Influence of Endogenous and Exogenous Estrogenic Endocrine on Intestinal Microbiota in Zebrafish

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Abstract

Gender is one of the factors influencing the intestinal microbial composition in mammals, but whether fish also have gender-specific intestinal microbial patterns remains unknown. In this decade, endocrine disrupting chemicals in surface and ground water of many areas and increasing observation of freshwater male fish displaying female sexual characteristics have been reported. Here we identified the difference in intestinal microbiota between male and female zebrafish, and revealed the influence of endocrine disrupting chemicals on zebrafish intestinal microbiota by using high-throughput sequencing. The results indicated that Fusobacteria, Bacteroidetes and Proteobacteria were dominant in the gut of zebrafish and there were no obvious gender-specific intestinal microbial patterns. Two endocrine disrupting chemicals, Estradiol (E2) and Bisphenol A (BPA), were selected to treat male zebrafish for 5 weeks. E2 and BPA increased vitellogenin expression in the liver of male zebrafish and altered the intestinal microbial composition with the abundance of the phylum CKC4 increased significantly. Our results suggested that because of the developmental character and living environment, gender did not influence the assembly of intestinal microbiota in zebrafish as it does in mammals, but exposure extra to endocrine disrupting chemicals disturbed the intestinal microbial composition, which may be related to changes in host physiological metabolism.

Introduction

For most terrestrial mammals, gut microbiota are involved in absorbing nutrition [1], defending against etiological microbes [2] and even modulating social behaviors [3]. Bacterial dysbiosis has been shown to be closely related to the risk of gastrointestinal disease [4], genitourinary inflammation [5], pre-malignant lesions in the colon [6] and breast cancers [7]. In 1984, Adlercreutz et al. found that antibiotics could reduce the estrogen level and assumed that intestinal microbiota may relate to estrogen metabolism [8]. Since then, more and more evidences have shown that microbiota of the female and male animals respond to the same stimulation in
distinct ways. For example, gender bias has been observed in numerous diseases such as inflammatory bowel disease and Type 1 diabetes (T1D) [9,10]. Removal of the microbiota reduced T1D incidence and increased the testosterone level in female mice [10], suggesting a complex but underlying interaction between microbiota and sex hormone level. However, the relationships among the intestinal microbiota, sex hormone and host health status have not been extensively investigated in animals other than terrestrial mammals [11].

In this decade, endocrine disrupting chemicals (EDCs) have been reported in surface and ground waters in many areas including Europe, Asia, South America and Oceania[12]. Freshwater male fish display female sexual characteristics due to the EDC pollution. The UK Government’s Environment Agency found in 2004 that 86% of male fish sampled at 51 sites around the country were intersex [13]. Considering the close relationship between hormone and gut microbiota in terrestrial mammals and the increasing observation of sexual disruption in fish [14], we wondered whether the “microgenderome”, i.e. sex hormone modulation associated with the microbiome [9], exists in fish and whether EDCs could modify the microbiota.

Estradiol (E2), as a natural EDC, functions via estrogen receptors (ERα and ERβ). It increases vitellogenin (VTG) level, which results in transsexual male fish and genital deformities [15]. Bisphenol A (BPA), a common synthetic EDC, is found in a multitude of products including food, beverage packaging, flame retardants, adhesives, building materials, electronic components and paper coatings [16]. BPA, a nonsteroidal xenoestrogen, exhibits approxi- mately 1/10^4 the activity of estradiol [17]. The effects of E2 and BPA on the microbiota of fish remain unknown.

In this study, zebrafish (Danio rerio) was selected as the model. Differences in intestinal microbiota between male and female fish were characterized. In addition, the influences of E2 and BPA on the intestinal microbial composition were identified.

**Methods and Materials**

**Experimental animals, design and facilities**

Zebrafish were purchased from Yuyi tropical fish farm (Shanghai, China). The average weight was 0.40±0.024g. BPA was purchased from Sangon Biotech (Shanghai, China). E2 was purchased from Fluka (Mo, USA). Acetone was purchased from Lingfeng Chemical Reagent Co. Ltd. (Shanghai, China) and acetone was the vehicle for E2 and BPA. Serial concentrations of E2 (500ng/L and 2000ng/L) and BPA (200μg/L and 2000μg/L) were used for pre-trial[18,19]. Because of the pre-trial data (S1 Fig) and the short exposing time (5 weeks in the present study) compared with the long exposing time in natural water, BPA at 2000μg/L and E2 at 2000ng/L were selected for subsequent experiments.

The zebrafish were divided into four groups according to the treatment: Group F, female zebrafish were reared with 3.33mL/L acetone; Group M, male fish were reared with 3.33mL/L acetone; Group E2, male zebrafish were reared with 2000ng/L E2; Group BPA, male zebrafish were reared with 2000μg/L BPA. Only the male fish were treated with EDCs to avoid the effect of the female own estrogen. Group F and Group M occupied five tanks which contained 30 fish in each tank while Group BPA and Group E2 occupied four tanks which contained 30 fish in each tank. All the fish were reared for 5 weeks. During the whole experiment, one-third of the tank water was exchanged once per day with aerated water supplied BPA, E2, or acetone, as appropriate. The water quality parameters across the whole experiment were pH 7.5–7.9, temperature 26–28°C, dissolved oxygen 4.8–6.4 mg/L, and total ammonia nitrogen <0.02 mg/L. Zebrafish were fed three times per day, at 08:00, 16:00 and 22:00 h. Based on the amount of feed remaining on the following day, daily rations were adjusted to a feeding level slightly over satiation. Uneaten feed was removed daily with a siphon tube. The initial body length and
weight of fish were similar and no significant difference was found at the end of the experiment (S2 Fig). All samples were collected at the end of the feeding trial following a 24 h starvation period. Etomidate was injected 0.3μg/g for anesthesia before the decapitation.

All experiments were conducted under the Guidance of the “Care and Use of Laboratory Animals in China”. This research was approved by the Committee on the Ethics of Animal Experiments of East China Normal University.

VTG gene expression level in male zebrafish liver

The liver of zebrafish in Groups M, E2 and BPA were carefully divided from each fish. Five fish from each tank were collected as one sample in each group. Trizol reagent (Invitrogen, Carlsbad, CA, USA) was used to extract total RNA from liver samples. A Nano Drop 2000 (Thermo Scientific, Waltham, MA, USA) was used to measure the amount of RNA. Visualization of the 28S/18S ribosomal RNA ratio on a 1% agarose gel was used to assess the RNA quality of each sample. Total RNA (1μg) was used to synthesize complementary DNA using the PrimeScript™ RT Reagent Kit (Takara, Dalian, China), according to the manufacturer’s instructions. The forward and reverse primers for VTG gene expression quantification were, VTGF: 5ʹ-TTTTGCTATCATCGCCCGT-3ʹ and VTGR: 5ʹ-TTCACCCAGGACACCAGCTT-3ʹ. The PCR program was run on a CFX Connect Real Time System (Bio-Rad, USA) with Ultra SYBR Mixture (CWBio, Beijing, China). The PCR program began with a 3 min denaturation step at 94°C; followed by 30 cycles of 1 min at 94°C (denaturation), a 1 min annealing and elongation collective step at 60°C. Housekeeping gene was β-actin, and the primers were ACTF: 5ʹ-TCTGGTGATGGGTGACCA-3ʹ and ACTR: 5ʹ-GGTGAAGCTGTAGCCACGCT-3ʹ. The quantification of gene expression was calculated by the comparative ΔΔCT method, VTG expression in each group was normalized to the endogenous reference β-actin level and reported as the fold difference relative to β-actin gene expression. All the assays were performed in triplicate and repeated at least twice. Data were expressed as mean ±standard error of the mean (SEM).

Triglyceride content of male zebrafish muscle

TG content, which was one of the most important parameters reflecting the host lipid metabolism, was detected in this study. Because of the limited available quantity of zebrafish livers, triglyceride (TG) content was determined in muscles of fish from Groups M, E2 and BPA. About 0.034±0.010 g muscle was prepared for TG content test. A TG test kit (BHKT Clinical Reagent Co. Ltd., Beijing, China) was used. In brief, 20mL ethanol was added to each gram of tissue homogenate. Then the mixture was centrifuged at 2,000g for 10min at 4°C. The GPO-AAP method was performed to test the supernatant [20].

Intestinal content collection and bacterial DNA extraction

Whole intestines of five zebrafish from each tank were aseptically dissected, and the intestinal content from each tank was pooled as one sample. Groups M and F contained five samples while Group E2 and BPA contained four samples. Total bacterial community DNA was isolated with an E.Z.N.A.™ Soil DNA Kit (Omega, USA). DNA yield was measured in a NanoDrop spectrophotometer. DNA quality was assessed by PCR amplification of the bacterial 16S rRNA gene.

Illumina high-throughput sequencing of barcoded 16S rRNA genes

Bacterial DNA was used as the template for 16S rRNA gene V4-V5 region amplification [21]. The forward and reverse primers were 515F: 5ʹ-GTGCCAGCMGGCGTAA-3ʹ and 907R:
5ʹ-CCGTCATTCTTTRAGTTT-3ʹ. Unique eight-base barcodes were added to each primer to distinguish PCR products. The PCR reaction mixture (25 μL) included 0.25 U of Platinum® Pfx DNA polymerase (Invitrogen), 2.5 μL of the corresponding 10×Pfx amplification buffer, 0.5 mM of MgSO₄, 0.25 mM of deoxynucleoside triphosphate (dNTP) mixture, 6.25 pmol of each primer, and 20 ng of extracted DNA. The PCR program began with a 3 min denaturation step at 94°C, followed by 20 cycles of 1 min at 94°C (denaturation), a 1 min annealing step (65°C to 57°C with a 1°C reduction every two cycles followed by one cycle at 56°C and one cycle at 55°C), and a 1 min elongation step at 72°C; there was a final 6 min extension at 72°C [22]. Thirty nanograms of each purified PCR product were subjected to Illumina-based high-throughput sequencing (Majorbio Bio-Pharm Technology Co., Ltd., Shanghai, China). The sequences obtained in this paper are available in GeneBank with the accession number PRJNA304186.

Bioinformatics and statistical analyzes

Raw fastq files were demultiplexed and quality-filtered using QIIME (version 1.17) [23]. Reads containing more than two mismatches to the primers or more than one mismatch to the barcode were discarded and reads with length <50 bp were removed. Reads (250 bp) were truncated at any site receiving an average quality score of <20 over a 50 bp sliding window.

Operational Taxonomic Units (OTUs) were clustered with a 97% similarity cutoff using UPARSE (version 7.1; http://drive5.com/uparse/) [24] and chimeric sequences were identified and removed using UCHIME [25]. The phylogenetic affiliation of each 16S rRNA gene sequence was analyzed by RDP Classifier (http://rdp.cme.msu.edu/) against the SILVA database using a confidence threshold of 70%.

Taxonomic richness and diversity estimators were determined for each library in Mothur. Rarefaction curves were created in Mothur to determine whether sequencing depth was sufficient to cover the expected number of OTUs at the level of 97% sequence similarity. ACE and Chao1 were used to reflect community richness [26,27]. Diversity was assessed using Shannon and Simpson indices [28,29]. All these indices were estimated based on OTU abundance matrices. The mean of the estimated parameters was used for comparisons among groups. Principal component analysis (PCA) was used to analyze all OTUs, affording information on microbial community differences among samples. Clustering analysis was based on the average OTU abundance in each group with Matlab R2013b. Network analysis was constructed by igraph package in R language to visualize the most abundant genus and to compare their abundance among groups. Statistical analysis was performed by one-way ANOVA followed by Tukey’s post hoc multiple comparison test. P value < 0.05 was considered statistically significant.

Results

VTG expression level in livers of male fish exposed to E2 or BPA

In order to avoid the influence of host estrogen, VTG expression level was detected on male zebrafish exposed to E2 or BPA at different concentrations for 5 weeks. As stated in the method, two concentration of BPA (2000μg/L) and E2 (2000ng/L) were used. Liver, as one of the most sexually dimorphic organs in terms of gene expression [30], and the main organ for production of VTG, was used for VTG expression level detection [31]. The results indicated that E2 and BPA significantly induced the expression of VTG in male zebrafish (F = 5.891, P = 0.0384) (Fig 1A), suggesting that E2 and BPA treatment caused female characters in male fish.
TG content in muscles of male fish exposed to E2 or BPA

TG content of male zebrafish increased significantly in E2 treatment from 0.67±0.10 to 1.97±0.25 mmol/L (P = 0.0235). However, BPA treatment increased TG content in muscle from 0.67±0.10 to 1.06±0.15 mmol/L (P = 0.0889) (Fig 1B). All these results showed that BPA and E2 accelerated the TG accumulation in male fish.

Microbial complexity of fish gut flora

A total of 236,006 quality reads were collected by Illumina high-throughput sequencing of bar-coded bacterial 16S rRNA genes and used for subsequent data refinement. The rarefaction curves reached a plateau, suggesting good sampling depth (S3 Fig). Group BPA had the largest alpha-diversity indices (Ace, 100; Chao, 88; Shannon 1.63), followed by Group F, and then Group E2. Group M had the lowest alpha-diversity values (Ace, 82; Chao, 79; Shannon 1.26) (Table 1).

Microbial community composition

Fig 2 illustrates bacterial community composition at the phylum level by pie charts. The predominant phyla in Groups M and F were Fusobacteria, Bacteroidetes and Proteobacteria. The abundances of these three phyla in Group M were 47.13%±8.45%, 31.00%±6.09% and 17.46%±0.46%, respectively, while in Group F, the abundance of these three phyla was 32.19%±5.26%, 45.28%±0.01% and 18.56%±3.19%, respectively. The predominant phyla in Group BPA were

Table 1. Summary of intestinal microbiotic species richness estimators for sample groups.

| Group | Reads | Gender | Ace | Chao | Shannon |
|-------|-------|--------|-----|------|---------|
| M     | 12592 | Male   | 82  | 79   | 1.26    |
| F     | 11735 | Female | 88  | 85   | 1.47    |
| BPA   | 11724 | Male   | 100 | 88   | 1.63    |
| E2    | 11201 | Male   | 87  | 79   | 1.05    |

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Fusobacteria, Proteobacteria and Bacteroidetes, representing 49.83%±6.59%, 17.77%±2.41% and 15.50%±3.48% abundance, respectively. The predominated phyla in Group E2 were CKC4, Fusobacteria, Bacteroidetes and Proteobacteria, representing 69.53%±5.44%, 15.77%±3.58%, 5.25%±0.77% and 5.21%±1.16%, respectively. Most interestingly, abundance of CKC4 increased after the treatment with E2 and BPA from 0.05%±0.01% (Group M and Group F) to 5.74%±0.92% (Group BPA) and 69.53%±5.45% (Group E2) (P<0.01). Other shared phyla in all groups included Firmicutes (1.22%±0.35%-9.38%±4.39%), Actinobacteria (0.44%±0.01%-0.84%±0.27%), Tenericutes (0.01%±0.01%-0.28%±0.01%) and Planctomycetes (0.00%±0.01%-0.19%±0.01%).

Principal component analysis was used to identify the intestinal microbial composition in four groups at the OTU level (Fig 3A). PC1 was 55.67% and PC2 was 36.27%. PCA plots showed that Groups F and M clustered together, consistent with our observation of a large shared microbiota between these two groups. Group E2 formed a distinct cluster and separated from the control group (Group M). Although Group BPA did not separate clearly from the control group, samples in Group BPA exhibited clear shifts from Groups M and F towards Group E2. These findings were also confirmed by the clustering analysis (Fig 3B).

Network analysis was used to summarize the proportion of genera in the intestinal microbiota of zebrafish subject to each treatment (Fig 4). Thirty-four genera with abundance >0.01%...
were selected for network analysis. Groups M and F shared a similar gut microbiome composition. The dominant genera in all the groups were *Cetobacterium* and *Flavobacterium* whose abundance ranged from 26.87%±3.58% to 47.13%±8.45% and 4.62%±0.76% to 47.13%±2.64%, respectively. Four genera in Proteobacteria, namely *Acinetobacter*, *Aquabacter*, *Bosea* and *Xanthobacter*, decreased in Groups E2 and BPA compared with Groups M and F, while the proportion of CKC4 increased significantly in Groups BPA and E2 compared with Groups M and Group F.

**Discussion**

Cardiovascular diseases [32], autoimmune diseases [10], human malaria [33] and many other diseases have been proven to be closely related to gender. Other evidences highlight the cross-reactivity among microbial flora, sex hormones and diseases in rodents and humans. For instance, in a tumor necrosis factor receptor 2 knock out (TNFR2−/−2D2) mouse model for central nervous system, demyelinating autoimmune disorders caused mostly female diseases development. Increased abundance of *Bacteroides* *sp.*, *B. uniformis*, and *Parabacteroides* *sp.* in female mice and increased abundance of *Akkermansia muciniphila*, *Oscillospira* *sp.*, *Bacteroides acidifaciens*, *Anaeroplasma* *sp.*, and *Sutterella* *sp.* in male 2D2 mice suggested possible microbial influences on disease causation and protection [34]. It has been demonstrated that the gut community in male and female rats metabolized an oligo-fructose supplemented diet differently with an increase in the content of phylotypes of the phylum Bacteroidetes in female rats, while there were significantly higher levels of the pro-inflammatory cytokines IL-6 and CINC-1 in male rats [35].

It was almost confirmed that the gut microbiota differs in men and women [36–38]. However, analysis of the intestinal microbiota in male and female zebrafish in the present study indicates that they share a similar bacterial composition (Fig 3). Consistent with the previous study [39], Fusobacteria and Proteobacteria are dominant members in both male and female
zebrafish, suggesting that gender does not show significant influence on the intestinal microbiota in zebrafish. The possible reasons are: 1) zebrafish are an undifferentiated gonochoristic species, with all individuals developing an immature ovarian tissue before the differentiation into mature ovaries or testes [40]; 2) zebrafish are exposed to the water environment from the embryo, so the microbial assembly is greatly impacted by water quality. Therefore, we assumed that water quality and the gonad development pathway played a more important role than host sex hormones in the assembly of gut microbiota in zebrafish.

In addition, we found that some studies in fish contained both genders also revealed the similar phenomenon, although no possible explanations had been given. Bacterial composition analysis in different intestine parts of Asian carp and indigenous American fish, with a sex ratio of half male and half female [41], showed no obvious gender-specific intestinal microbiota.

Fig 4. Network analysis visualizing the dominant genera in Groups M, F, BPA and E2. Node sizes correspond to the mean relative abundance of each genus. The proportions of genera shared by different groups are represented by the different colors in the pie charts.

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patterns. Another study conducted in zebrafish during different developmental stages also indicate that there is no significant effect of sex on the microbial community [42].

The powerful influence of water quality was not only restricted to gut microbiota but also applicable to skin microbiota in other aquatic animals including mammals. The data from a study on humpback whales (*Megaptera novaeangliae*) surface microbiota showed that there were non-significant distinctions between male and female whales [43], while it was found that men and women harbored distinct skin bacterial communities, even when controlling for hand hygiene [44].

After exposure of zebrafish to E2 and BPA, a transition to intersex happened in the treatment groups because of an increase in VTG content (Fig 1B). Surprisingly, both BPA and E2 altered the microbial constitution in a distinct manner from gender in PCA plots since female and male fish have similar microbiota to each other (Fig 3A). BPA and E2 treatment lead to a sharp increase in CKC4 abundance (Fig 2). It is a pity that the functional study of CKC4, a phylum in SLIVA database, is very limited, and our results suggested that CKC4 may be sensitive to estrogen and its analogue. The influence of E2 and BPA on the intestinal microbiota was similar, with higher estrogenic effect (E2: BPA = 10:1) resulting in greater distance from the “normal” microbiota in the PCA plot (Fig 3A). Furthermore, the triglyceride content in zebrafish muscle was increased by the BPA and E2 treatment (Fig 1A), which suggested that endocrine exposure and alteration of the intestinal microbiota may be related to changes in host lipid metabolism.

In conclusion, these findings showed that gut microbiota of male and female zebrafish were similar. Application of BPA and E2 resulted in bacterial dysbiosis, which may be related to changes in host lipid metabolism. These observations also suggest that intestinal microbiota may be one indicator for world-wide pollution by EDCs, although the exact mechanism remains unclear. Intestinal microbiota should be considered when we evaluate the influence of environment pollution or stress on the host health.

### Supporting Information

S1 Fig. The mRNA relative expression of in male zebrafish liver with gradient concentration of EDCs. a) The treatment by 500ng/L E2 and 2000ng/L E2. b) The treatment by 200μg/L BPA and 2000μg/L BPA.

(SDOCX)

S2 Fig. The weight and the body length of sampled zebrafish.

(SDOCX)

S3 Fig. Rarefaction analysis of Group M, Group F, Group BPA and Group E2.

(SDOCX)

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