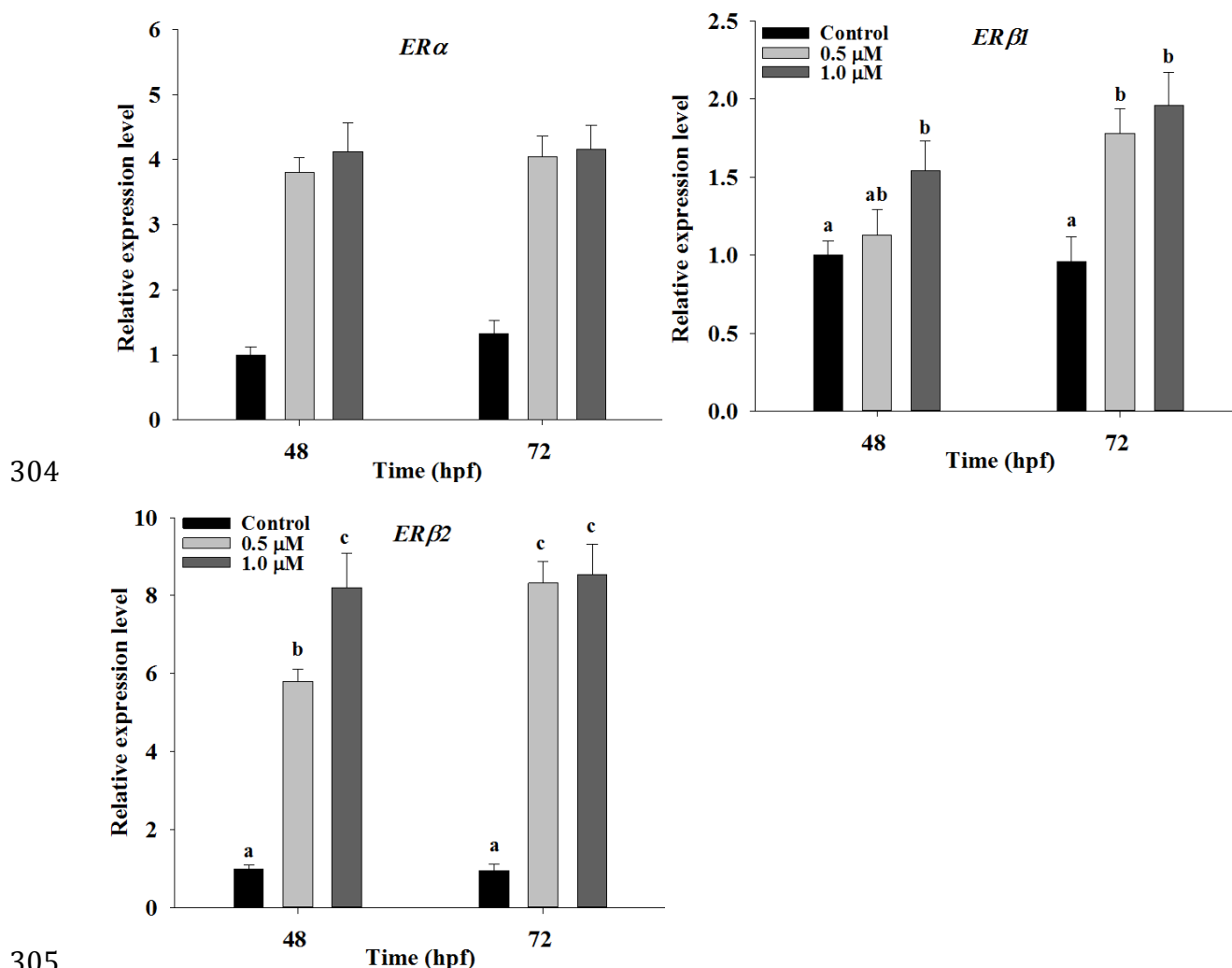
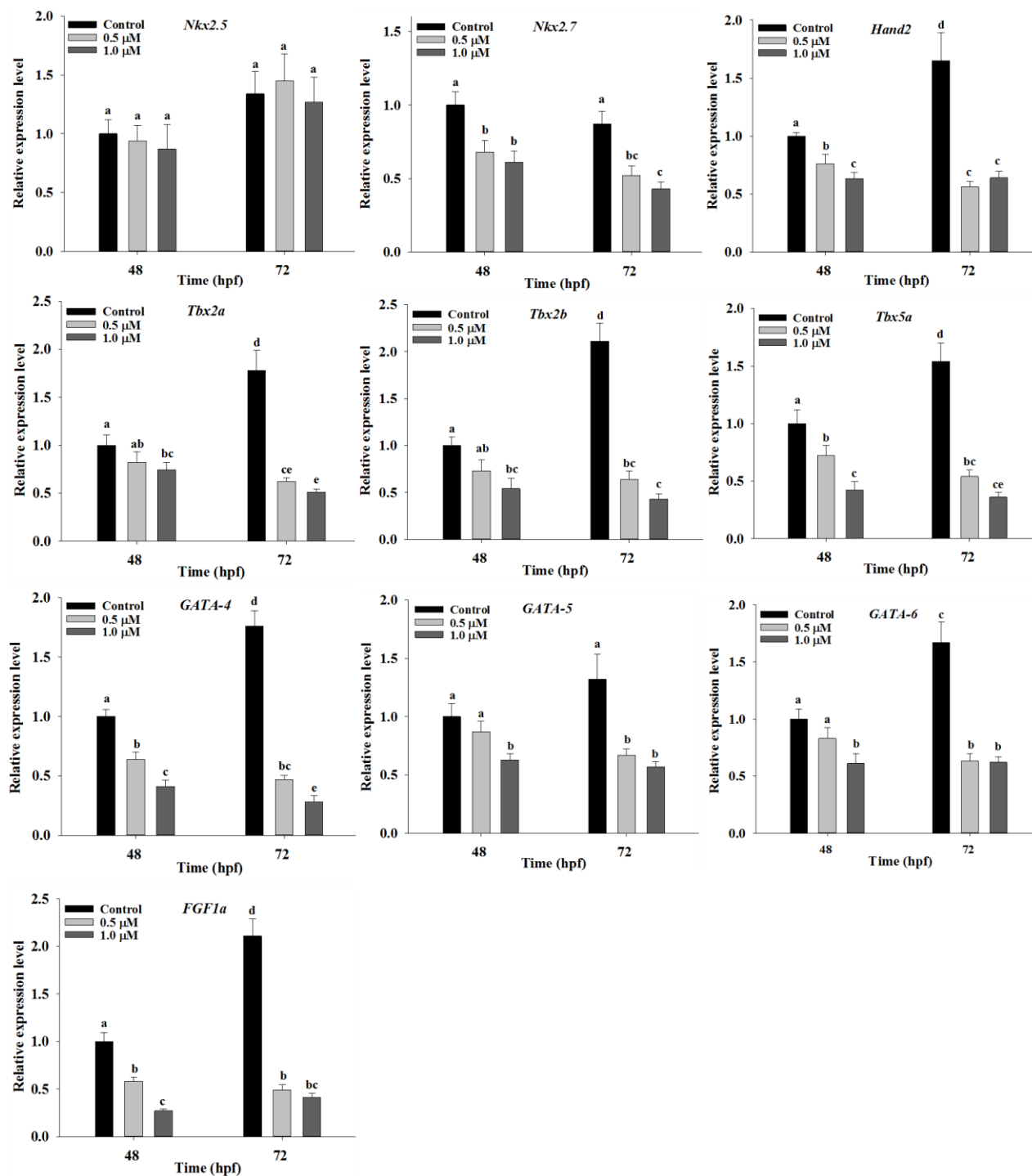


Figure 3 Quantitative PCR analyses of gene expression in 4-t-OP exposed Tg(fil-1:EGFP) zebrafish embryos at 48 and 72 hpf. The

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expression of ER, ER1 and ER2 were quantified. The data of the control group at 48 hpf was set as 1. The results represented triplicated experiments, and the different letters were considered significantly at $p < 0.05$.



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Figure 4 Relative expression level of selected genes in 4-t-OP exposed Tg(fil-1:EGFP) zebrafish embryos were determined by real time PCR at 48 and 72 hpf. The expressions of the cardiovascular-related genes Nkx2.5, Nkx2.7, Hand2, Tbx2a, Tbx2b, Tbx5a, FGF1a, GATA-

312 4, -5 and -6 was quantified. The data of the control group at 48 hpf was set as 1. The results represented triplicated experiments, and the
313 different letters were considered significantly at $p < 0.05$.

314 **4 Discussion**

315 Alkylphenol polyethoxylate (APEs) as nonionic surfactants has been widely used in a
316 variety of industrial and surfactant applications. However, several investigations have
317 reported that the unstable property of APEs in environment cause rapid degradation to
318 hydrophobic and more toxic alkylphenols including 4-nonylphenol (4-NP) and 4-t-OP.
319 Based on hematological and biochemical parameters examined, the study elucidated that
320 OP had a relatively greater effect than NP and affected hematological enzymes leading to
321 serious impairment of the metabolism and physiology in African sharptooth catfish (*C.*
322 *gariepinus*) (Senthil et al., 2011). Other study reported that zebrafish embryos exposed to
323 1 M of 4-t-OP developed normally (Chandrasekar et al., 2011), however our results
324 showed that zebrafish exposure to 1 M of 4-t-OP resulted in cardiovascular defect. These
325 results also suggest that the toxicity effect of 4-t-OP was higher than 4-NP, and affect
326 blood circulation of fish. In the present study the developmental toxicity of 4-t-OP on
327 zebrafish embryos was first demonstrated that 4-t-OP disrupts zebrafish cardiovascular
328 system. 4-t-OP exposure at 1 M significantly decreased heart rate in zebrafish hatchlings.
329 4-t-OP and other endocrine-disrupting compounds has been linked to endocrine disruption
330 mediated via interference with the estrogen and thyroid hormone systems (Ghisari and
331 Bonefeld, 2009). A strong positive correlation between levels of thyroid hormone and
332 heart rate has been demonstrated (Roef et al., 2013). Thus, based on those studies, we
333 assume that 4-t-OP may reduce heart rate through its effects on reducing thyroid hormone.

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335 Transgenic biosensor zebrafish embryos which express the green florescent protein (GFP)
336 under the control of estrogen-inducible promoter had been developed for studying

337 potential health effects of environmental estrogens (Petersen et al., 2013). Exposure of the
338 transgenic biosensor of zebrafish to 4-t-OP induced GFP expressed demonstrating that 4-
339 t-OP possesses ability to act as natural estrogen activity in zebrafish (Brion et al., 2012).
340 Moreover, exposure to alkylphenol induced GFP expressed in heart of transgenic
341 biosensor zebrafish suggesting 4-t-OP act action in cardiovascular system (Lee et al.,
342 2012). It is well-known that estrogen mediates estrogen receptors (ERs) to activate
343 transcription factors (TFs) that modulating estrogen target gene activity. Exposure of
344 zebrafish embryos to 4-t-OP caused carodio vascular defects can be done through 4-t-OP
345 binding of ERs. In zebrafish, the three estrogen receptors, ER, ER 1 and ER 2, had been
346 characterized, and three ERs with a distinct feature in gene structure and tissue distribution
347 pattern (Menuet et al., 2002). In the present study, the expression of estrogen receptors
348 including ER, ER 1 and ER 2 were analyzed. The presence of 4-t-OP at 0.5 M and 1 M
349 significantly induced ER, ER 1 and ER 2 expression in zebrafish, and higher induction
350 level was revealed in ER and ER 2. Reports have showed that ERs expression can be
351 induced by diverse estrogens or estrogen analog, and different type of ER have a different
352 binding affinity for the different ligands. Using HELN assay, which ERE-driven full-
353 length zebrafish ER, ER 1 and ER 2 expression in HeLa cells, 4-t-OP has been
354 demonstrated to have greater affinity towards zebrafish ER and ER 2 relative to ER 1
355 (Pinto et al., 2014). Our study present higher expression level induced by 4-t-OP in ER
356 and ER 2 also potentially suggesting that zebrafish ER and ER 2 have higher affinity for
357 4-t-OP.

358

359 The heart is the first organ to form and function during embryogenesis and its circulatory
360 function is critical for the viability of zebrafish embryos. The presence of 4-t-OP in
361 zebrafish cause cardiovascular defects including incomplete looping of ventricle and

362 atrium, defects in formation of intersegmental vessels and organization of caudal vein,
363 and these indicators signifying that the heart development and circulation function were
364 injured. Several genes encoding transcription factors are required for normal heart and
365 blood vessel development. The *tinmen* gene encodes a NK-class of homeobox
366 transcription factor which plays key roles in the establishment of myogenic lineages. In
367 zebrafish *Nkx2.5* and *Nkx2.7* are expressed in heart field of lateral plate mesoderm and
368 required for cardiac morphogenesis (Stainier, 2001). Report has showed that morpholino
369 (MO) knockdown *Nkx2.5* in zebrafish did not affect heart development. Furthermore
370 *Nkx2.7* has been demonstrated to play a critical function in the lateral development of the
371 heart and normal cardiac looping and chamber formation (Tu et al., 2009). The *hand2*
372 gene encodes bHLH transcription factor that regulate differentiation and the
373 morphogenesis of the myocardial cells and involved in cardiac chamber formation. In the
374 present study, the expression of *NKx2.7* and *hand2* is significantly declined in 4-t-OP
375 exposed zebrafish at 48 and 72 hpf; however expression of *Nkx2.5* without significant
376 difference. This result potentially indicated the 4-t-OP induced incomplete looping of
377 ventricle and atrium, and chamber shape through suppressing *Nkx2.7* and *hand2*
378 expression. GATA family act important transcription factors for the development of
379 diverse tissues. *Tbx2* encodes a T box factor is required for regulating heart chamber
380 development. Report has demonstrated that two genes, *tbxa* and *tbxb*, were retained in
381 zebrafish and both are required for the development of atrioventricular canal (ACV)
382 (Sedletcaia and Evans, 2011). Study also report that homozygous mutation of *tbx5a* gene
383 in zebrafish leads to defects in cardiac looping morphogenesis (Parrie et al., 2013). The
384 three members of GATA family, transcription factor GATA-4, -5, and GATA-6 play a
385 critical role for heart development. GATA-5 is specifically expressed in endocardium and
386 GATA-4 and -6 are present in the myocardium. GATA-5 and GATA-6 involved in

387 regulating endocardial and myocardial cell differentiation (Heicklen et al., 2005). GATA-
388 4 is required for heart tube formation and ventral morphogenesis (Molkentin et al., 1997).
389 In the present study, the expression of *tbx2a*, *tbx2b*, *tbx5a*, *gata-4*, -5 and -6 is significantly
390 declined in zebrafish exposure to 4-t-OP at 48 and 72 hpf suggesting that 4-t-OP
391 suppresses the expression of these critical transcription factors and leads to defects in
392 development and morphogenesis of heart chamber formation. Fibroblast growth factors
393 (FGFs) are considered as important angiogenic factors for vascular development (Javerzat
394 et al., 2002). Other investigators have demonstrated that FGF signaling affects vascular
395 outgrowth and is required for the maintenance of blood vessel integrity in zebrafish (De
396 Smet et al., 2014). In addition to FGF, GATA-4 has been demonstrated to regulate
397 development of the caudal vascular plexus in zebrafish through the chemokine *sdf1a*
398 mediation (Torregroza et al., 2012). The present result showed that downregulation of
399 *EGF* and *GATA-4* expression in the presence of 4-t-OP suggesting 4-t-OP may suppress
400 *EGF* and *GATA-4* expressions in zebrafish resulting in the absence of intersegmental
401 vessel and parachoral vessel and links in the caudal vein.

402

403 In conclusion, the present study is the first report representing that the exposure of
404 zebrafish embryos to 4-t-octylphenol resulting in highly incidence of cardiovascular
405 defects. The presence of 4-t-OP in zebrafish embryos that induced expression level of
406 ER α and ER β 2 suggesting the 4-t-OP mimicking estrogen which act highly binding
407 affinity with both ER. The 4-t-OP exposed zebrafish embryos resulted in suppression of
408 transcription factor *NKX2.7*, *hand2*, *Tbx2*, *Tbx5*, *FGF*, *GATA-4*, -5 and -6 expression may
409 be the cause of cardiovascular defects. The susceptibility of zebrafish model exposed to
410 4-t-OP during early life suggests its role in injuring cardiovascular development and
411 function, which is a health-risk concern of early life exposure in humans.

412

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419

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