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Risk assessment of cardiotoxicity to zebrafish (Danio rerio) by environmental exposure to triclosan and its derivatives

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Abstract
Triclosan (TCS) and its two derivatives (2,4-dichlorophenol and 2,4,6-trichlorophenol) are priority pollutants that coexist in aquatic environments. Joint exposure of TCS, 2,4-dichlorophenol and 2,4,6-trichlorophenol, hereafter referred to as TCS-DT, contributes severe toxicity to aquatic organisms. There is currently a paucity of data regarding TCS-DT molecular toxicity, especially on cardiac diseases. We used zebrafish (Danio rerio) as a model organism, and evaluated the molecular-level cardiotoxicity induced by TCS-DT from embryonic to adult stages. TCS-DT exposure prominently led to phenotypic malformations, such as pericardial cysts, cardiac bleeding, increased SV-BA distance, decreased heart rate and reduced ejection fraction, as well as abnormal swimming behavior. Analyses of the GO and KEGG pathways revealed enrichment pathways related to cardiac development and screened for significantly down-regulated adrenaline signaling in cardiomyocytes. The cardiac marker genes (amhc, cmlc2, vmhc, and nkx2.5) were obtained through protein-protein interaction (PPI) networks, and expressed as down-regulation by WISH. After chronic exposure to TCS-DT from 30 to 90-dpf, both body mass and heart indexes prominently increased, showing myocardial hypertrophy, abnormal heart rate and histopathological injury. Heart tissue damage included disordered and ruptured myocardial fibers, broken and dissolved myofilaments, nuclear pyknosis, mitochondrial injury and inflammatory cell infiltration. Further, abnormal changes in a series of cardiac functions-related biomarkers, including superoxide dismutase, triglyceride, lactate dehydrogenase and creatinine kinase MB, provided evidence for cardiac pathological responses. These results highlight the molecular mechanisms involving TCS-DT induced cardiac toxicity, and provide theoretical data to guide prevention and treatment of pollutant-induced cardiac diseases.

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1. Introduction

Triclosan (TCS) is widely used in medical and personal care products owing to its broad-spectrum antibacterial effect (Dann and Hontela, 2011). However, it may pose serious health risk to human, wildlife and aquatic organisms (Olaniyan et al., 2016). TCS levels in human tissues, urine, blood, and breast milk had been reported to be related to the usage of this antimicrobial agent (Ruszkiewicz et al., 2017); for example, it was even detected in the range of 2.3–3620 μg/L in approximately 80% of urine samples in the USA population (Adgent and Rogan, 2015). Although TCS is a stable and lipophilic compound, it can be transformed into more toxic and bioaccumulative derivatives once in the aquatic environment (Morales et al., 2005), namely 2,4-dichlorophenol (2,4-DCP) and 2,4,6-trichlorophenol (2,4,6-TCP). These intermediates are by-products of water chlorination and combustion processes, contributing to their widespread occurrence in the environment (Jin et al., 2012). Both 2,4-DCP and 2,4,6-TCP are classified as highly carcinogenic chemicals, and thus treated as priority pollutants by many countries (Du et al., 2016; Jiang et al., 2015). TCS and its main
derivatives were frequently detected in soil and municipal sewage at the level of ng/L–μg/L. They cause high bioaccumulation not only in the human body, but also in aquatic organisms such as fish and shellfish (Wang and Tian, 2015; Meador et al., 2018; Liu et al., 2019). In 15 European countries, the concentrations of TCS in effluents of sewage treatment plants reached up to 47800 ng/L (Thomaïdi et al., 2017). In China surface water, 2,4-DCP (1.1–19960 ng/L) and 2,4,6-TCP (1.4–28650 ng/L) were detected in the Yellow River, Huaihe and Haihe River watersheds (Gao et al., 2008). In the surface water of Taihu Lake, the concentrations of 2,4-DCP ranged from 0 to 143 ng/L, while those of 2,4,6-TCP varied between 0 and 840 ng/L (Zhong et al., 2010).

Chronic TCS exposure may cause biological genotoxicity, hepatotoxicity, immunotoxicity, neurotoxicity and cardiotoxicity, as well as impairment of lipid metabolism (Ho et al., 2016; Yueh and Tukey, 2016). TCS leads to craniofacial morphosis in zebrafish (Kim et al., 2018), and acute TCS exposure induces subtle cardiotoxicity in developing fish (Saley et al., 2016). Also, TCS results in changes in fertility function in different animal models. Exposure of pregnant rats and mice to TCS can impair placental development and nutritional transfer (Feng et al., 2016; Cao et al., 2017). In addition to reproductive toxicity, in the C. elegans (Lin et al., 2016), TCS leads to craniofacial morphosis and heart malformation.

2. Materials and methods

2.1. Ethics statement

Our zebrafish-use protocols were approved by the Institutional Animal Care & Use Committee (IACUC) at Wenzhou Medical University. We conducted all experiments in strict compliance with IACUC guidelines. Dissection was performed on ice to minimize zebrafish suffering.

2.2. Chemicals reagents

TCS, 2,4,6-TCP and 2,4-DCP (chemical purity ≥ 98.0%) were acquired from Sigma-Aldrich (St. Louis, USA), and their CAS No. and purities were summarized in Table S1. The stock solutions of three chemicals were prepared in acetone and stored at −20 °C for use. The mixture of TCS, 2,4-DCP and 2,4,6-TCP is hereafter referred to as TCS-DT.

2.3. Zebrafish maintenance and exposure protocols

Wild-type (AB strain) zebrafish maintenance followed previously reported protocols (Li et al., 2016). Our previous study determined LC50 and EC50 values for TCS-DT of 2.28 and 1.16 mg/L, respectively. TCS-DT exposure concentrations were based on the 1/8 to 1/3 increments of the LC50 value (Li et al., 2018; Zhang et al., 2018) resulting in four TCS-DT concentrations (0, 0.28, 0.56 and 0.84 mg/L) for 6–120 h post-fertilization (hpf) zebrafish embryo exposure. Control group included 0.0084% acetone, which was referred to the highest 0.84 mg/L TCS-DT-exposure treatment. This study referred to the optimal concentration ratio of TCS, 2,4,6-TCP and 2,4-DCP (1: 2: 4) for evaluation (Zhang et al., 2018), and their individual concentrations in each concentration of TCS-DT are listed in Table S1. In the control group and 0.84 mg/L TCS-DT-exposure treatment group, 100 larval zebrafish at 120-hpf were employed for extracting the RNA-seq sample for each replicate, and each group was composed of two biological replicates (n = 2). As a consequence, a total of 400 larvae were used for four RNA-seq samples. Additionally, adult zebrafish were exposed to TCS-DT concentrations of 0, 0.14, 0.28 and 0.56 mg/L from 30 to 90 days post-fertilization (dpf). These nominal (fortified) TCS-DT-exposure concentrations simulated real-world environmental levels in aquatic ecosystems. In order to maintain stable water quality and TCS-DT concentration, the exposure solution was renewed daily (6–120 hpf and 30–90 dpf). A schematic representation of overall experimental design is depicted in Fig. S1.

2.4. Assessment of TCS-DT-induced cardiotoxicity

To evaluate the effects of TCS-DT exposure on larval heart development, we assessed cardiac malformations, including heart rate, ejection fraction and SV-BA distance.

(i) Heart rate and ejection fraction measurements

Subsequent to exposure to TCS-DT from 6 hpf, we observed the 48-hpf embryos using a stereoscope coupled to a digital camera (SZX16, Olympus, Japan) to record video images for determination of heartbeat rate for 2 min at 28.5 °C (Li et al., 2019). We used area subtraction to estimate ejection fraction according to Matrone and coworkers’ report (Matrone et al., 2014). Briefly, the ventricular contraction area was subtracted from the diastolic area of the heart, which were expressed as a percentage of the diastolic area.

(ii) SV-BA distance measurement
After TCS-DT exposure, changes in larval cardiac morphology were detected by acquiring lateral images to determine the SV-BA distance between sinus venous (SV) and balloon arterial (BA). Imaging software was utilized to compare the length of lines connecting the two structures on the acquired images (Antkiewicz et al., 2005).

2.5. Behavioral assessment

In this study, 120-hpf zebrafish larvae were transferred to 96-well plates and placed directly in the DanioVision Observation Chamber (Noldus IT, Wageningen, Netherlands) to evaluate the larval responses to external stimuli and spontaneous movements (without any irritation). Zebrafish were preadapted for 5 min prior to light stimulation. In the light-dark stimulation process, we used video analysis to determine average swim distance using EthoVision XT software (Noldus IT).

2.6. Illumina mRNA-seq and bioinformatics analysis

We extracted total RNA using Trizol Reagent (Invitrogen, CA, USA). Bioanalyzer 2100 and RNA 1000 Nano Lab Chip Kit (Agilent, CA, USA) were used to detect the quantity and purity of total RNA, and the RIN number ≥ 9.6, respectively. Poly(A) RNA was purified from total RNA (5 μg) using poly-T oligo-attached magnetic beads. After two rounds of purification, the divergent cations were used to fragment the mRNA into small pieces under elevated temperature. Subsequently, the final cDNA library was created by reverse transcription of the cleaved RNA fragments. The detailed procedures were carried out according to the mRNA-Seq sample preparation kit (Illumina, San Diego, USA), and the average insert size for the paired-end library was 300 bp (±50 bp). Paired-end sequencing performed using an Illumina_HiSeq4000 (LC Sciences, Houston, USA) followed the vendor’s recommended protocols (Mortazavi et al., 2008; Wang et al., 2012). We used StringTie to analyze the expression levels of mRNAs by calculating Fragments Per Kilobase per Million (FPKM) (Trapnell et al., 2010). Differentially expressed mRNAs/genes were screened on the basis of log2 fold change ≥ 0.5 or < -0.5 at p < 0.05 using Feature Count software (Liao et al., 2014). Functional annotation of differentially expressed genes (DEG) was conducted by Gene Ontology (GO) (http://www.geneontology.org/) and “Kyoto Encyclopedia of Genes and Genomes” (KEGG) (http://www.genome.jp/kegg/) functional classification (Kanehisa et al., 2007; Young et al., 2010) and Database for Annotation, Visualization and Integrated Discovery (DAVID, http://david.abcc.ncifcrf.gov/ (Jiang et al., 2008). We imported each protein list encoded by DEGs into the “Search Tool for Retrieving Interacting Genes” (STRING 11.0) database (http://string-db.org/) for the sake of building a PPI network. We used Cytoscape 3.6.1 software (http://www.cytoscape.org/) to map the protein-protein interaction (PPI) network and pathway localization of genes involved in cardiac development.

2.7. qRT-PCR and WISH for heart-related genes

We performed qRT-PCR and WISH to evaluate the genes related to heart development. When exposed to TCS-DT from 6 to 120 hpf, total RNA of 50 larval zebrafish homogenized for each replicate of different groups (0, 0.28, 0.56 and 0.84 mg/L) was isolated using TRIzol reagent, and elf5a and β-actin were used as the endogenous reference (McCurley and Callard, 2008). The primers for qRT-PCR are listed in Tables S2—S3. The cDNA probe sequences of total four genes (amhc, cmil2, vmhc and nkx2.5) in WISH were labeled with digoxigenin (DIG) (Fig. S2). The 6-hpf embryos were exposed to TCS-DT (0, 0.28, 0.56 and 0.84 mg/L), and the control and treatment groups were treated with 0.5% N-phenylthiourea (PTU, Aladdin, Shanghai, China). The 24, 48 and 72-hpf larvae were collected for WISH (Thise and Thise, 2008) and the results recorded using a microscope camera (SZX16, Olympus, Japan). Each group was performed in triplicate, and the primers for WISH are listed in Table S4.

2.8. Measurements of TG, LDH, CK-MB and SOD

We recorded myocardial cellular damage by measuring triglyceride (TG), lactate dehydrogenase (LDH) and creatinine kinase MB (CK-MB) levels in heart tissue using a fully automated biochemical analyzer and a commercial assay kit (Rayto, Shenzhen, China). Superoxide dismutase (SOD) activity is an indicator of reactive oxygen radical and lipid superoxide levels (Lubrano and Balzan, 2015), which was detected in the 90-dpf adult heart using the hydroxylamine method (Nanjing Jiancheng Bioengineering Institute, Nanjing, China).

2.9. Histopathological observation of heart tissue

After TCS-DT exposure from 30 to 90 dpf, zebrafish heart tissue was dissected for histopathological observation. We performed H&E staining of the isolated heart tissue using a HE-Staining kit (Solarbio, Beijing, China) following manufacturer’s instructions. Paraffin sections (4 μm) were Masson stained to differentiate between normal and diseased heart tissues; muscle fibers were red and collagen fibers were blue. Microscopic observation of heart damage was assessed using transmission electron microscopy (TEM7500, Hitachi, Tokyo, Japan).

2.10. Statistical analysis

Experimental data are reported as the mean ± standard deviation (SD). Table S5 summaries the number of biological replicates, technological replicates and zebrafish evaluated at different developmental stages. We assessed the TCS-DT treatment effects using one-way analysis of variance (ANOVA). All statistical analyses were conducted with SPSS 18.0 (SPSS, Chicago, USA) using p < 0.05 (*), p < 0.01 (**) or p < 0.001 (***) significance levels, unless otherwise stated.

3. Results

3.1. Effects of TCS-DT exposure on zebrafish morphology and swimming behavior

Three TCS-DT concentrations were implemented to evaluate the exposure effects on cardiac development and swim behavior. Prominent heart phenotypic defects occurred for 72-hpf larvae with a dose-response relationship. The 0.28 mg/L treatment produced a slight pericardial cyst (PC), and the 0.56 mg/L treatment resulted in severe yolk retention, PC, and SV bleeding. In the 0.84 mg/L treatment, dysplasia became more pronounced, which included small eye deformities, yolk cyst, severe PC, SV bleeding, closure of swim sac (SS), ear capsule and otolith differentiation defects (Fig. S3A).

The standard for assessing whether hearing and vision are developing normally is the stress response of zebrafish larvae to external sound or optical stimuli (Einhorn et al., 2012). Average swim distance after stimulation served as a response sensitivity and vitality metric. There was a slight increase in spontaneous movement distance for embryos in the 0.28 and 0.84 mg/L treatments at 24-hpf (fetal movement) (p > 0.05). In contrast, the 0.56 mg/L TCS-DT treatment produced a significant increase (p < 0.05) in spontaneous movement distance (Fig. S3B). Light-dark
rhythm activities of 120-hpf larvae were evaluated using a 30-min alternating photoperiod stimulation (Chen and Wang, 2012). In the control group, larval swim activity showed an initial burst after a rapid transition from dark-to-light; followed by regular motion rhythms consisting of more exercise during dark conditions and less exercise in light conditions. In contrast, the 0.28 mg/L treatment yielded excessive excitement, although there were also regular reaction rhythms to light-dark stimulation. Finally, the 0.56 and 0.84 mg/L treatments resulted in insensitive responses to light-dark stimulation (Fig. S3C).

3.2. Heart rate, ejection fraction and SV-BA distance response to TCS-DT exposure

As compared to the control group, the 48-hpf larval heart rates were significantly reduced (p < 0.05) in the 0.56 and 0.84 mg/L treatments (Fig. S3D). The SV-BA distance is used to mark the looping of the heart tube into the atrium and ventricle, which determines the normal angle between the atria and ventricle (Antkiewicz et al., 2005; Lin et al., 2007), and reflects the change in cardiac looping (Huang et al., 2011). At 120 hpf, TCS-DT exposure caused the SV-BA distance to significantly increase (p < 0.05) in a dose-dependent relationship (Fig. S3E). Ejection fraction is often adopted as an indicator of contraction rate, that is, the ratio of the volume of blood drawn from the ventricle in each cardiac cycle to the maximum volume of the entire heart after diastole (Cikes and Solomon, 2016). At 48 hpf, the TCS-DT-exposed ejection fraction showed a significant concentration-dependent decrease compared to the control group (Fig. S3F, p < 0.05 or p < 0.01). Therefore, we posit that TCS-DT exposure affects cardiac looping and leads to abnormal ventricles, thereby influencing heart function in zebrafish.

3.3. Functional annotation of DEGs and prediction of heart development

To probe the molecular mechanism on phenotypic malformation, we performed deep RNA-seq analysis of the 120-hpf TCS-DT-exposed larvae. Because the expression level of a gene is different in varying tissues and organs of zebrafish, the integrative (synergistic/subtractive) expression effect of this gene reflected in RNA-seq of the entire juvenile body will cause omission of some heart development-related DEGs. For this reason, a lowered threshold of differential fold was adopted for screening DEGs, i.e., |log2 (fold change)| > 0.5 and p < 0.05. Comparison of TCS-DT treatments to the control group revealed 898 up-regulated and 1134 down-regulated DEGs (Fig. S4A). To clarify the function and location in the metabolic pathway of the DEGs, we annotated DEGs through the GO and KEGG pathways. GO annotation indicated that the DEGs affect cardiac development and differentiation or polarization of maternal cells in the yolk by a complex regulatory network.

3.4. Expression pattern clustering of cardiac development-related genes

Several important genes (17 up-regulation and 22 down-regulation) were retrieved from these GO and KEGG pathways, and they were highly correlated with heart development, heart looping, heart morphogenesis, determination of heart left/right asymmetry, regeneration and adrenergic signaling in cardiomyocytes (Fig. S5 and Fig. 1A). Table S6 lists the detailed information for 39 genes, such as functional description, KEGG pathway, chromosome location, transcript ID, change multiple and p-value (DEGs). Among them, 15 genes had identified KEGG pathways while the remaining 24 genes did not (Fig. 1B). To display the underlying mechanism of heart dysgenesis more succinctly, we selected four representative cardiac marker genes (amhc, cmlc2, vmhc and nkd2.5) that constitute a network mapping of protein interactions among these DEGs (3 up-regulated and 12 down-regulated genes; Fig. 1C). The pathway interaction network showed the candidate gene interactions split into three branches according to the interaction distance. Seven genes on one branch interacted closely with cardiac markers genes. In contrast, the other two branches had little evidence to support interactive relationships with cardiac marker genes. In the heart, CAMP signaling controls many basic cell functions, such as automatism, contractility and relaxation (Lezoualc’h et al., 2016). Pentose and glucuronate interconversions are related to ventricular development (Singh et al., 2016). For example, the up-regulated udgh is mainly distributed in the pathway of pentose and glucuronate metabolism in chromosome 1 for regulation of myocardial oxygen supply, while the up-regulated sox9b is located in the CAMP signaling pathway of chromosome 2 (Table S6). Expressions of mgaa, camk2a and ppp2r3a were all down-regulated based on qRT-PCR verification and RNA sequencing (Fig. 1D–E, p < 0.05). These results indicate that the DEGs affect cardiac development and differentiation or polarization of maternal cells in the yolk by a complex regulatory network.

3.5. Expression of cardiac marker genes by qRT-PCR and WISH

We evaluated the changes in expression levels and sites of four cardiac marker genes induced by TCS-DT in zebrafish larvae. Compared to the control group, the expression of amhc was down-regulated with increasing TCS-DT dose-dependent concentrations. vmhc and cmlc2 were significantly up-regulated at low concentration (0.28 mg/L, p < 0.05), while down-regulated at the higher concentrations (0.56 and 0.84 mg/L, p < 0.05). In contrast, nkd2.5 was significantly down-regulated at 0.28 mg/L (p < 0.01), while up-regulated at 0.56 and 0.84 mg/L (p < 0.05) (Fig. 1F). This irregular pattern in expression change for nkd2.5 may be due to universal expression in the entire juvenile zebrafish body. In addition to the heart, nkd2.5 was expressed in the notochord, pharyngeal arch and spleen (Searcy et al., 1998; Sehonova et al., 2019). Therefore, the level of nkd2.5 mRNA in whole mount larvae did not represent its expression changes exclusively in heart tissue.

To elucidate the distribution and changes of cardiac marker genes, we performed WISH analysis for the four genes in 24, 48 and 72-hpf zebrafish embryos and larvae following exposure to TCS-DT. Expression of the amhc gene mainly occurred in the head and heart tissues (Fig. S6A). At 24-hpf, amhc gene expression did not show any prominent change (Fig. S6B, p > 0.05). At 48 and 72-hpf, the amount of amhc expression decreased slowly with increasing TCS-DT concentrations (Figs. S6C–F, p < 0.05). For the nkd2.5 gene, there was no significant difference in the hybrid signal between 24 and 48-hpf, but differences did arise in the 72-hpf embryonic developmental stage. In comparison with the control group, the
expression of nlx2.5 was significantly decreased in the 0.56 and 0.84 mg/L treatments, but it was not the same case in the 0.28 mg/L treatment (Fig. S6G, S6G', p < 0.01).

The cmlc2 gene showed an obvious down-regulation at 24 and 48-hpf in the 0.56 and 0.84 mg/L treatments relative to the control (Fig. 2A–D, p < 0.05). At 72-hpf, the expression level of cmlc2 was significantly reduced in the 0.84 mg/L treatment group (Fig. 2E–F, p < 0.001). Expression of the vmhc gene was slightly decreased in response to TCS-DT stress at the 24-hpf stage of early cardiac development, primarily in the heart (Fig. 2G–H, p < 0.01). The vmhc expression trend continued through the whole embryonic developmental period and remained stable to 72-hpf (Fig. 2I–L, p < 0.05). Integration of these results provides compelling evidence that TCS-DT exposure affects atrial and ventricular differentiation.

3.6. Effects of chronic TCS-DT exposure on cardiac development in adult zebrafish

3.6.1. Changes in zebrafish growth and development after chronic TCS-DT exposure

The effects of chronic TCS-DT exposure on zebrafish growth and development were investigated in detail (Fig. 3A–D), such as effects on body length and width. Body mass index (BMI) is often used as a measure of obesity (Okorodudu et al., 2010). There was no significant difference in cardiac morphology (Fig. 3E–H) and adult body length (Fig. 3Q). Compared to the control group, the BMI and body width of 90-dpf zebrafish in the TCS-DT treatments significantly increased (p < 0.05; Fig. 3Q and S7A). In terms of sexual differentiation, the chronic TCS-DT exposure to zebrafish resulted in increasing female percentage (>58%), demonstrating their
Fig. 2. Differential expression of two cardiac marker genes (cmlc2 and vmhc) by WISH. Note: (1) A, C and E, WISH of cmlc2 expression in 24, 48 and 72-hpf larvae, respectively; (2) B, D and F, IODs of cmlc2 expression in 24, 48 and 72-hpf zebrafish larvae, respectively; (3) G, I, K and K', WISH of vmhc expression in 24, 48 and 72-hpf larvae, respectively; (4) H, J and L, IODs of vmhc expression in 24, 48 and 72-hpf zebrafish larvae, respectively; (5) Arrows indicate the heart; (6) Scale bars: 100 μm; and (7) **, *** and **** indicate significance levels of p < 0.05, p < 0.01 and p < 0.001, respectively.
prominent estrogen disrupting effects (Fig. 3R). Meanwhile, both cardiac organ index and heart weight showed slightly increasing trends with increasing TCS-DT concentrations when compared to the control group (Figs. 3S and S7B).

Compared to the control group, CK-MB activity and triglyceride (TG) levels in adult heart were significantly higher in the TCS-DT treatments ($p < 0.05$; Figs. S7C–D). SOD levels were significantly increased in the 0.14 and 0.28 mg/L treatments, but decreased in the 0.56 mg/L treatment ($p < 0.001$; Fig. S7E). Notably, LDH levels were increased in the 0.14 mg/L treatment ($p < 0.05$), but decreased significantly in the 0.28 and 0.56 mg/L treatments ($p < 0.001$; Fig. S7F). Changes in these enzymatic activities indicate damage to cardiomyocytes (Zhou et al., 2008), suggesting the presence of myocarditis. In general, these findings demonstrate that chronic TCS-DT exposure affects the normal growth and development of zebrafish, especially with respect to triggering heart diseases.

### 3.6.2 TCS-DT induced myocardial ablation and mitochondrial damage

In this study, three TCS-DT concentrations (0.14, 0.28, and 0.56 mg/L) were selected as sublethal doses to assess chronic toxicity effect. Following continuous TCS-DT exposure from 30 to 90-dpf, zebrafish heart tissues were dissected and H&E-stained (Fig. 3I–P). In the control group, myocardial cells were intact (Fig. 3M), the nucleus was ellipsoidal and neatly arranged, and myofilament fiber was continuous. However, some myofilaments were damaged in the 0.14 mg/L treatment, and severely myofibrillar fiber rupture and inflammatory cell infiltration were observed in the 0.28 and 0.56 mg/L treatments (Fig. 3N–P). Maas’ trichrome staining showed cardiomyocytes with well-preserved cytoplasm and prominent nuclei in the control group. In contrast, TCS-DT exposure led to varying degrees of myofilament rupture with a prominent dose-dependence. For example, the
compacted nuclei in the 0.56 mg/L treatment indicated inflammatory infiltration, and a small amount of collagen fibers, suggesting possible cardiac fibrosis (Fig. 4A).

We examined ultrastructural sections of zebrafish heart tissue by TEM to assess the effects of long-term TCS-DT exposure on subcellular structure injury to cardiomyocytes. The control group had a nucleus that was clear and complete, the matrix electron density was uniform. Complete discs were clearly visible in myocardial fibers, the intercellular filaments were connected tightly (Fig. 4B), and myocardial fibers were arranged in a continuous and regular manner. Further, the Z line, the H band and the light and dark bands were clearly distinguishable (Fig. 4C). Additionally, the electron density of mitochondria was clear, the crest neatly arranged and the double-layer membrane clearly visible (Fig. 4D). In the 0.14 mg/L treatment group, the nuclear membrane was slightly wrinkled, the number of intercalated disks decreased, the gap between the filaments was some evidence of edema (Fig. 4B). Further, the H band and the light and dark bands were difficult to distinguish, and there was myofilament breakage (Fig. 4C). Additionally, there was structural dissolution in mitochondria (Fig. 4D). As the TCS-DT concentration increased to 0.28 and 0.56 mg/L, and even the discs were indistinguishable, the myocardial cells showed signs of edema (Fig. 4B). The Z line was also difficult to distinguish (Fig. 4C). These results indicate that exposure to increasing concentrations of TCS-DT causes serious damage to the myocardial subcellular structure that can lead to myocardial inflammation and heart failure.

4. Discussion

The first organ to form and function in vertebrate embryos is the heart (Yin and Pacifici, 2001), which develops from endocardial progenitors and cardiomyocytes (Bakkers, 2011). The nkx2.5 gene is often regarded as a cardiac progenitor cell marker, ensuring the qualitative or quantitative characteristics of the ventricles during the initial differentiation of cardiac muscle cells (Targoff et al., 2013). In this study, we demonstrated that the expression of nkx2.5 was significantly reduced after exposure to TCS-DT. The
change in the expression level of nkx2.5 led to an imbalance of cardiac progenitor cell differentiation, further initiating abnormal myocardial growth and destruction of ventricular morphology. Zebrafish heart is composed of a single atrium and ventricle separated by atrial valves (Sarmah and Marrs, 2016). The vmhc is a ventricular myosin heavy chain gene that plays an important role in heart construction (Zhang and Xu, 2009; Shih et al., 2015). Further, cmlc2 is used as a fluorescent cardiac marker in the construction of transgenic zebrafish (Chen et al., 2008). Additionally, amhc is related to the contraction of the atrial myofibrils that participate in heart contraction and construction of the cardiac muscle skeleton (Berdougo et al., 2003; Abu-Daya et al., 2009). Both atria and ventricles were damaged by TCS-DT exposure, suggesting the jointly toxic effects of TCS and its derivatives.

Based on the qRT-PCR and RNA-seq results, we constructed a regulatory network to illustrate the adverse effects of TCS-DT on heart development (Fig. 5). In brief, TCS-DT induced cardiotoxicity resulted from disruption of several signaling pathways: (i) cAMP signaling pathway, (ii) adrenergic signaling in cardiomyocytes, and (iii) pentose and glucuronate interconversions. Moreover, when traced back to the regulatory mechanism, the transcription levels of cardiac development-related genes, including ugdh, sox9b, ppp2r3a and camk2a, were dysregulated following exposure to TCS-DT. These genes play critical roles in lipid balance and cardiac morphogenesis. For example, ugdh functions in cardiac valve formation (Walsh and Stainier, 2001), and sox9b in cardiomyocytes for cardiac morphogenesis (Gawdzik et al., 2018). mgaa as a MAX dimerization protein, is crucial for heart organ development (Rikin and Evans, 2010). This study demonstrated that sox9b was closely related to the formation of pericardial walls, and the formation and 

Fig. 5. Predictive regulatory mechanism for TCS-DT exposure on zebrafish cardiotoxicity.
migration of epicardial layers around the heart. After exposure to TCDD, the expression of sox9b was decreased, which prevented the formation of heart (Hofsteen et al., 2013). However, our experimental results showed the transcription level of sox9b was up-regulated by TCS-DT exposure, highlighting the need for further molecular-level research to resolve expression trends of sox9b upon exposure to environmental pollutants. Although a certain number of DEGs related to heart development were identified, their detected rate was relatively low due to only two biological replicates for RNA-seq. Under the same RNA-seq depth, the increase of biological replicate number is in concomitant with increase in the detection rate of DEGs (Khang and Lau, 2015). In further research, we will adopt three or more biological replicates for RNA-seq samples in each control or treatment group. It is worth noting that because the larval heart is very difficult to collect, single-cell sequencing technology can be used to evaluate the toxic effect of TCS-DT exposure on larval heart development (Weinberger et al., 2020).

Recent studies revealed that adrenergic signaling enhanced overall cardiac myocyte cohesion. Adrenaline stimulated the heart muscle and contributed to the heart function adapt to sympathetic overall cardiac myocyte cohesion. Adrenaline stimulated the heart sequencing technology can be used to evaluate the toxic effect of TCS-DT exposure on larval heart development (Weinberger et al., 2020).

Herein, we evaluated the cardiac developmental toxicity of TCS-DT to zebrafish larvae and found that it caused a series of cardiac malformations: pericardial edema, heart bleeding and hypertrophy as well as myocardial fibrosis. Based on these phenotype abnormalities, TCS-DT induced cardioxic molecular mechanism was investigated in detail. Cardiac development marker genes (vhmc, nkx2.5, amhc and cmnc2) and adrenergic signaling in cardiomyocytes showed the down-regulated trends upon exposure to TCS-DT. Besides, chronic TCS-DT exposure resulted in prominently abnormal changes in a series of cardiac functions-related biomarkers (TG, LDH, CK-MB and SOD), consequently providing strong evidence for cardiac pathological responses. Due to the coexistence of TCS and its derivatives in real-world aquatic environment, these data are conducive to reveal the molecular mechanism of cardiac toxicity under the joint exposure to TCS, 2,4-DCP and 2,4,6-TCP, also providing a more comprehensive risk assessment.

Declaration of competing interest

The author reports no conflicts of interest in this work.

Credit authorship contribution statement

Danting Wang: Writing - original draft, Writing - review & editing, Methodology, Formal analysis, Data curation, Conceptualization. Yuhuan Zhang: Conceptualization, Methodology, Investigation, Formal analysis, Data curation. Jieyi Li: Data curation, Methodology. Randy A. Dahlgren: Writing - review & editing. Xuedong Wang: Methodology, Investigation, Writing - review & editing. Haishan Huang: Supervision, Project administration. Huili Wang: Conceptualization, Supervision, Project administration, Funding acquisition.

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Appendix A. Supplementary data

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References


