Chronic low-dose dioxin exposure accelerates high fat diet-induced hyperglycemia in female mice

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Abstract:

Human studies consistently show an association between exposure to persistent organic pollutants, including 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD, aka “dioxin”), and increased diabetes risk. We previously showed that acute high-dose (20 µg/kg) TCDD exposure decreased plasma insulin levels in both male and female mice in vivo, however effects on glucose homeostasis were sex-dependent. The current study investigated the impact of chronic low-dose TCDD exposure (20 ng/kg/d, 2x/week) on beta cell function and glucose homeostasis in male and female mice fed either a chow diet or high-fat diet (HFD) for 12 weeks. Low-dose TCDD exposure did not lead to adverse metabolic consequences in chow-fed male or female mice, or in HFD-fed males. However, TCDD accelerated the onset of HFD-induced hyperglycemia and impaired glucose-induced plasma insulin levels in female mice. In addition, islet RNAseq analysis showed that TCDD exposure promoted abnormal changes to endocrine and metabolic pathways in HFD-fed females. Our data supports a ‘double-hit’ hypothesis that TCDD exposure combined with HFD-feeding is more deleterious than either in isolation in female mice, and that chronic TCDD exposure increases diabetes susceptibility in females.

Keywords: dioxin, diabetes, sex differences, beta cells, insulin, hyperglycemia

Abbreviations

AhR      Aryl Hydrocarbon Receptor
CO       Corn oil
ELISA    Enzyme-linked immunofluorescence assay
GTT      Glucose tolerance test
GSIS     Glucose-stimulated insulin secretion
HFD      High-fat diet
ITT      Insulin tolerance test
PFA      Paraformaldehyde
POPs     Persistent organic pollutants
qPCR     Quantitative real-time PCR
TCDD     2,3,7,8-tetrachlorodibenzo-p-dioxin
T2D      Type 2 diabetes
Introduction

Global diabetes incidence is on the rise, yet the underlying cause for this increase remains to be elucidated (1). Genetics and lifestyle factors, such as physical inactivity and poor diet, are known risk factors for type 2 diabetes (T2D) (2,3). For example, high fat diet (HFD) feeding promotes hyperglycemia, hyperinsulinemia, and insulin resistance in both humans and rodents (4). However, these risk factors alone cannot account for the rapid increase in diabetes burden, but rather other environmental factors, such as environmental pollutants, may also be contributing.

Persistent organic pollutants (POPs) are of particular concern to human health since these compounds resist degradation and bioaccumulate in our environment. High serum POP concentrations are positively associated with a modest relative risk of developing T2D (5–10) and insulin resistance (11,12) both in populations exposed to high doses of pollutants (e.g. war veterans, victims of chemical disasters, and occupational workers) (13–15) and the general population exposed to chronic low-doses of pollutants (16–18). Most epidemiological studies investigating the association between POP exposure and diabetes risk only report data from males. Interestingly, the few studies that considered both males and females suggest that females may be at greater risk of developing diabetes following POP exposure. For example, follow up studies on the Seveso population exposed to 2,3,7,8-tetrachlorodibenzo- p-dioxin (TCDD, aka “dioxin”) found a higher relative risk of developing diabetes in females than males (14,19). In addition, high serum PCB levels in the Michigan cohort (20) and Yucheng cohort (10) were associated with significantly increased diabetes incidence in women only. A causal link between POP exposure and T2D pathogenesis remains to be established and potential sex differences in pollutant-induced diabetes risk requires further investigation.

Dioxin/TCDD is a potent aryl hydrocarbon receptor (AhR) agonist and an excellent model chemical for investigating whether dioxin-like compounds increase diabetes risk. Although TCDD is now well regulated worldwide, there are hundreds of dioxin-like chemicals in our environment that remain a global concern for human health (21). We have shown that TCDD exposure induces AhR-regulated genes, including cytochrome P450 1a1 (Cyp1a1), in pancreatic islets in vitro and in vivo, demonstrating that TCDD reaches the endocrine pancreas (22). Direct TCDD exposure also significantly reduced glucose-stimulated insulin secretion in mouse and human islets in vitro (22). Interestingly, a single high-dose injection of TCDD (20 µg/kg) in vivo reduced plasma insulin levels in both male and female mice, but overall metabolic effects differed drastically between sexes (23). TCDD-exposed males had modest fasting hypoglycemia for ~4 weeks post-injection, increased insulin sensitivity, and decreased beta cell area compared to vehicle-exposed controls. Conversely, TCDD did not alter insulin sensitivity or islet composition in females, but did induce transient hyperglycemia during a glucose tolerance test (GTT) at 4 weeks post-injection (23). Whether similar sex-specific effects occur with a more physiologically relevant dose of TCDD remains to be investigated. However, we found that transient low-dose TCDD exposure during pregnancy/lactation in female mice did not impact glucose homeostasis in chow-fed mice, but promoted HFD-induced obesity and hyperglycemia post-exposure (24). These results suggest that chronic low-dose TCDD exposure on its own may not cause diabetes but may increase diabetes susceptibility when challenged with a secondary metabolic stressor, such as HFD feeding. Whether this is specific to female mice or also occurs in males, remains to be determined.
The effects of concurrent TCDD exposure and HFD feeding, termed a ‘double-hit’, are not well understood. One study showed that TCDD exposure promoted hepatic fatty acid deposition in HFD fed female mice (25). In addition, intestinal and hepatic lipid transport genes were significantly upregulated in chow-fed female mice exposed to TCDD compared to vehicle controls, suggesting that TCDD exposure increases hepatic lipid storage, a hallmark of insulin resistance (25). These data support that a ‘double-hit’ of concurrent TCDD exposure and HFD-feeding is more deleterious than either alone in hepatic tissue. However, the potential impact on systemic glucose homeostasis and the endocrine pancreas have not been reported. The purpose of this study was to determine whether chronic-low dose TCDD exposure impairs beta cell function and/or glucose homeostasis in a sex-dependent manner in either chow-fed or HFD-fed mice.

Methods

Animals
Male and female C57BL/6 mice, 6-8 weeks old (Charles River; Raleigh, NC, USA), were maintained on a 12-hour light/dark cycle. All mice received ad libitum access to a standard chow diet for 1 week prior to starting experimental treatments. All experiments were approved by Carleton University and University of Ottawa Animal Care Committees and carried out in accordance with Canadian Council on Animal Care guidelines. All experimental groups were matched for mean body weight and fasting blood glucose levels prior to starting treatments to ensure that the groups were not significantly different.

As outlined in Fig. 1A, male and female mice received intraperitoneal (i.p.) injections of CO (vehicle control, 25 ml/kg) (#48599, Sigma Aldrich, St. Louis, MO, USA) or a low-dose of TCDD (20 ng/kg/d) (Sigma Aldrich, # 48599) 2x/week for 12 weeks. Simultaneously, mice were fed either standard rodent chow (20% fat, 66% carbohydrate, 14% protein; #2918, Harlan Laboratories, Madison, WI, USA) or a HFD (45% fat, 35% carbohydrate, 20% protein; #D12451, Cedarlane, Burlington, ON, Canada), generating the following experimental groups (n=10 per group per sex): COChow, COHFD, TCDDChow, TCDDHFD. Whole pancreas was isolated from a subset of mice at week 12 of the study and stored in 4% paraformaldehyde (PFA; #AAJ19943K2, Thermo Fisher Scientific, Waltham, MA, USA) for 24 hrs, followed by long-term storage in 70% EtOH for histological analysis (n=4-5 per group per sex). Pancreatic islets were isolated from a different subset of female mice at week 12 and stored in RNAlater (#76106, Qiagen, Hilden, Germany) for analysis by quantitative real-time PCR (qPCR) and RNAseq (n=3-5 per group per sex).

Metabolic Assessments
All metabolic assessments were performed on conscious, restrained mice. Blood samples were collected via the saphenous vein using heparinized microhematocrit tubes at the indicated time points and measurements were performed using a handheld glucometer (Johnson & Johnson, New Brunswick, NJ, USA).

Body weight and blood glucose measurements were performed weekly following a 4-hour morning fast. For all metabolic assessments, time 0 indicates the blood collected prior to glucose or insulin administration. For GTTs, mice received an i.p. bolus of glucose (2 g/kg) following a
4-hour fast. Blood samples were collected at 0, 15, and 30 minutes for measuring plasma insulin levels by ELISA (#80-INSMSU-E01, ALPCO, Salem, NH, USA). For insulin tolerance tests (ITT), mice received an i.p. bolus of insulin (0.7 IU/kg, #02024233, Novo Nordisk, Toronto, Canada) following a 4-hour morning fast. Mice from different treatment groups were randomly distributed throughout all experiments to ensure that timing of blood collection did not affect our analysis.

**Immunofluorescence staining and image quantification**

Tissues were processed and embedded in paraffin wax blocks (University of Ottawa Heart Institute, Ottawa, ON, Canada) and then sectioned into 5 µm slices (Thermo Scientific Microm HM 3558). Immunofluorescence staining was performed as previously described (22). In brief, slides were deparaffinized with sequential incubations in xylene and ethanol. Heat-induced epitope retrieval was performed in 10 mM citrate buffer at 95°C for 10-15 minutes, and slides were incubated with Dako Serum Free Protein Block (#X090930-2, Agilent, Santa Clara, CA, USA) for 30 minutes at room temperature. Slides were incubated overnight at 4°C with primary antibodies, and then incubated with secondary antibodies for 1-hour at room temperature. Coverslips were mounted with Vectashield® hardset mounting medium with DAPI (#H-1500, Vector Laboratories, Burlingame, CA, USA).

The following primary antibodies were used: rabbit anti-insulin (1:200, C27C9, #3014, Cell Signaling Technology), and mouse anti-glucagon (1:1000; #G2654, Sigma Aldrich). The following secondary antibodies were used: goat anti-mouse IgG (H+L) secondary antibody, Alexa Fluor 488 (1:1000, Invitrogen, #A11029); and goat anti-rabbit IgG (H+L) secondary antibody, Alexa Fluor 594 (1:1000, Invitrogen, #A11037).

Whole pancreas sections were imaged using a Zeiss Axio Observer 7 microscope, and immunofluorescence was manually quantified using Zen Blue 2.6 software (Zeiss, Oberkochen, Germany). For all measurements, a range of 4-36 islets per mouse were quantified and the average was reported for each biological replicate. The % hormone+ area per islet was calculated as [(hormone+ area / islet area) x 100].

**Islet Isolation**

Islets were isolated by pancreatic duct injections with collagenase (1000 U/ml; #C7657, Sigma-Aldrich) as previously described (22). In brief, inflated pancreas tissues were excised and incubated at 37°C for 12 min, and the collagenase reaction quenched with cold Hank’s balanced salt solution (HBSS) with 1mM CaCl2. The pancreas tissue was washed in HBSS+CaCl2 and resuspended in Ham’s F-10 (#SH30025.01, HyClone, GE Healthcare Bio-sciences, Pittsburgh, PA, USA). Pancreas tissue was filtered through a 70 µm cell strainer and islets were handpicked under a dissecting scope to >95% purity.

**Quantitative Real Time PCR**

RNA was isolated from pancreatic islets using the RNeasy Micro Kit (#74004, Qiagen) as per the manufacturer’s instructions, with the following amendment: 7 volumes of buffer RLT + DTT were added to the samples prior to lysing with 70% EtOH. DNase treatment was performed prior to cDNA synthesis with the iScript™ gDNA Clear cDNA Synthesis Kit (#1725035, Biorad, Mississauga, ON, Canada). qPCR was performed using SsoAdvanced Universal SYBR Green
Supermix (Biorad, #1725271) and run on a CFX96 or CFX394 (Biorad). *Ppia* was used as the reference gene since it displayed stable expression under control and treatment conditions. Data were analyzed using the $2^{\Delta\Delta CT}$ relative quantitation method. Primer sequences are listed in Supp. Table 1.

**Tempo-Seq®**
Gene expression in whole islets was measured using the TempO-Seq Mouse Whole Transcriptome panel (BioSpyder Technologies Inc, Carlsbad, CA). 2 µl of RNA was diluted in 2x TempO-Seq lysis buffer diluted with an equal amount of PBS. Lysates, positive controls (100 ng/µl of Universal Mouse Reference RNA (UMRR) Quantitative PCR Mouse Reference Total RNA Agilent, cat # 750600), and no-cell negative controls (1X TempO-Seq lysis buffer alone) were hybridized to the detector oligo mix following the manufacturer’s instructions (TEMPO-SEQ Mouse Mouse Whole Transcriptome Assay Transcriptome Kit (96 Samples) BioSpyder Technologies, Inc. Carlsbad, CA, USA). Hybridization was followed by nuclease digestion of excess oligos, detector oligo ligation, and amplification of the product with the tagged primers according to manufacturer’s instructions. Each sample’s primers were also ligated to a sample-specific barcode that allows multiplexing for sequencing purposes. Labelled amplicons were pooled and purified using NucleoSpin Gel and PCR Clean-up kits (Takara Bio USA, Inc, Mountain View, CA USA). Libraries were sequenced in-house using a NextSeq 500 High-Throughput Sequencing System (Illumina, San Diego, CA, USA) using 50 cycles from a 75-cycle high throughput flow cell. A median read depth of 2 million reads/sample was achieved.

Reads were extracted from the bcl files, demultiplexed (i.e., assigned to respective sample files) and were processed into fastq files with bcl2fastq v.2.17.1.14. The fastq files were processed with the “pete. star. script_v3.0” supplied by BioSpyder. Briefly, the script uses star v.2.5 to align the reads and the qCount function from QuasR to extract the feature counts specified in a gtf file from the aligned reads. The data were then passed through internal quality control scripts. Boxplots of the log2 CPM (counts per million) were plotted to ensure a reproducible distribution between replicates within a group. Hierarchical clustering plots were generated (hclust function: default linkage function of hclust function in R; complete-linkage) using a distance metric defined as 1-Spearman correlation in order to identify potential outliers. Probes with low counts (i.e. less than a median of 5 counts in at least one group) were flagged as absent genes and eliminated from the dataset. DEG analysis was conducted using the R software (26) on the counts using the default parameters of DESeq2 (27) with respective control and exposure groups. A shrinkage estimator was applied to the fold change estimates using the apeglm method (28) using the lfcShrink() function.

Probes reaching the threshold of an adjusted p-value < 0.05 and an absolute fold change > 1.5 (on a linear scale) were defined as differentially expressed genes (DEGs) and were retained for pathway analysis. Gene set analysis using KEGG pathways was conducted using DAVID (29,30). Pathways with a modified Fisher Extract p-value < 0.05 were considered enriched. Data are presented as bar graphs showing pathway fold enrichment, and secondly, as bar graphs with DEG counts for pathways that were trending (p = 0.05 - 0.1) or significantly enriched (p <0.05). Pathways that were not trending or significantly enriched are indicated on bar graphs as not having the DEG count minimum threshold for significance.

**Statistical analysis**
Statistical analysis was performed using GraphPad Prism Software (GraphPad Software Inc., La Jolla, CA). Specific statistical tests are indicated in the figure legends and p<0.05 was considered statistically significant for all analyses. Statistically significant outliers were detected by a Grubbs’ test with α=0.05. All data was tested for normality using a Shapiro-Wilk test and for equal variance using either a Brown-Forsyth test (for one-way ANOVAs) or an F test (for unpaired t tests). Non-parametric statistics were used in cases where the data failed normality or equal variance tests. Parametric tests were used for all two-way ANOVAs, but normality and equal variance were tested on area under the curve values and by one-way ANOVAs. Data in line graphs are presented as mean ± SEM. Data in box and whisker plots are displayed as median, with whiskers representing maximum and minimum values. Each individual data point represents a different biological replicate (i.e. individual mouse).

Data and resource availability
The datasets generated and/or analyzed during the current study are available from the corresponding author upon reasonable request. Raw RNAseq data have been submitted to GEO (GEO: GSE144765).

Results

Chronic low-dose TCDD exposure does not impact body weight or fasting blood glucose levels in chow-fed or HFD-fed mice
Male and female mice were exposed to CO (vehicle) or 20 ng/kg/d TCDD 2x/week for 12 weeks, and simultaneously fed either a chow diet or 45% HFD (Fig. 1A). We were particularly interested in whether chronic low-dose TCDD exposure alone would promote weight gain (i.e. COChow vs TCDDChow) and/or accelerate diet-induced obesity (i.e. COHFD vs TCDDHFD) as previously seen in female mice exposed to TCDD during pregnancy (24). Interestingly, TCDD exposure had no effect on body weight in chow-fed females in this study (Fig. 1B). In addition, female mice were resistant to HFD-induced weight gain within the 12-week study timeframe, regardless of chemical treatment (Fig. 1B). Likewise, TCDD exposure did not alter body weight in chow-fed or HFD-fed male mice, but HFD feeding promoted significant weight gain compared to chow feeding in male mice from days 73 to 86 (Fig. 1C). TCDD exposure did not influence fasting blood glucose levels in any group, but HFD feeding caused fasting hyperglycemia in both CO- and TCDD-exposed females (Fig. 1D,E) and males (Fig. 1F,G).

TCDD exposure accelerates the onset of HFD-induced hyperglycemia in female mice
TCDD had no effect on glucose tolerance in chow-fed female or male mice at weeks 4 and 8 of the study (COChow vs TCDDChow; Fig. 2i). However, we observed an interaction between TCDD exposure and HFD feeding in female mice (Fig. 2i). At week 4, HFD feeding had no impact on glucose tolerance in CO-exposed females (COChow vs COHFD; Fig. 2Aii,2B), but TCDDHFD females were significantly hyperglycemic at 15- and 30-minutes post-glucose bolus and had a significantly elevated overall glucose excursion compared to TCDDChow females (TCDDChow vs TCDDHFD; Fig. 2Aiii,2B). By week 8, both COHFD and TCDDHFD females were glucose intolerant compared to chow-fed females (Fig. 2C,D). In contrast, HFD-fed male mice were significantly hyperglycemic following a glucose stimulus after 4 and 8 weeks of exposure compared to chow-fed males, irrespective of chemical exposure (Fig. 2E-H). In other words, both TCDD-exposed and CO-exposed males showed a similar response to HFD feeding.
(Fig. 2E-H), whereas HFD-fed female mice had accelerated onset of hyperglycemia in the presence of TCDD (Fig. 2A-D).

**TCDD-exposed female mice lack a compensatory hyperinsulinemic response to HFD feeding but show normal insulin tolerance**

We next assessed whether TCDD altered circulating insulin levels during a GTT. After 8 weeks of HFD feeding, COHFD females showed an expected compensatory increase in plasma insulin levels during the GTT (Fig. 3A) ~2.3-fold more insulin overall than COChow females (Fig. 3A-AUC). In contrast, TCDDHFD females did not show the same compensatory response, and instead plasma insulin levels during the GTT were comparable to chow-fed females (Fig. 3A). TCDD did not impact insulin levels in either chow-fed or HFD-fed male mice (Fig. 3B), but rather diet had an overall effect on insulin levels in males (Fig. 3B, p_{diet}<0.01). These results suggest that TCDD exposure may cause abnormal insulin secretion in HFD-fed females only.

To determine whether the accelerated hyperglycemia in TCDDHFD female mice was driven in part by changes in peripheral insulin sensitivity, we performed ITTs at weeks 5 and 9. HFD-fed male and female mice were not overtly insulin resistant in this study (Fig. 3C-F). At week 5, COHFD females were slightly more insulin sensitive than COChow females at 60 minutes after the insulin bolus (Fig. 3C), and at week 9 TCDDChow females were modestly insulin sensitive at 15 minutes post bolus compared to COChow females (Fig. 3E), but otherwise there were no overall changes in blood glucose levels during ITTs, as measured by area under the curve (Fig. 3C-F).

We also measured expression of genes involved in insulin-dependent lipogenesis and gluconeogenesis pathways in the liver at week 12 as another indicator of insulin sensitivity. In females, HFD feeding significantly downregulated genes involved in gluconeogenesis (Supp. Fig. 1A, p_{diet}<0.05), including *G6pc* (glucose-6-phosphatase catalytic subunit), *Ppargc1a* (peroxisome proliferator-activated receptor gamma coactivator 1 alpha), and *Pck1* (phosphoenolpyruvate carboxykinase 1). There was a significant interaction between TCDD and HFD exposure on expression of *Acacb* (acetyl-CoA carboxylase beta; Supp. Fig. 1A, p_{interaction}<0.05), but this was modest and unlikely to explain the differences observed in plasma insulin levels or glucose homeostasis in female mice. There were no differences in any of the measured genes in the liver of male mice following TCDD and/or HFD exposure (Supp. Fig. 1B).

**Islet size and endocrine composition were unchanged by TCDD exposure**

To investigate why TCDD-exposed female mice did not develop hyperinsulinemia following HFD feeding, we examined islet size and endocrine cell composition. We predicted that HFD-fed mice might show increased islet size compared to chow-fed mice, but this was not the case in either male or female mice (Fig. 4A,D). Surprisingly, there was a significant overall effect of TCDD exposure to increase average islet size in male mice (Fig. 4D, p_{chemical}<0.05). We did not observe any changes in the % of islet area that was immunoreactive for either insulin (Fig. 4B, E,G) or glucagon (Fig. 4C,F,G) in either sex.

**TCDDHFD female mice have enriched endocrine-related pathways in islets relative to COChow females**
Since our histological findings did not explain the phenotype observed in TCDDHFD females, we performed RNAseq analysis on isolated islets from a subset of females at week 12 of the study to assess whole transcriptomic changes. We first compared all experimental groups to COChow females to investigate the effect of TCDD or HFD feeding alone (i.e. TCDDChow and COHFD, respectively), and in combination (TCDDHFD). TCDDChow females had 956 differentially expressed genes (DEGs), whereas COHFD females had 2121 DEGs and TCDDHFD females had 2667 DEGs compared to COChow females (Fig. 5A). Interestingly, only 50 DEGs were uniquely common between TCDDChow and TCDDHFD females, and only 196 were unique to TCDDChow (Fig. 5A). These results suggest that TCDD alone has very minor effects on gene expression in whole islets, which supports our in vivo observations (Figs. 1-4). Furthermore, 893 DEGs were uniquely common between HFD groups; 1180 DEGs were unique to TCDDHFD females and 518 DEGs unique to COHFD females (Fig. 5A), suggesting that TCDD may be driving abnormal changes in gene expression in response to HFD feeding.

Next, we performed KEGG pathway analysis on DEGs in Fig. 5A (TCDDChow, COHFD, and TCDDHFD females were each compared to COChow) using DAVID. Since pathway enrichment is determined based on total number of DEGs and the number of DEGs varied between our experimental groups (Fig. 5A), we presented pathway analysis as both pathway fold enrichment and number of DEGs within each pathway to avoid misinterpretation of our results. We first assessed pathways that were uniquely enriched in TCDDChow females to determine the effects of TCDD alone on the islet transcriptome (Fig. 5Bi). As expected, “Xenobiotic Metabolism by CYP450” was highly enriched in TCDDChow compared to COChow females, and 50% of the pathways enriched in TCDDChow females were involved in drug/chemical exposure, suggesting that TCDD alone mainly alters drug metabolism in islets. It is important to note that although “Xenobiotic Metabolism by CYP450” was not significantly enriched in COHFD or TCDDHFD females, a greater number of genes in the pathway were altered in HFD-fed than chow-fed females (Fig. 5Bi), suggesting that both TCDD exposure and HFD feeding alter this pathway.

We also assessed pathways that were enriched in COHFD and TCDDHFD females compared to COChow controls to identify transcriptomic differences that may explain why TCDDHFD females lacked a compensatory increase in insulin secretion (Fig. 3A). “Maturity Onset Diabetes of the Young” (MODY) was the most enriched pathway in COHFD compared to COChow females, but the number of DEGs was similar in both HFD groups irrespective of chemical treatment (Fig. 5Bii). In addition, “Insulin Secretion” was enriched in both HFD groups, although more genes were altered in TCDDHFD females compared to COHFD females (Fig. 5Biv). Interestingly, several pathways involved in beta cell function were differentially enriched in COHFD and TCDDHFD females. First, “Regulation of Actin Cytoskeleton” was significantly altered in COHFD females but not TCDDHFD females, with 35 DEGs in COHFD females (Fig. 5Bii). Therefore, changes in the actin cytoskeleton in islets may be a normal response to HFD feeding that is absent in the TCDDHFD condition. Second, “Circadian Rhythm” was the most enriched pathway in TCDDHFD compared to COChow females, and more genes within this pathway were changed in TCDDHFD females compared to COHFD females (Fig. 5Bv). Likewise, “FoxO1 Signaling Pathway” was enriched in TCDDHFD females, and more genes were altered in this pathway in TCDDHFD females compared to COHFD females (Fig. 5Bv). Lastly, thyroid hormone signaling pathway was significantly altered in TCDDHFD females only (Fig. 5Bv).
**TCDDHFD females have inappropriate Cyp1a1 and circadian rhythm gene expression**

To better understand the effect of TCDD exposure on the islet transcriptome, we generated both hierarchical heatmaps (Fig. 6) and heatmaps organized by experimental group (Supp. Fig. 2) for pathways of interest that were enriched in TCDDChow and TCDDHFD females compared to COChow controls (Fig. 5). We were particularly interested in the “Xenobiotic Metabolism by CYP450” pathway since we have previously shown that TCDD induces CYP1A1 enzymes in islets (22,23). We expected to see distinct clustering between CO- and TCDD-exposed females, but instead TCDD-exposed and COHFD females had a similar gene expression profile compared to COChow females (Fig. 6A, Supp. Fig. 2A). However, TCDDChow females formed a sub-cluster, indicating slight differences in gene expression compared to HFD-fed females (Fig. 6A). We further assessed the effect of TCDD on xenobiotic metabolism by specifically looking at Cyp1a1 expression as a marker of AhR activation by dioxin exposure (22,23). Interestingly, Cyp1a1 was upregulated ~3.5-fold and ~3.2-fold in TCDDChow and COHFD females, respectively, compared to COChow females (Fig. 6C). In contrast, TCDDHFD islets showed a non-significant trend towards having only ~2-fold higher Cyp1a1 expression compared to COChow females, and Cyp1a1 levels remained significantly lower than in TCDDChow females (Fig. 6C). These results suggest that crosstalk between TCDD exposure and HFD feeding prevents normal Cyp1a1 induction in islets. These findings were validated by qPCR (Fig. 6D).

We also assessed circadian rhythm gene expression patterns since this pathway was the most enriched in TCDDHFD females (Fig. 5Bv). As expected, changes in circadian rhythm gene expression were mainly driven by diet, with chow-fed and HFD-fed females forming two distinct clusters (Fig. 6B). However, within both diet clusters we observed subclusters that were driven by chemical exposure (Fig. 6B, Supp. Fig 2B), suggesting that TCDD may be altering circadian rhythm in islets.

**Chronic low-dose TCDD exposure promotes diet-induced changes in amino acid metabolism**

Our in vivo results (Figs. 2-3) and previous findings from a separate study in pregnant mice (24) suggest that in females, background exposure to TCDD accelerates HFD-induced changes in metabolism. As such, we next investigated the effects of HFD feeding on the islet transcriptome of females with either CO or TCDD background exposure (i.e. COHFD vs COChow compared to TCDDHFD vs TCDDChow). Interestingly, COHFD females had 1674 unique DEGs relative to COChow females, whereas TCDDHFD had 706 unique DEGs relative to TCDDChow females (Fig. 7A), indicating that the effect of HFD varies depending on chemical exposure. In fact, KEGG pathway analysis revealed significant differences between COHFD and TCDDHFD females when compared to their respective chow-fed controls (Fig. 7B). As with our findings in Fig. 5B, MODY was the most enriched pathway in COHFD females; this pathway was also altered in TCDDHFD females but to a lesser extent (Fig. 7Bi). “Regulation of Actin Cytoskeleton” and “PI3K-Akt Signaling Pathway” were only altered by HFD feeding when mice were on a CO background (Fig. 7Bi), suggesting that TCDD exposure may prevent these normal responses to HFD feeding. Most interestingly, the top two most enriched pathways in TCDDHFD compared to TCDDChow females were “Alanine, Aspartate & Glutamate metabolism”, and “Tyrosine Metabolism” (Fig. 7Bii). In fact, 40% of significantly enriched pathways in TCDDHFD females are involved in amino acid metabolism (Fig. 7Bii), suggesting
that background exposure to TCDD may promote abnormal changes to amino acid metabolism in HFD-fed females.

**HFD-induced changes in MODY genes and Alanine, Aspartate, and Glutamate metabolism are promoted by TCDD exposure.**

We generated heatmaps of the most enriched pathway in COHFD and TCDDHFD females with respect to their chow-fed controls (Fig. 7B). Interestingly, the “MODY” heatmap showed that TCDDHFD females formed a unique cluster compared to all other groups (Fig. 8A, Supp. Fig. 2C), suggesting an interactive effect of HFD feeding and TCDD exposure on expression of islet-specific genes. We next looked at specific genes essential for maintaining beta cell function and found a distinct pattern of gene expression in TCDDHFD females compared to COHFD females (Fig. 8A, 8C-H, Supp. Fig. 2C). Expression of MafA and Slc2a2 were significantly downregulated in TCDDHFD females compared to COChow and TCDDChow females, whereas expression was only trending towards being downregulated in COHFD females (Fig. 8C, D, H). Similarly, Hnf4a was downregulated by HFD feeding, but this effect was worsened in TCDD-exposed females (Fig. 8G), whereas Pax6 was only downregulated in TCDDHFD females compared to COChow females (Fig. 8F). Interestingly, Nkx6.1 was upregulated in COHFD compared to COChow females, but expression was downregulated in TCDDHFD females compared to both COHFD and TCDDChow females (Fig. 8E). These results suggest that TCDD exposure may alter diet-induced changes in beta cell specific gene expression.

Lastly, the “Alanine, Aspartate and Glutamate Metabolism” pathway heatmap showed that COHFD and TCDD-exposed females formed a separate cluster compared to COChow females, suggesting that both HFD feeding and TCDD exposure alter amino acid metabolism (Fig. 8B, Supp. Fig. 2D). Interestingly, TCDDHFD females formed a separate subcluster compared to both TCDDChow and COHFD females, suggesting that TCDD may alter HFD-induced changes in amino acid metabolism (Fig. 8B).

**Discussion**

In this study, 12 weeks of low-dose TCDD exposure alone did not impact glucose homeostasis or plasma insulin levels in male or female mice, suggesting that background level TCDD exposure may not be sufficient to induce diabetes. TCDD exposure also did not affect the metabolic response to HFD feeding in male mice. However, when challenged with HFD feeding, TCDD exposure accelerated the onset of HFD-induced hyperglycemia in female mice by at least 4 weeks. An interaction between TCDD exposure and HFD feeding was also evident in the insulin response to a glucose challenge in females. HFD feeding induced hyperinsulinemia in CO-exposed female mice and both TCDD- and CO-exposed male mice, but failed to induce compensatory hyperinsulinemia in TCDD-exposed females. RNAseq analysis showed that TCDD exposure promoted abnormal changes to endocrine and metabolic pathways in HFD-fed females.

Our study demonstrates that the effects of chronic low-dose TCDD exposure on glucose metabolism are sex-dependent. TCDD accelerated the onset of diet-induced hyperglycemia and prevented diet-induced hyperinsulinemia, but only in female mice. There were no differences in
islet size or pancreatic endocrine cell composition to explain the discrepancy in insulin levels in TCDDHFD females. Instead, our RNASeq data points to a possible defect in beta cell function and impaired metabolic adaptability in females. For example, expression of MODY genes differed between TCDDHFD and COHFD females, indicating an interaction between TCDD and HFD on beta-cell specific genes. In fact, the diet-induced decrease in MafA, Hnf4α, and Slc2a2 was worsened in TCDD-exposed females, whereas Pax6 was downregulated only in TCDDHFD compared to COChow females. Further, the compensatory increase in Nkx6.1 in COHFD females compared to COChow was absent in TCDDHFD females. These findings are in line with our previous work showing that female mice exposed to low-dose TCDD during pregnancy/lactation and subsequently fed HFD later in life had accelerated onset of hyperglycemia, transient low glucose-induced plasma insulin levels, and reduced MafA+ beta cells compared to COHFD females (24). The beta cell transcription factors MAFA, PDX1, HNF4α, and PAX6 form a complex network essential for maintaining beta cell identity and regulating genes involved in insulin secretion (e.g. Slc2a2); inactivation of these genes is associated with beta cell dedifferentiation and metabolic inflexibility (31). Under conditions of nutrient overload, beta cells must compensate for changes in metabolic demand by altering the rate and amount of insulin secreted; failure to adapt is associated with diabetes (32). Our results suggest that low-dose TCDD exposure may promote diet-induced beta cell dedifferentiation, which could contribute to impaired metabolic adaptability and increase diabetes risk in female mice, but further research is required to support this hypothesis.

RNASeq analysis also revealed unique changes in metabolic and endocrine pathways in TCDDHFD females compared to chow-fed females that further support a defect in metabolic adaptability. We focus on two pathways of interest here, however future studies should further investigate the interactive effect of TCDD and HFD on other pathways that were enriched in our study, including “Thyroid Signalling”, “Foxo1 Signaling”, “Regulation of Actin Cytoskeleton”, amongst others. Our analysis revealed changes in the “Circadian Rhythm” pathway in islets from TCDDHFD females. Clock genes were generally altered by diet, however hierarchal clustering revealed overall differences in gene expression between CO- and TCDD-exposed females fed either a chow or HFD diet. These results suggest that both TCDD and HFD alter circadian rhythm, in line with previous data (33,34), but also that TCDD may impair normal alterations to islet circadian rhythms during HFD feeding. These data are interesting given that alterations to the circadian clock has been associated with diabetes, obesity, and metabolic syndrome in humans (35–37). Abnormal expression of clock genes has also been reported in islets from T2D donors compared to non-diabetic donors, and insulin content was positively correlated with expression of clock genes, suggesting that disruptions to the islet circadian clock may contribute to beta cell dysfunction and diabetes pathogenesis (38). In fact, knockdown of Clock in human islets also led to dysfunctional glucose-stimulated insulin secretion and altered the expression of genes involved in insulin secretion (39). In addition, islet-specific Clock knockout mice and pancreas-specific Bmal1 knockout mice exhibit hypoinsulinemia, hyperglycemia, glucose intolerance, and an impaired insulin secretory response (40). Lastly, experimental disruption of circadian rhythms accelerates the onset of hyperglycemia and loss of functional beta cells in diabetes-prone rats (41). Taken together, our data suggest that the glucose intolerance and reduced plasma insulin levels in TCDDHFD female mice may involve altered islet circadian rhythms, but more detailed analysis is required to understand the mechanism involved.
Chronic low-dose TCDD exposure also promoted alterations to amino acid metabolism pathways in HFD-fed females. Exposure to TCDD has been previously associated with changes to circulating amino acid concentrations (42–44) and hepatic amino acid metabolism genes in vivo (45) and in vitro (46,47). Our data indicates that TCDD may also alter amino acid metabolism in islets under conditions of HFD feeding, which may have important implications for diabetes risk. Amino acids play an important role in maintaining beta cell function (48). Several studies have shown that amino acids enhance glucose-stimulated insulin secretion in islets ex vivo and in beta cell lines (49–51), and may indirectly regulate lipid-stimulated insulin secretion by activating enzymes involved in lipid metabolism (52,53). Amino acids have also been shown to promote the expression of genes involved in beta cell signalling, metabolism, and insulin secretion (e.g. Pdx1) (54,55). As such, perturbations to amino acid metabolism have been associated with T2D, obesity, and insulin resistance (56). Our data suggests that impaired amino acid metabolism in islets may contribute to the reduced plasma insulin levels and hyperglycemia observed in TCDDHFD female mice, but further investigation into amino acid metabolism and amino acid-stimulated insulin secretion following TCDD exposure is required.

Lastly, our RNAseq analysis revealed interesting changes in “Xenobiotic Metabolism by CYP450”. We have previously shown that acute high-dose and chronic low-dose TCDD exposure induced Cyp1a1 expression in islets from male mice (22,23). The present study demonstrates that Cyp1a1 is also upregulated in female islets following low-dose TCDD exposure. We also found that HFD feeding induces Cyp1a1 expression in islets to a similar degree as TCDD exposure, suggesting a role for Cyp1a1 in islets beyond xenobiotic metabolism. Interestingly, we have previously shown that male Cyp1a1/1a2 knockout mice had suppressed glucose-stimulated insulin secretion compared to wildtype controls, pointing to a non-conventional role of CYP enzymes in maintaining beta cell function (22). Furthermore, the TCDD- and HFD-induced increase in Cyp1a1 was suppressed in TCDDHFD females, suggesting cross-talk between TCDD and HFD feeding. Given that HFD feeding increases cytokine signalling (57), we speculate that there may be an interaction between AhR and cytokine signalling pathways in this mouse model. In fact, we’ve previously shown that exposure of human islets to high-dose TCDD in vitro induced a ~26-fold increase in Cyp1a1, which was completely prevented when cotreated with cytokines (22). Whether this impaired Cyp1a1 induction contributes to the metabolic phenotype observed in TCDDHFD females remains unclear.

To our knowledge this is the first study to do a head-to-head comparison of the effects of chronic low-dose TCDD exposure on diabetes risk in male and female mice. Our findings are consistent with epidemiological evidence showing that females with high serum pollutant levels have a higher risk of developing diabetes than males. However, few epidemiological studies examined sex-differences in the general population with background level of pollutant exposure, and most do not consider diet as a confounding variable when studying the association between pollutant exposure and diabetes risk. Our mouse study shows that 12 weeks of chronic low-dose TCDD alone does not cause diabetes but does increase susceptibility to diet-induced diabetes in female mice, emphasizing the need to study the interaction of pollutants with other environmental factors.
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Author Contributions
J.E.B. and G.M. conceived the experimental design. J.E.B., G.M., and M.P.H. wrote the manuscript. G.M., M.P.H., H.L.B., J.Z., S.F.B, K.R.C.R., A.W., R.G., J.K.B., C.Y., and J.E.B. were involved with acquisition, analysis, and interpretation of data. All authors contributed to manuscript revisions and approved the final version of the article.

Declaration of Interest
The authors declare no competing interests.

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Fig. 1 Chronic low-dose TCDD exposure does not impact body weight or fasting blood glucose levels in chow-fed or HFD-fed mice. (A) Male and female mice were exposed to corn oil or 20 ng/kg/d TCDD 2x/week for 12 weeks, and simultaneously fed either a chow diet or a 45% HFD. Body weight and blood glucose was tracked throughout the 12 week study. BW = body weight; BG = blood glucose; GTT = glucose tolerance test; GSIS = glucose-stimulated insulin secretion; ITT = insulin tolerance test. (B-C) Body weight and (D-G) blood glucose were measured weekly following a 4-hour morning fast in (B, D, E) females and (C, F, G) males. Blood glucose data is presented as (D, F) all groups compared to COChow, COHFD versus COChow, TCDDChow compared to TCDDHFD, and (E, G) area under the curve for all groups. All data are presented as mean ± SEM in line graphs or min/max values in box and whisker plots. Individual data points on box and whisker plots represent biological replicates (different mice). *p <0.05, **p <0.01, coloured stars are versus COChow. The following statistical tests were used: (B, C, D, F) two-way REML-ANOVA with Tukey’s multiple comparisons test; (E, G) two-way ANOVA with Tukey’s multiple comparisons test.
Fig. 2 TCDD exposure accelerates the onset of HFD-induced hyperglycemia in female mice. Glucose tolerance tests (GTTs) were performed after 4 and 8 weeks of TCDD exposure with/without HFD feeding (see Fig. 1A for study timeline). Blood glucose levels in (A-D) females and (E-H) males at week 4 (A, B, E, F) and 8 (C, D, G, H). (A, C, E, G) Blood glucose data is presented as (i) all groups compared to COChow, (ii) COHFD versus COChow, (iii) TCDDHFD compared to TCDDChow. All data are presented as mean ± SEM in line graphs or min/max values in box and whisker plots. Individual data points on box and whisker plots represent biological replicates (different mice). *p <0.05, **p <0.01. The following statistical tests were used: (A, C, E, G) two-way RM ANOVA with Tukey’s multiple comparison test; (B, D, F, H) two-way ANOVA with Tukey’s multiple comparison test.
Fig. 3 TCDD-exposed female mice lack an appropriate compensatory hyperinsulinemic response to HFD feeding but had normal insulin tolerance. Glucose-stimulated insulin secretion (GSIS) was assessed in vivo at week 8, and insulin tolerance (ITT) at week 5 and 9 (see Fig. 1A for study timeline). (A, B) Plasma insulin levels during a GSIS at week 8 of the study in females (A) and males (B). (C-F) Blood glucose levels during an ITT at (C, D) 5 weeks and (E, F) 9 weeks in females (C, E) and males (D, F). All data are presented as mean ± SEM in line graphs or min/max values in box and whisker plots. Individual data points on box and whisker plots represent biological replicates (different mice). *p <0.05, **p <0.01, coloured stars are versus COChow. The following statistical tests were used: (A-F) line graphs, two-way RM ANOVA with Tukey’s multiple comparison test; box and whisker plots, two-way ANOVA with Tukey’s multiple comparison test.
**Fig. 4** Female islet size and endocrine composition were unchanged by TCDD exposure. Whole pancreas was harvested at week 12 of the study for analysis by immunofluorescence staining (see **Fig. 1A** for study timeline). (**A, D**) Average islet area, (**B, E**) % islet^+^ area / islet area, (**C, F**) % glucagon^+^ area / islet area in females (**A-C**) and males (**D-F**). (**G**) Representative images of pancreas sections showing immunofluorescence staining for insulin/glucagon. Scale bar = 100 µm. All data are presented as median with min/max values. Individual data points on box and whisker plots represent biological replicates (different mice). The following statistical tests were used: (**A-F**) two-way ANOVA with Tukey’s multiple comparison test.
Fig. 5 TCDDHFD female mice have enriched endocrine-related pathways relative to COChow females. Islets were isolated from females at week 12 of the study for RNAseq analysis (n=3-4/group). (A) Venn diagram displaying differentially expressed genes (DEGs) (adjusted p < 0.05, absolute fold change ≥ 1.5) in all experimental groups relative to COChow females. (B) DAVID KEGG pathway analysis was performed on the DEGs in (A) for each experimental group. Results are displayed as a Venn diagram to show pathway overlap between experimental groups, as a bar graph showing pathway fold enrichment in each group relative to COChow (DAVID modified Fisher Extract p-value < 0.05 relative to COChow females), and as a bar graph comparing differentially expressed gene counts for significantly enriched pathways (coloured * = modified Fisher Extract p-value < 0.05 versus COChow).
Fig. 6 TCDDHFD females have inappropriate Cyp1a1 and circadian rhythm gene expression. Islets were isolated from females at week 12 of the study for RNAseq and qPCR analysis (n=3-4/group). (A-B) Hierarchal heatmaps showing expression levels of gene involved in (A) xenobiotic metabolism by CYP450 and (B) circadian rhythms. (C) Cyp1a1 expression in counts per million measured by RNAseq analysis. (D) Cyp1a1 gene expression measured by qPCR analysis. Individual data points on box and whisker plots represent biological replicates (different mice). All data are presented as median with min/max values. *p <0.05, **p <0.01. The following statistical tests were used: (C-D) two-way ANOVA with Tukey’s multiple comparison test.
Fig. 7 Chronic low-dose TCDD exposure promotes diet-induced changes in amino acid metabolism. Islets were isolated from females at week 12 of the study for RNAseq analysis (n=3-4/group). (A) Venn diagram displaying differentially expressed genes (DEGs) (adjusted p < 0.05, absolute fold change ≥ 1.5) in HFD fed females relative their respective chow-fed control female. (B) DAVID KEGG pathway analysis was performed on the DEGs in (A). Results are displayed as a Venn diagram to show pathway overlap between experimental groups, as a bar graph showing pathway fold enrichment in each comparison (DAVID modified Fisher Extract p-value < 0.05 relative to respective chow-fed control), and as a bar graph comparing differentially expressed gene counts for significantly enriched pathways (coloured * = modified Fisher Extract p-value < 0.05 versus chow-fed control).
Fig. 8 HFD-induced changes in MODY genes and alanine, aspartate, and glutamate metabolism are promoted by TCDD exposure. Islets were isolated from females at week 12 of the study for RNAseq and qPCR analysis (n=3-4/group). (A-B) Hierarchical heatmaps showing expression levels of gene involved in (A) maturity onset diabetes of the young and (B) alanine, aspartate and glutamate metabolism. (C, F-H) Gene expression of (C) MafA, (E) Nkx6.1, (F) Pax6, (G) Hnf4α, and (H) Slc2a2 in counts per million measured by RNAseq analysis. (D) MafA gene expression measured by qPCR analysis. Individual data points on box and whisker plots represent biological replicates (different mice). All data are presented as median with min/max values. *p <0.05, **p <0.01. The following statistical tests were used: (C-H) two-way ANOVA with Tukey’s multiple comparison test.