

1 **TITLE:** Sex differences in islet stress responses support female beta cell resilience

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3 **SHORT TITLE:** Sex-specific beta cell stress responses

4

5 **AUTHORS:** George P. Brownrigg<sup>1</sup>, Yi Han Xia<sup>1</sup>, Chieh Min Jamie Chu<sup>1</sup>, Su Wang<sup>1</sup>, Charlotte Chao<sup>1</sup>,  
6 Jiashuo Aaron Zhang<sup>1</sup>, Søs Skovsø<sup>1</sup>, Evgeniy Panzhinskiy<sup>1</sup>, Xiaoke Hu<sup>1</sup>, James D. Johnson<sup>1§</sup>,  
7 Elizabeth J. Rideout<sup>1§</sup>

8

9 **AFFILIATION:** <sup>1</sup>Department of Cellular and Physiological Sciences, Life Sciences Institute, The  
10 University of British Columbia, 2350 Health Sciences Mall, Vancouver, BC, V6T 1Z3, Canada

11

12 **§ CO-CORRESPONDING AUTHORS:** Department of Cellular and Physiological Sciences, Life  
13 Sciences Institute, The University of British Columbia, 2350 Health Sciences Mall, Vancouver, BC,  
14 V6T 1Z3, Canada. Email: [elizabeth.rideout@ubc.ca](mailto:elizabeth.rideout@ubc.ca) & [james.d.johnson@ubc.ca](mailto:james.d.johnson@ubc.ca) Phone: (604) 822-  
15 0623 & (604) 822-7187. Fax: (604) 822-2316.

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## 17 **AUTHOR CONTRIBUTIONS**

18 G. P. B. conceived studies, conducted experiments, interpreted experiments, wrote the manuscript

19 Y. X. performed bioinformatic analysis and data visualization

20 J. C. created custom R scripts (single-cell GFP tracking)

21 S. W. analyzed data (human RNAseq)

22 C. C. created custom R scripts (mouse RNAseq analysis)

23 J. A. Z. analyzed data (HPAP perfusions)

24 S. S. conducted experiments (*in vivo* physiology)

25 E. P. conducted experiments (islet western blots)

26 X. H. conducted experiments (dissections)

27 J. D. J. conceived studies, interpreted experiments, edited the manuscript

28 E. J. R. conceived studies, interpreted experiments, edited the manuscript, and is the guarantor of this  
29 work

30

31 **KEYWORDS:** pancreatic islets,  $\beta$  cells, diabetes mellitus, endoplasmic reticulum stress, protein  
32 synthesis, transcriptomics

33

34

35 **ABSTRACT**

36

37 **Objective:** Pancreatic  $\beta$  cells play a key role in glucose homeostasis; dysfunction of this  
38 critical cell type causes type 2 diabetes (T2D). Emerging evidence points to sex differences in  
39  $\beta$  cells, but few studies have examined male-female differences in  $\beta$  cell stress responses  
40 and resilience across multiple contexts, including diabetes. Here, we address the need for  
41 high-quality information on sex differences in  $\beta$  cell/islet gene expression and function using  
42 both human and rodent samples.

43

44 **Methods:** We compared  $\beta$  cell gene expression and insulin secretion in donors living with  
45 T2D to non-diabetic donors in both males and females. In mice, we generated a well-powered  
46 islet RNAseq dataset from 20-week-old male and female siblings with equivalent insulin  
47 sensitivity. Because our unbiased analysis of gene expression pointed to sex differences  
48 in endoplasmic reticulum (ER) stress response, we subjected islets isolated from age-  
49 matched male and female mice to thapsigargin treatment and monitored protein synthesis,  
50 cell death, and  $\beta$  cell insulin production and secretion. Transcriptomic and proteomic analyses  
51 were used to characterize sex differences in islet responses to ER stress.

52

53 **Results:** Our single-cell analysis of human  $\beta$  cells revealed sex-specific changes to gene  
54 expression and function in T2D, correlating with more robust insulin secretion in islets isolated  
55 from female donors living with T2D compared to male T2D donors. In mice, RNA sequencing  
56 revealed differential enrichment of unfolded protein response pathway-associated genes,  
57 where female islets showed higher expression of genes linked with protein synthesis, folding,  
58 and processing. This differential expression was biologically significant, as female islets were  
59 more resilient to ER stress induction with thapsigargin. Specifically, female islets maintained  
60 better insulin secretion and showed a distinct transcriptional response under ER stress  
61 compared with males.

62

63 **Conclusions:** Our data demonstrate that physiologically significant sex differences in  $\beta$  cell  
64 gene expression exist in both humans and mice, and that female  $\beta$  cells maintain better  
65 insulin production and secretion across multiple physiological and pathological contexts.

66

67 **1. Introduction**

68 Pancreatic  $\beta$  cells make and secrete insulin, an essential hormone required to maintain  
69 whole-body glucose homeostasis. Emerging evidence from multiple species suggests  
70 biological sex is an important, but often overlooked, factor that affects  $\beta$  cell biology (1–6).  
71 Large-scale surveys of gene expression in mice and humans show that differences exist  
72 between the sexes in the pancreas (7–9), in islets (10), and in  $\beta$  cells specifically (4,11).  
73 Humans also have a sex-specific  $\beta$  cell gene expression response to aging (12), and show  
74 male-female differences in pancreatic  $\beta$  cell number (6). With respect to  $\beta$  cell function, most  
75 data from rodent and human studies suggests glucose-stimulated insulin secretion is higher in  
76 females than in males (5,10,13–16). While male-female differences in peripheral insulin  
77 sensitivity (15,17–27) may contribute to these differences, sex-biased insulin secretion in  
78 humans persists in the context of equivalent insulin sensitivity between males and females  
79 (5). Whether sex differences in additional aspects of  $\beta$  cell gene expression and function  
80 similarly persist remains unclear, as insulin sensitivity is not routinely monitored across  
81 datasets showing sex differences in  $\beta$  cell biology.

82 Biological sex also affects the risk of developing T2D. Across many population groups,  
83 men are at a higher risk of developing T2D than women (28–31). Some of the differential risk  
84 is explained by lifestyle and cultural factors (31–33). Biological sex also plays a role, however,  
85 as the male-biased risk of developing diabetes-like phenotypes exists across multiple animal  
86 models (22,34–39). Despite a dominant role for  $\beta$  cell function in T2D pathogenesis (40,41),  
87 T2D- and stress-associated changes to  $\beta$  cell gene expression and function in each sex  
88 remain largely unexplored, as many studies on this topic did not include biological sex as a  
89 variable in their analysis (42–49). More detailed knowledge of  $\beta$  cell gene expression and  
90 function in physiological and pathological contexts is therefore a key first step toward  
91 understanding how sex differences in this important cell type may contribute to T2D risk.

92 The overall goal of our study was to provide detailed knowledge of  $\beta$  cell gene expression  
93 and function in both males and females across multiple contexts to advance our  
94 understanding of sex differences in this important cell type. Collectively, our data show  
95 significant sex differences in islet and  $\beta$  cell gene expression and stress responses in both  
96 humans and mice. These differences contribute to sex differences in  $\beta$  cell resilience, where  
97 we find female  $\beta$  cells maintain better insulin secretion in response to stress and T2D.  
98 Importantly, these differences cannot be fully explained by differential peripheral insulin

99 sensitivity between the sexes, suggesting biological sex is an important variable to consider in  
100 studies on islet and  $\beta$  cell function.

101

102 **2. Materials and methods**

103 **2.1. Animals**

104 Mice were bred in-house or purchased from the Jackson Laboratory. Unless otherwise stated  
105 mouse islets were isolated from C57BL/6J mice aged 20-24 weeks. Animals were housed  
106 and studied in the UBC Modified Barrier Facility using protocols approved by the UBC Animal  
107 Care Committee and in accordance with international guidelines. Mice were housed on a 12-  
108 hour light/dark cycle with food and drinking water *ad libitum*. Mice were fed a regular chow  
109 diet (LabDiet #5053); 24.5% energy from protein, 13.1% energy from fat, and 62.4% energy  
110 from carbohydrates.

111

112 **2.2. Islet Isolation, Culture, Dispersion and Treatment**

113 Mouse islet isolations were performed by ductal collagenase injection followed by filtration  
114 and hand-picking, using modifications of the protocol described by Salvalaggio (50). Islets  
115 recovered overnight, in islet culture media (RPMI media with 11.1 mM D-glucose  
116 supplemented with 10% vol/vol fetal bovine serum (FBS) (Thermo: 12483020) and 1% vol/vol  
117 Penicillin-Streptomycin (P/S) (GIBCO: 15140-148)) at 37°C with 5% CO<sub>2</sub>. After four washes  
118 with Minimal Essential Medium [L-glutamine, calcium and magnesium free] (Corning: 15-015  
119 CV) islets were dispersed with 0.01% trypsin and resuspended in islet culture media. Cell  
120 seedings were done as per the experimental procedure (protein synthesis: 20,000 cells per  
121 well, live cell imaging: 5,000 cells per well). ER stress was induced by treating islets with the  
122 SERCA inhibitor thapsigargin. For assays less than 24 hours, we used (11.1 mM D-glucose  
123 RPMI, 1% vol/vol P/S). For assays greater than 24 hours we used (11.1 mM D-glucose RPMI,  
124 1% vol/vol P/S, 10% vol/vol FBS).

125

126 **2.3. Analysis of protein synthesis**

127 Dispersed islets were seeded into an optical 96-well plate (Perkin Elmer) at a density of  
128 approximately 20,000 cells per well islet culture media (11.1 mM D-glucose RPMI, 1% vol/vol  
129 P/S, 10% vol/vol FBS). 24 hours after seeding, treatments were applied in fresh islet culture  
130 media (11.1 mM D-glucose RPMI, 1% vol/vol P/S). After incubation, fresh culture media was  
131 applied (11.1 mM D-glucose RPMI, 1% vol/vol P/S), supplemented with 20  $\mu$ M OPP

132 (Invitrogen) and respective drug treatments. The assay was performed according to  
133 manufacturer's instructions; cells were then imaged at 10x with an ImageXpress<sup>MICRO</sup> high-  
134 content imager and analyzed with MetaXpress (Molecular Devices) to quantify the integrated  
135 staining intensity of OPP-Alexa Fluor 594 in cells identified by NuclearMask Blue Stain.  
136

137 **2.4. Live cell imaging**

138 Dispersed islets were seeded into 384-well plates (Perkin Elmer) at a density of  
139 approximately 5,000 cells per well and allowed to adhere for 48 hours in islet culture media  
140 (11.1 mM D-glucose RPMI, 1% vol/vol P/S, 10% vol/vol FBS). Cells were stained with  
141 Hoechst 33342 (Sigma-Aldrich) (0.05 µg/mL) and propidium iodide (Sigma-Aldrich) (0.5  
142 µg/mL) for one hour in islet culture media (11.1 mM D-glucose RPMI, 1% vol/vol P/S, 10%  
143 vol/vol FBS) prior to the addition of treatments and imaging. 384-well plates were placed into  
144 environmentally controlled (37°C, 5% CO<sub>2</sub>) ImageXpress<sup>MICRO</sup> high content imaging system.  
145 To measure cell death, islet cells were imaged every 2 hours for 84 hours, and MetaXpress  
146 software was used to quantify cell death, defined as the number of Propidium Iodide-  
147 positive/Hoechst 33342-positive cells. To measure *Ins2* gene activity, dispersed islets from  
148 *Ins2*<sup>GFP/WT</sup> mice aged 21-23 weeks were used (51). Islet cells were imaged every 30 minutes  
149 for 60 hours. MetaXpress analysis software and custom R scripts were used to perform  
150 single-cell tracking of *Ins2*<sup>GFP/WT</sup> β cells as previously described (51).  
151

152 **2.5. Western blot**

153 After a 24-hour treatment with 1 µM Tg in islet culture media (11.1 mM D-glucose RPMI, 1%  
154 vol/vol P/S, 10% vol/vol FBS), mouse islets were sonicated in RIPA lysis buffer (150 mM  
155 NaCl, 1% Nonidet P-40, 0.5% DOC, 0.1% SDS, 50 mM Tris (pH 7.4), 2 mM EGTA, 2 mM  
156 Na<sub>3</sub>VO<sub>4</sub>, and 2 mM NaF supplemented with complete mini protease inhibitor cocktail (Roche,  
157 Laval, QC)). Protein lysates were incubated in Laemmli loading buffer (Thermo, J61337AC) at  
158 95°C for 5 minutes and resolved by SDS-PAGE. Proteins were then transferred to PVDF  
159 membranes (BioRad, CA) and probed with antibodies against HSPA5 (1:1000, Cat. #3183,  
160 Cell Signalling), eIF2α (1:1000, Cat. #2103, Cell Signalling), phospho-eIF2α (1:1000, Cat.  
161 #3398, Cell Signalling), IRE1α (1:1000, Cat. #3294, Cell Signalling), phospho-IRE1α (1:1000,  
162 Cat. #PA1-16927, Thermo Fisher Scientific), CHOP (1:1000, #ab11419, Abcam), β-actin  
163 (1:1000, NB600-501, Novus Biologicals). The signals were detected by secondary HRP-  
164 conjugated antibodies (Anti-mouse, Cat. #7076; Anti-rabbit, Cat. #7074; CST) and either

165 Pierce ECL Western Blotting Substrate (Thermo Fisher Scientific) or Forte (Immobilon).  
166 Protein band intensities were quantified using Image Studio (LI-COR).

167

168 **2.6. Islet Secretion and Content**

169 Glucose-stimulated insulin/proinsulin production and secretion were assessed using size-  
170 matched islets (five islets per well, in triplicate) seeded into 96-well V-bottom Tissue Culture  
171 Treated Microplates (Corning: #CLS3894). Islets were allowed to adhere for 48 hours in  
172 culture media (11.1 mM D-glucose RPMI, 1% vol/vol P/S, 10% vol/vol FBS). Islets were  
173 washed with Krebs-Ringer Buffer (KRB; 129 mM NaCl, 4.8 mM KCl, 1.2 mM MgSO<sub>4</sub>, 1.2 mM  
174 KH<sub>2</sub>PO<sub>4</sub>, 2.5 mM CaCl<sub>2</sub>, 5 mM NaHCO<sub>3</sub>, 10 mM HEPES, 0.5% bovine serum albumin)  
175 containing 3 mM glucose then pre-incubated for 4 hours in 3 mM glucose KRB. 1 µM Tg was  
176 added to the 3 mM low glucose pre-incubation buffer 4 hours prior, 2 hours prior, or at the  
177 start of the low glucose incubation period. Islets were incubated in KRB with 3 mM glucose  
178 then 20 mM glucose for 45 minutes each. Supernatant was collected after each stimulation.  
179 Islet insulin and proinsulin content was extracted by freeze-thawing in 100 µL of acid ethanol,  
180 then the plates were shaken at 1200 rpm for 10 minutes at 4°C to lyse the islets. Insulin was  
181 measured by Rodent Insulin Chemiluminescent ELISA (ALPCO: 80-INSMR) and proinsulin by  
182 Rat/Mouse Proinsulin ELISA (Mercodia: 10-1232-01). Measurements were performed on a  
183 Spark plate reader (TECAN).

184

185 **2.7. Blood collection and in vivo analysis of glucose homeostasis and insulin secretion**

186 Mice were fasted for 6 hours prior to glucose and insulin tolerance tests. During glucose and  
187 insulin tolerance tests, tail blood was collected for blood glucose measurements using a  
188 glucometer (One Touch Ultra 2 Glucometer, Lifescan, Canada). For intraperitoneal (i.p.)  
189 glucose tolerance tests, the glucose dose was 2 g glucose/kg of body mass. For insulin  
190 tolerance tests, the insulin dose was 0.75U insulin/kg body mass. For measurements of *in*  
191 *vivo* glucose-stimulated insulin secretion, femoral blood was collected after i.p. injection of 2 g  
192 glucose/kg body mass. Blood samples were kept on ice during collection, centrifuged at 2000  
193 rpm for 10 minutes at 4°C and stored as plasma at -20°C. Plasma samples were analysed for  
194 insulin using Rodent Insulin Chemiluminescent ELISA (ALPCO: 80-INSMR).

195

196 **2.8. RNA sequencing**

197 To assess basal transcriptional differences islets from male and female mice (n=9, 8) were  
198 snap frozen and stored at -80°C until RNA extraction. To assess Tg-induced transcriptional  
199 changes islets from each mouse were treated with DMSO or Tg for 6- or 12-hours in culture  
200 media (11.1 mM D-glucose RPMI, 1% vol/vol P/S) (8 groups, n=3-4 per group, each n  
201 represents pooled islet RNA from two mice). Islets were frozen at -80°C in 100 µL of RLT  
202 buffer (Qiagen) with beta mercaptoethanol (1%). RNA was isolated using RNeasy Mini Kit  
203 (Qiagen #74106) according to manufacturer's instructions. RNA sequencing was performed at  
204 the UBC Biomedical Research Centre Sequencing Core. Sample quality control was  
205 performed using the Agilent 2100 Bioanalyzer System (RNA Pico LabChip Kit). Qualifying  
206 samples were prepped following the standard protocol for the NEBNext Ultra II Stranded  
207 mRNA (New England Biolabs). Sequencing was performed on the Illumina NextSeq 500 with  
208 Paired End 42bp × 42bp reads. Demultiplexed read sequences were then aligned to the  
209 reference sequence (UCSC mm10) using STAR aligner (v 2.5.0b) (52). Gene differential  
210 expression was analyzed using DESeq2 R package (53). Pathway enrichment analysis were  
211 performed using Reactome (54). Over-representation analysis was performed using  
212 NetworkAnalyst3.0 ([www.networkanalyst.ca](http://www.networkanalyst.ca)) (55).  
213

## 214 **2.9. Proteomics**

215 Islets were treated with DMSO or Tg for 6 hours in islet culture media (11.1 mM D-glucose  
216 RPMI, 1% vol/vol P/S) (4 groups, n=5-7 per group, each n represents pooled islets from two  
217 mice). Islet pellets were frozen at -80°C in 100 µL of SDS lysis buffer (4% SDS, 100 mM Tris,  
218 pH 8) and the proteins in each sample were precipitated using acetone. University of Victoria  
219 proteomics service performed non-targeted quantitative proteomic analysis using data-  
220 independent acquisition (DIA) with LC-MS/MS on an Orbitrap mass spectrometer. A mouse  
221 FASTA database was downloaded from Uniprot (<http://uniprot.org>). This file was used with  
222 the 6 gas phase fraction files from the analysis of the chromatogram library sample to create  
223 a mouse islet specific chromatogram library using the EncyclopeDIA (v 1.2.2) software  
224 package (Searle et al, 2018). This chromatogram library file was then used to perform  
225 identification and quantitation of the proteins in the samples again using EncyclopeDIA with  
226 Overlapping DIA as the acquisition type, trypsin used as the enzyme, CID/HCD as the  
227 fragmentation, 10 ppm mass tolerances for the precursor, fragment, and library mass  
228 tolerances. The Percolator version used was 3.10. The precursor FDR rate was set to 1%.  
229 Protein abundances were log2 transformed, imputation was performed for missing values,

230 then proteins were normalized to median sample intensities. Gene differential expression was  
231 analyzed using limma in Perseus (56).

232

233 **2.10. Data from HPAP**

234 To compare sex differences in dynamic insulin secretion, data acquired was from the Human  
235 Pancreas Analysis Program (HPAP-RRID:SCR\_016202) Database  
236 (<https://hpap.pmacs.upenn.edu>), a Human Islet Research Network (RRID:SCR\_014393)  
237 consortium (UC4-DK-112217, U01-DK-123594, UC4-DK-112232, and U01-DK-123716).

238

239 **2.11. Statistical Analysis**

240 Statistical analyses and data presentation were carried out using GraphPad Prism 9  
241 (GraphPad Software, San Diego, CA, USA) or R (v 4.1.1). Student's *t*-tests or two-way  
242 ANOVAs were used for parametric data. A Mann-Whitney test was used for non-parametric  
243 data. Statistical tests are indicated in the figure legends. For all statistical analyses,  
244 differences were considered significant if the p-value was less than 0.05. \*: p<0.05; \*\* p<0.01;  
245 \*\*\* p<0.001. Data were presented as means ± SEM with individual data points from biological  
246 replicates.

247

248

249 **3. Results**

250 **3.1. Sex differences in  $\beta$  cell transcriptional and functional responses in ND and T2D  
251 human islets**

252 To define  $\beta$  cell-specific gene expression changes in T2D in each sex, we used a recently  
253 compiled meta-analysis of publicly available scRNASeq datasets from male and female  
254 human islets (57). In line with prior reports (12), non-diabetic (ND) and T2D  $\beta$  cells showed  
255 significant transcriptional differences. In  $\beta$  cells isolated from female T2D donors, mRNA  
256 levels of 127 genes were significantly different from ND female donors (77 downregulated, 50  
257 upregulated in T2D) (Figure 1A-C). In  $\beta$  cells isolated from male T2D donors, 462 genes were  
258 differentially expressed compared with male ND donors (138 downregulated, 324 upregulated  
259 in T2D) (Figure 1A-C). Of the 660 genes that were differentially regulated in T2D, 71 were  
260 differentially regulated in both males and females (15 downregulated, 56 upregulated in T2D)  
261 (Figure 1A-C); however, the fold change for these 71 shared genes was different between  
262 males and females (Figure S1A; Supplementary file 1). This suggests that for shared genes,

263 the magnitude of gene expression changes in T2D was not the same between the sexes.  
264 Beyond shared genes, we observed that the majority of differentially expressed genes in T2D  
265 (589/660) were unique to either males or females (Figure S1B, C; Supplementary file 1).  
266 Given that the most prominent gene expression changes in T2D were found in genes that  
267 were unique to one sex (Figure S2A, B; Supplementary file 1), these data suggest there are  
268 important sex differences in the  $\beta$  cell gene expression response to T2D.

269 To determine which biological pathways were altered in  $\beta$  cells of T2D donors in each sex,  
270 we performed pathway enrichment analysis. Genes that were upregulated in  $\beta$  cells isolated  
271 from T2D donors included genes involved in Golgi-ER transport and the unfolded protein  
272 response (UPR) pathways (Figure 1D-F; Supplementary file 1). While these biological  
273 pathways were significantly upregulated in T2D in both males and females, ~75% of the  
274 differentially regulated genes in these categories were unique to each sex (Table 1). Genes  
275 that were downregulated in  $\beta$  cells from T2D donors revealed further differences between the  
276 sexes: biological pathways downregulated in  $\beta$  cells from female T2D donors included cellular  
277 responses to stress and to stimuli (Figure 1E; Supplementary file 1), whereas  $\beta$  cells from  
278 male T2D donors showed downregulation of pathways associated with respiratory electron  
279 transport and translation initiation (Figure 1F; Supplementary file 1). Thus, our analysis  
280 suggests that sex-biased  $\beta$  cell gene expression responses to T2D may influence different  
281 cellular processes in males and females.

282 The sex-biased  $\beta$  cell transcriptional response in T2D prompted us to compare glucose-  
283 stimulated insulin secretion in each sex from ND and T2D human islets using data from the  
284 Human Pancreas Analysis Program database (58). In ND donors, islets from males and  
285 females showed similar patterns of insulin secretion in response to various stimulatory media  
286 (Figure 1G, H). In donors living with T2D, we found that insulin secretion was impaired to a  
287 greater degree in islets from males than in females (Figure 1G-K). Indeed, in male but not  
288 female islets, insulin secretion was significantly lower in donors with T2D following stimulation  
289 with both high glucose and IBMX (Figure 1I, J), which potentiates insulin secretion by  
290 increasing cAMP levels to a similar degree as the incretins (59). Human islets from female  
291 donors living with T2D therefore show better  $\beta$  insulin release than islets from males living  
292 with T2D (Figure 1K). Indeed, while diabetes status was the main donor characteristic that  
293 correlated with changes in insulin secretion (Figure S3A), we noted that in T2D sex and age  
294 were two donor characteristics showing trends toward an effect on insulin secretion (Figure

295 S3B). Combined with our  $\beta$  cell gene expression data, these findings suggest that  $\beta$  cell  
296 transcriptional and functional responses in T2D are not shared between the sexes.  
297

### 298 **3.2. Sex differences in UPR-associated gene expression in mouse islets**

299 Our unbiased analysis of human  $\beta$  cell gene expression and function in T2D revealed  
300 differences between male and female donors living with T2D. Because human  $\beta$  cell gene  
301 expression and function can be affected by factors such as peripheral insulin sensitivity,  
302 disease processes, and medication (31,33), we investigated sex differences in  $\beta$  cell gene  
303 expression and function in another context. We generated a well-powered islet RNAseq  
304 dataset from 20-week-old male and female C57BL/6J mice, an age where we show insulin  
305 sensitivity is equivalent between the sexes (Figure S4A). Principal component analysis and  
306 unsupervised clustering clearly separated male and female islets on the basis of gene  
307 expression (Figure 2A; Figure S5A). We found that 17.7% (3268/18938) of genes were  
308 differentially expressed between the sexes (1648 upregulated in females, 1620 upregulated in  
309 males), in line with estimates of sex-biased gene expression in other tissues (60,61).  
310 Overrepresentation and pathway enrichment analysis both identified UPR-associated  
311 pathways as a biological process that differed significantly between the sexes, where the  
312 majority of genes in this category were enriched in female islets (Figure 2B, C;  
313 Supplementary file 2). Additional genes that were enriched in female islets were those  
314 associated with the gene ontology term “Cellular response to ER stress” (GO:0034976),  
315 which included many genes involved in regulating protein synthesis (Figure 2D). For example,  
316 females showed significantly higher levels of most ribosomal protein genes (Figure 2E).  
317 Further genes enriched in females included those associated with protein folding, protein  
318 processing, and quality control (Figure 2D). Given that protein synthesis, processing, and  
319 folding capacity are intrinsically important for multiple islet cell types (62–65), including  $\beta$  cells  
320 (66,67), this suggests female islets may have a larger protein production and folding capacity  
321 than male islets.  
322

### 323 **3.3. Female islets are more resilient to endoplasmic reticulum stress in mice**

324 The burden of insulin production causes endoplasmic reticulum (ER) stress in  $\beta$  cells (68–70).  
325 ER stress is associated with an attenuation of mRNA translation (71), and, if ER stress is  
326 prolonged, can lead to cell death (72–74). Given that female islets exhibited higher  
327 expression of genes associated with protein synthesis, processing, and folding than males,

328 and higher expression of genes associated with the UPR, which is activated in response to  
329 ER stress (75), we examined global protein synthesis rates in male and female islets under  
330 basal conditions and under ER stress. We incubated islets with O-propargyl-puromycin  
331 (OPP), which is incorporated into newly-translated proteins and can be ligated to a  
332 fluorophore. Using this technique, we monitored the accumulation of newly-synthesized islet  
333 proteins with single-cell resolution (Figure S6A). In basal culture conditions, male islet cells  
334 had significantly greater protein synthesis rates compared with female islet cells (Figure S6B).  
335 To investigate islet protein synthesis under ER stress in each sex, we treated islets with  
336 thapsigargin (Tg), a specific inhibitor of the sarcoplasmic/endoplasmic reticulum  $\text{Ca}^{2+}$ -ATPase  
337 (SERCA) that induces ER stress and the UPR by lowering ER calcium levels (72,76). At 2  
338 hours post-Tg treatment, protein synthesis was repressed in both male and female islet cells  
339 (Figure 3A, B; Figure S6C). At 24 hours post-Tg treatment, we found that protein synthesis  
340 was restored to basal levels in female islet cells, but not in male islet cells (Figure 3A, B;  
341 Figure S6C). Importantly, recovery from protein synthesis repression was significantly  
342 different in males and females (sex:treatment interaction  $p=0.0115$ ). This suggests that while  
343 protein synthesis repression associated with ER stress was transient in female islets, this  
344 phenotype persisted for longer in male islets. Because insulin biosynthesis accounts for  
345 approximately half the total protein production in  $\beta$  cells (77), one potential explanation for the  
346 sex-specific recovery from protein synthesis repression is a sex difference in transcriptional  
347 changes to insulin. To test this, we quantified GFP levels in  $\beta$  cells isolated from male and  
348 female mice with GFP knocked into the endogenous mouse *Ins2* locus (*Ins2*<sup>GFP/WT</sup>) (51,78).  
349 While ER stress induced a significant reduction in *Ins2* gene activity, this response was  
350 equivalent between the sexes. This suggests *Ins2* transcriptional changes cannot fully explain  
351 the sex difference in recovery from protein synthesis repression during ER stress (Figure S7).

352 Given the prolonged protein synthesis repression in males following ER stress, we next  
353 quantified cell death, another ER stress-associated phenotype (75), in male and female islets.  
354 Using a kinetic cell death analysis, we observed clear sex differences in Tg-induced cell death  
355 at 0.1  $\mu\text{M}$  and 1.0  $\mu\text{M}$  Tg doses throughout the time course of the experiment (Figure 3C, D).  
356 After 84 hours of Tg treatment, no significant increase in female islet cell apoptosis was  
357 observed with either 0.1  $\mu\text{M}$  or 1.0  $\mu\text{M}$  Tg treatment compared with controls (Figure 3E). In  
358 contrast, cell death was significantly increased at both the 0.1  $\mu\text{M}$  and the 1.0  $\mu\text{M}$  doses of Tg  
359 in male islet cells compared with vehicle-only controls (Figure 3F). Importantly, our analysis  
360 shows the magnitude of Tg-induced cell death was larger in male islet cells compared with

361 female islet cells (sex:treatment interaction  $p=0.0399$  [0.1  $\mu$ M],  $p=0.0007$  [1.0  $\mu$ M]). While one  
362 possible explanation for these data is that female islets are resistant to Tg-induced cell death,  
363 we found a significant increase in apoptosis in both female and male islet cells treated with 10  
364  $\mu$ M Tg (Figure 3G, H, sex:treatment interaction  $p=0.0996$  [0.1  $\mu$ M]). This suggests female  
365 islets were more resilient to mild ER stress caused by low-dose Tg than male islets.

366 To determine whether this increased ER stress resilience was caused by differential UPR  
367 signaling, we monitored levels of several protein markers of UPR activation including binding  
368 immunoglobulin protein (BiP), phosphorylated inositol-requiring enzyme 1 (pIRE1),  
369 phosphorylated eukaryotic initiation factor alpha (p $\epsilon$ IF2 $\alpha$ ), and C/EBP homologous protein  
370 (CHOP) (79,80) after treating male and female islets with 1  $\mu$ M Tg for 24 hours. We found no  
371 sex difference in UPR protein markers between male and female islets without Tg treatment  
372 (Figure S8A-D) and observed a significant increase in levels of pIRE1 $\alpha$  and CHOP in islets  
373 from both sexes and BiP in female islets after a 24-hour Tg treatment (Figure S8A-D). Lack of  
374 a sex difference in protein markers suggests UPR activation by Tg treatment was similar  
375 between male and female islets at 20 weeks of age. While this finding differs from other  
376 studies showing male-biased UPR activation (37), we reproduced the male-biased induction  
377 of BiP in islets isolated from 60-week-old male and female mice (Figure S8E-G), suggesting  
378 that age plays a role in the sex difference in UPR activation. Together, our data indicate that  
379 despite equivalent UPR activation in male and female islets treated with Tg, significant sex  
380 differences exist in ER stress-associated protein synthesis repression and cell death.

381

### 382 **3.4. Female islets retain greater $\beta$ cell function during ER stress in mice**

383 We next examined glucose-stimulated insulin secretion in islets cultured under basal  
384 conditions and after Tg treatment (Figure 4A). In all conditions tested, high glucose  
385 significantly stimulated insulin secretion in both sexes (Figure S9A); however, we identified  
386 sex differences in how well islets sustained glucose-stimulated insulin secretion during longer  
387 Tg treatments (Figure 4B, C, Figure S9A). Female islets, in both low and high glucose,  
388 maintained robust insulin secretion during Tg treatment (Figure 4B). Specifically, we observed  
389 a significant increase in insulin secretion after short Tg treatment (0 and 2 hours post-Tg),  
390 with a return to basal secretion levels 4 hours post-Tg (Figure 4B). In contrast, male islets  
391 showed no significant increase in insulin secretion after short Tg treatment, and there was a  
392 significant drop in insulin secretion at 4 hours post-Tg treatment (Figure 4C). This suggests  
393 female islets sustained insulin secretion for a longer period than male islets during ER stress.

394 Given that insulin content measurements showed insulin content significantly increased  
395 during the 4-hour Tg treatment in female islets, but not male islets (Figure 4D), our data  
396 suggest one reason female islets maintain insulin secretion during ER stress is by  
397 augmenting islet insulin content. Proinsulin secretion followed similar trends to what we  
398 observed with insulin secretion (Figure S9B), but Tg treatment reduced proinsulin content to a  
399 greater degree in male islets (Figure 4E). This suggests that in addition to females  
400 maintaining better insulin secretion during ER stress, they also show a larger increase in  
401 insulin content and a smaller decrease in proinsulin content.

402 To determine whether female islets show improved  $\beta$  cell function under ER stress in other  
403 contexts, we next monitored glucose-stimulated insulin secretion and glucose tolerance in  
404 mice at 20 weeks, an age where we show insulin sensitivity was equivalent between the  
405 sexes (Figure 4F-H; Figure S4). Despite higher fasting plasma insulin levels in males (Figure  
406 4F), and similar glucose tolerance (Figure 4H), we found that the magnitude of glucose-  
407 stimulated insulin secretion was greater in females (Figure 4G). Given that ER stress exists  
408 even in normal physiological conditions due to the burden of insulin production (81), this adds  
409 further support to a model in which female  $\beta$  cells maintain better insulin production than male  
410  $\beta$  cells under ER stress.

411

### 412 **3.5. Sex differences in islet transcriptional and proteomic responses to ER stress in 413 mice**

414 To gain insight into the differential ER stress-associated phenotypes in male and female  
415 islets, we investigated global transcriptional changes after either a 6- or 12-hour Tg treatment  
416 in each sex. Principal component analysis and unsupervised clustering shows that islets  
417 clustered by sex, treatment, and treatment time (Figure 5A; Figure S10A). The majority of the  
418 variance was explained by treatment (Figure 5B), and pathway enrichment analysis confirms  
419 the UPR as the top upregulated pathway in Tg-treated male and female islets at both 6- and  
420 12-hours after treatment (Figure S11A, B; Supplementary file 3). While some UPR-associated  
421 genes differentially regulated by Tg treatment were shared between the sexes (6-hour: 29/36,  
422 12-hour: 25/31), biological sex explained a large proportion of variance in the gene  
423 expression response to ER stress. This suggests the transcriptional response to ER stress  
424 was not fully shared between the sexes. Indeed, after a 6-hour Tg treatment, 32.6%  
425 (2247/4655) of genes that were differentially expressed between DMSO and Tg were unique

426 to one sex (881 to females, 1376 to males). After a 12-hour Tg treatment, 29% (2259/7785)  
427 were unique to one sex (1017 to males, 1242 to females).

428 To describe the transcriptional response of each sex to Tg treatment in more detail, we  
429 used a two-way ANOVA to identify genes that were upregulated, downregulated, or  
430 unchanged in male and female islets between 6- and 12-hours post-Tg (Supplementary file  
431 4). By performing pathway enrichment analysis, we were able to determine which processes  
432 were shared, and which processes differed, between the sexes during Tg treatment. For  
433 example, we observed a significant increase in mRNA levels of genes corresponding to  
434 pathways such as cellular responses to stimuli, stress, and starvation in both male and female  
435 islets between 6- and 12-hour Tg treatments (Figure 5C; Supplementary file 4), suggesting Tg  
436 has similar effects on genes related to these pathways in both sexes. In contrast, there was a  
437 male-specific increase in mRNA levels of genes associated with translation during Tg  
438 treatment (Figure 5C; Supplementary file 4). In females, there was a decrease in mRNA  
439 levels of genes associated with  $\beta$  cell identity, such as *Pklr*, *Rfx6*, *Hnf4a*, *Slc2a2*, *Pdx1*, and  
440 *MafA* (Figure S12A), and in genes linked with regulation of gene expression in  $\beta$  cells (Figure  
441 5C). Neither of these categories were altered between 6- and 12-hour Tg treatments in males  
442 (Figure 5C; Figure S12B). While this data suggests some aspects of the gene expression  
443 response to ER stress were shared between the sexes, we found that many genes  
444 corresponding to important cellular processes were differentially regulated during Tg  
445 treatment in only one sex.

446 Beyond sex-specific transcriptional changes following Tg treatment, ER stress also had a  
447 sex-specific effect on the islet proteome. Although the majority of proteins were  
448 downregulated by Tg treatment due to the generalized repression of protein synthesis under  
449 ER stress (Figure 5D), we identified 47 proteins (35 downregulated, 12 upregulated in Tg)  
450 that were differentially expressed in female islets and 82 proteins (72 downregulated, 10  
451 upregulated in Tg) that were differentially expressed after Tg treatment in male islets  
452 (Supplementary table 1). Proteins downregulated only in females include proteins associated  
453 with GO term 'endoplasmic reticulum to Golgi vesicle-mediated transport' (GO:0006888)  
454 (BCAP31, COG5, COG3, GOSR1), whereas proteins downregulated only in males include  
455 proteins associated with GO terms 'insulin secretion' (GO:0030073) (PTPRN2, CLTRN,  
456 PTPRN) and 'lysosome pathway' (KEGG) (NPC2, CTSZ, LAMP2, PSAP, CLTA). Importantly,  
457 only seven differentially expressed proteins were in common between the sexes (Figure 5D).

458 This suggests that as with our phenotypic and transcriptomic data, the proteomic response to  
459 Tg treatment was largely not shared between the sexes.

460

461

#### 462 **4. Discussion**

463 Emerging evidence shows biological sex affects many aspects of  $\beta$  cell gene expression and  
464 function. Yet, many studies on  $\beta$  cells do not include both sexes, or fail to analyze male and  
465 female data separately. To address this gap in knowledge, the goal of our study was to  
466 provide detailed information on sex differences in islet and  $\beta$  cell gene expression and  
467 function in multiple contexts. In humans, we used a large scRNASeq dataset from ND and  
468 T2D donors to reveal significant male-female differences in the magnitude of gene expression  
469 changes, and in the identity of genes that were differentially regulated, between ND and T2D  
470 donors. This suggests  $\beta$  cell gene expression changes in T2D are not fully shared between  
471 the sexes. Given that our analysis shows  $\beta$  cells from female donors living with T2D maintain  
472 better insulin production than  $\beta$  cells from male donors living with T2D, our findings suggest  
473 female  $\beta$  cells are more resilient than male  $\beta$  cells in the context of T2D. In mice, our  
474 unbiased analysis of gene expression in islets from males and females with equivalent insulin  
475 sensitivity revealed sex differences in genes associated with the UPR under normal  
476 physiological conditions. This differential gene expression was significant, as female islets  
477 were more resilient to phenotypes caused by ER stress and UPR activation than male islets,  
478 showed sex-specific transcriptional and proteomic changes in this context, and maintained  
479 better insulin secretion. Collectively, these data suggest that in rodents,  $\beta$  cells from females  
480 are more resilient to ER stress. Considering the well-established links between ER stress and  
481 T2D (79,82–84), our data suggests a model in which female  $\beta$  cells maintain better function in  
482 T2D because they are more resilient to ER stress and UPR activation. While future studies  
483 are needed to test this model, and to assess the relative contribution of sex differences in  $\beta$   
484 cells to the sex-biased risk of T2D, our findings highlight the importance of including both  
485 sexes in islet and  $\beta$  cell studies.

486 With respect to gene expression, including both sexes in our analysis of  $\beta$  cell gene  
487 expression in human ND and T2D allowed us to uncover genes that were differentially  
488 regulated in T2D in each sex. Because many of these genes may have been missed if the  
489 scRNASeq data was not analyzed by sex, our findings advance knowledge of  $\beta$  cell changes  
490 in T2D by identifying additional genes that are differentially regulated in this context. This

491 knowledge adds to a growing number of studies that identify sex differences in  $\beta$  cell gene  
492 expression during aging in humans (12), and in mice fed either a normal (4,11) or a high fat  
493 diet (11). Further, given that our RNAseq on islets from male and female mice with equivalent  
494 insulin sensitivity identifies genes and biological pathways that align with previous studies on  
495 sex differences in murine  $\beta$  cell gene expression (4,11), our data suggests that sex  
496 differences in islet and  $\beta$  cell gene expression cannot be explained solely by a male-female  
497 difference in peripheral insulin resistance. Instead, there is likely a basal sex difference in  $\beta$   
498 cell gene expression that forms the foundation for sex-specific transcriptional responses to  
499 perturbations such as ER stress and T2D. By generating large gene expression datasets from  
500 islet from male and female mice with equivalent peripheral insulin sensitivity and from islets  
501 subjected to pharmacological induction of ER stress, our studies provide a foundation of  
502 knowledge for future studies aimed at studying the causes and consequences of sex  
503 differences in islet ER stress responses and  $\beta$  cell function following UPR activation. This will  
504 provide deeper mechanistic insight into the sex-specific phenotypic effects reported in animal  
505 models of  $\beta$  cell dysfunction (35–39,85–88) and the sex-biased risk of diseases such as T2D  
506 that are associated with  $\beta$  cell dysfunction (12,22,89,90).

507 Beyond gene expression, our sex-based analysis of mouse islets allowed us to uncover  
508 male-female differences in ER stress-associated phenotypes (e.g. protein synthesis  
509 repression, cell death). While previous studies identify a sex difference in  $\beta$  cell loss in  
510 diabetic mouse models (37,39,91), and show that estrogen plays a protective role via  
511 estrogen receptor  $\alpha$  (ER $\alpha$ ) against ER stress to preserve  $\beta$  cell mass and prevent apoptosis in  
512 cell lines, mouse models, and human islets (39,91,92), we extend prior findings by showing  
513 that differences in ER stress-induced cell death were present in the context of equivalent  
514 insulin sensitivity between the sexes. This suggests sex differences in ER stress-associated  
515 phenotypes occur prior to male-female differences in peripheral insulin sensitivity. Indeed,  
516 islets isolated from males and females with equivalent sensitivity also show a sex difference  
517 in protein synthesis repression, a classical ER stress-associated phenotype (75). While  
518 estrogen affects insulin biosynthesis via ER $\alpha$  (93), future studies will need to determine  
519 whether estrogen also allows female islets to restore protein synthesis to basal levels faster  
520 than male islets following ER stress. We currently lack this knowledge, as most studies on  
521 UPR-mediated recovery from protein translation repression use single- and mixed-sex animal  
522 groups, or cultured cells (94–99).

523 Assessing whether the recovery of protein synthesis contributes to reduced cell death in  
524 female islets following ER stress will also be an important task for future studies, as prior  
525 studies suggest the inability to recover from protein synthesis repression increases ER-stress  
526 induced apoptosis (94). Ideally, this type of study would also monitor the activity of pathways  
527 known to regulate protein synthesis repression during ER stress. For example, while we did  
528 not detect any changes in levels of phosphorylated eIF2 $\alpha$  (also known as Eif2s1), which is  
529 known to mediate UPR-induced protein synthesis repression (75), our chosen timepoints did  
530 not overlap with the rapid changes in phospho-eIF2 $\alpha$  following ER stress published in other  
531 studies (100,101). A more detailed time course will therefore be necessary to assess p-eIF2 $\alpha$   
532 levels during ER stress in both sexes, and to test a role for phospho-eIF2 $\alpha$  in mediating  
533 differences in protein synthesis repression. Ultimately, a better understanding of sex  
534 differences in ER stress-associated phenotypes in  $\beta$  cells will provide a mechanistic  
535 explanation for the strongly male-biased onset of diabetes-like phenotypes in mouse models  
536 of  $\beta$  cell ER stress (e.g. Akita, KINGS, Munich mice) (37,38,85). Given the known relationship  
537 between ER stress,  $\beta$  cell death, and T2D, studies on the male-female difference in  $\beta$  cell ER  
538 stress-associated phenotypes may also advance our understanding of the male-biased risk of  
539 developing T2D in some population groups.

540 A further benefit of additional studies on the sex difference in  $\beta$  cell ER stress responses  
541 will be to identify mechanisms that support  $\beta$  cell insulin production. In rodents, we found that  
542 female islets maintained high glucose-stimulated insulin secretion and increased insulin  
543 content following ER stress, whereas male islets showed significant repression of high  
544 glucose-stimulated insulin secretion under the same conditions. In humans, while a study  
545 using a mixed-sex group of T2D donors shows  $\beta$  cells experience ER stress associated with  $\beta$   
546 cell dysfunction (102), we found that changes to  $\beta$  cell insulin secretion in T2D were not the  
547 same between the sexes. Specifically, the magnitude of the reduction in insulin release by  $\beta$   
548 cells from female donors living with T2D was smaller than in  $\beta$  cells from male donors living  
549 with T2D. Together with our data from rodents, this suggests female  $\beta$  cells maintain  
550 enhanced insulin production and/or secretion in multiple contexts, and the increased  $\beta$  cell  
551 function cannot be solely attributed to a sex difference in peripheral insulin sensitivity.

552 Clues into potential ways that female  $\beta$  cells maintain improved insulin production and  
553 secretion emerge from our examination of the transcriptional response to ER stress in each  
554 sex. Our data shows that Tg treatment induces gene expression changes characteristic of ER  
555 stress (103), and revealed similar biological pathways that were upregulated in T2D donors.

556 Furthermore, we identified significant differences between male and female islets in the  
557 transcriptional response to ER stress over time. One notable finding was that a greater  
558 number of  $\beta$  cell identity genes were downregulated between 6- and 12-hour Tg treatments in  
559 females, but not in males. Because most studies on the relationship between  $\beta$  cell identity  
560 and function used a mixed-sex pool of islets and  $\beta$  cells (68,104,105), more studies will be  
561 needed to test whether there are sex-specific changes to  $\beta$  cell identity during ER stress, and  
562 to determine the functional consequences of this sex-specific effect. Ultimately, a better  
563 understanding of changes to  $\beta$  cell gene expression and function in males and females will  
564 suggest effective ways to reverse disease-associated changes to this important cell type in  
565 each sex, improving equity in health outcomes (106).

566

#### 567 **4.1. Conclusions**

568 Our study reports significant sex differences in islet and  $\beta$  cell gene expression and stress  
569 responses in both humans and mice. These differences likely contribute to sex differences in  
570  $\beta$  cell resilience, allowing female  $\beta$  cells to maintain better insulin production across multiple  
571 contexts. This knowledge forms a foundation for future studies aimed at understanding how  
572 sex differences  $\beta$  cell function affect physiology and the pathophysiology of diseases such as  
573 T2D.

574

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590

591

## 592 **Conflict of interest**

593 The authors declare no competing interest.

594

595

## 596 **Data Availability**

597 Details of all statistical tests and *p*-values are provided in Supplementary file 5. All raw data  
598 generated in this study are available in Supplementary file 6. RNAseq data is available in  
599 Supplementary file 7 and Supplementary file 8.

600

601

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## 894 **FIGURE LEGENDS**

895 **Figure 1. Sex differences in human islet transcriptomic and functional responses in**  
896 **type 2 diabetes.** scRNAseq data from male and female human β cells. For donor metadata  
897 see Supplementary file 6. (A-C) Venn diagrams compare the number of significantly  
898 differentially expressed genes between ND and T2D donors ( $p\text{-adj}<0.05$ ). All differentially  
899 expressed genes (A), downregulated genes (B), upregulated genes (C) in T2D human β cells.

900 For complete gene lists see Supplementary file 1. (D-F) Top 10 significantly enriched  
901 Reactome pathways (ND vs T2D) from non-sex-specific (D), female (E), or male (F)  
902 significantly differentially expressed genes ( $p\text{-adj} < 0.05$ ). Gene ratio is calculated as  $k/n$ ,  
903 where  $k$  is the number of genes identified in each Reactome pathway, and  $n$  is the number of  
904 genes from the submitted gene list participating in any Reactome pathway. For complete  
905 Reactome pathway lists see Supplementary file 1. (G-K) Human islet perfusion data from the  
906 Human Pancreas Analysis Program in ND and T2D donor islets in females (F, I) and males  
907 (G, H). 3 mM glucose (3 mM G); 16.7 mM glucose (16.7 mM G); 0.1 mM  
908 isobutylmethylxanthine (0.1 mM IBMX); 30 mM potassium chloride (30 mM KCl); 4 mM amino  
909 acid mixture (4 mM AAM; mM: 0.44 alanine, 0.19 arginine, 0.038 aspartate, 0.094 citrulline,  
910 0.12 glutamate, 0.30 glycine, 0.077 histidine, 0.094 isoleucine, 0.16 leucine, 0.37 lysine, 0.05  
911 methionine, 0.70 ornithine, 0.08 phenylalanine, 0.35 proline, 0.57 serine, 0.27 threonine,  
912 0.073 tryptophan, and 0.20 valine, 2 mM glutamine). (I-K) Quantification of area under the  
913 curve (AUC) is shown for the various stimulatory media in females (I), males (J) and donors  
914 with T2D (K). (I) In females, insulin secretion from ND islets was not significantly higher than  
915 T2D islets. (J) In males, insulin secretion from ND islets was significantly higher than T2D  
916 islets under 4 mM AAM +16.7 mM glucose (HG) + 0.1 mM IBMX stimulation ( $p=0.0442$ ;  
917 unpaired Student's  $t$ -test). (K) Total insulin secretion was lower in T2D male islets than ND  
918 male islets ( $p=0.0503$ ; unpaired Student's  $t$ -test). \* indicates  $p < 0.05$ ; ns indicates not  
919 significant; error bars indicate SEM.  
920

921 **Figure 2. Sex-biased gene expression in mouse islet bulk RNAseq.** (A) Principal  
922 component analysis (PCA) of RNAseq data from male and female mouse islets. (B) Over-  
923 representation analysis (ORA) of all significantly differentially expressed genes ( $p\text{-adj} < 0.01$ )  
924 from male and female mouse islets. Top 30 enriched KEGG pathways (large nodes; size =  
925 proportional to connections, darker red color = greater significance) and associated genes  
926 (small nodes; green = male enriched, yellow = female enriched). (C) Top significantly  
927 enriched Reactome pathways from the top 1000 significantly differentially expressed genes.  
928 ( $p\text{-adj} < 0.01$ ) for males and females. Gene ratio is calculated as  $k/n$ , where  $k$  is the number of  
929 genes identified in each Reactome pathway, and  $n$  is the number of genes from the submitted  
930 gene list participating in any Reactome pathway. For complete Reactome pathway lists see  
931 Supplementary file 1. (D) All transcripts of differentially expressed genes under the gene  
932 ontology term "Cellular response to ER stress" (GO:0034976) and genes labeled by their role

933 in transcription, translation, protein processing, protein folding, secretion and protein quality  
934 control. (E) All transcripts of differentially expressed ribosomal genes.

935

936 **Figure 3. Sex differences in mouse islet ER stress-associated phenotypes.** (A) Protein  
937 synthesis was quantified in dispersed islet cells from 20-week-old male and female B6 mice  
938 after treatment with 1  $\mu$ M Tg for 2- or 24-hours. In female islet cells, protein synthesis was  
939 significantly lower after a 2-hour Tg treatment compared to control ( $p=0.0152$ ; paired  
940 Student's  $t$ -test) and significantly higher after a 24-hour Tg treatment compared to a 2-hour Tg  
941 treatment ( $p=0.0027$ ; paired Student's  $t$ -test). In male islet cells, protein synthesis was  
942 significantly lower after a 2- and 24-hour Tg treatment compared to control ( $p=0.0289$  [0-2  
943 hour] and  $p=0.0485$  [0-24 hour]; paired Student's  $t$ -test). (B) In both male and female islet  
944 cells protein synthesis was repressed after 2 hours. By 24 hours, protein synthesis repression  
945 was resolved in female, but not male islet cells. (C-H) Quantification of propidium iodide (PI)  
946 cell death assay of dispersed islets from 20-week-old male and female B6 mice treated with  
947 thapsigargin (0.1  $\mu$ M, 1  $\mu$ M or 10  $\mu$ M Tg) or DMSO for 84 hours.  $n=4-5$  mice,  $>1000$  cells per  
948 group. Percentage (%) of PI positive cells was quantified as the number of PI-  
949 positive/Hoechst 33342-positive cells in female (C) and male (D) islet cells. Relative cell death  
950 at 84 hr in Tg treatments compared with DMSO treatment in females (E, G) and males (F, H).  
951 In female islet cells, cell death was significantly higher in 10  $\mu$ M Tg compared to control  
952 ( $p<0.0001$ ; unpaired Student's  $t$ -test). In male islet cells, cell death was significantly higher in  
953 0.1, 1.0 and 10  $\mu$ M Tg compared to control ( $p=0.0230$  [0.1  $\mu$ M],  $p<0.0001$  [1  $\mu$ M] and  
954  $p<0.0001$  [10  $\mu$ M]; unpaired Student's  $t$ -test) (D). For E-H, at 84 hours the % of PI positive  
955 cells for each treatment was normalized to the DMSO control avg for each sex. \* indicates  
956  $p<0.05$ , \*\* indicates  $p<0.01$ , \*\*\*\* indicates  $p<0.0001$ ; ns indicates not significant; error bars  
957 indicate SEM.

958

959 **Figure 4. Sex differences in ex vivo and in vivo insulin secretion.** (A) Experimental  
960 workflow of static glucose-stimulated insulin secretion. (B, C) Relative high glucose (20 mM;  
961 high glucose, HG) in treatments compared with DMSO in female (B) and male (C) islets.  
962 Female islet HG secretion was significantly higher compared with control after 0- and 2-hour  
963 Tg pre-treatments ( $p=0.0083$  [0-hour] and  $p=0.0371$  [2-hour]; Mann Whitney test). Male islet  
964 HG secretion was significantly lower compared with control after a 4-hour Tg pre-treatment  
965 ( $p=0.0013$ ; Mann Whitney test). (D) Insulin content. Female islet insulin content was

966 significantly higher compared with control after a 4-hour Tg pre-treatment (p=0.0269; Mann  
967 Whitney test). (E) Proinsulin content. Female islet proinsulin content was significantly lower  
968 compared with control after a 2-hour Tg pre-treatment (p=0.0437; Mann Whitney test). Male  
969 islet proinsulin content was significantly lower compared with control after 2- and 4-hour Tg  
970 pre-treatments (p=0.0014 [2-hour] and p=0.0005 [4-hour]; Mann Whitney test). (F-H)  
971 Physiology measurements after a 6-hour fast in 20-week-old male and female B6 mice. (F, G)  
972 Insulin levels from glucose-stimulated insulin secretion tests (F: nM, G: % basal insulin)  
973 following a single glucose injection (2 g glucose/kg body weight, i.p). Area under the curve  
974 (AUC) calculations (n=13 females, n=18 males). (F) Insulin levels were significantly higher in  
975 male mice at 0 minutes and 30 minutes post injection (p=0.0063 [0 minutes] and p=0.0009  
976 [30 minutes]; Student's *t*-test). AUC was significantly higher in males (p=0.0159; Student's *t*-  
977 test). (G) Insulin levels (% baseline). Glucose-stimulated insulin secretion was significantly  
978 higher in female mice 15 minutes post injection (p=0.0279; Student's *t*-test). (H) Glucose  
979 levels from glucose tolerance tests following a single glucose injection (2 g glucose/kg body  
980 weight). AUC calculations (n=11 females, n=11 males). For B-E, grey triangles indicate the  
981 concentration of insulin or proinsulin from five islets, black circles indicate the average values  
982 per mouse. For B, ## indicates p<0.01 and ### indicates p<0.001 for comparisons between  
983 treatments and DMSO in low glucose. For all other figures, \* indicates p<0.05, \*\* indicates  
984 p<0.01, \*\*\* indicates p<0.001; ns indicates not significant; error bars indicate SEM.  
985

986 **Figure 5. Sex-specific transcriptomic and proteomic profiles following ER stress in**  
987 **mouse islets.** (A) Principal component analysis (PCA) of RNAseq data from male and female  
988 mouse islets treated with DMSO or 1  $\mu$ M Tg for 6- or 12-hours. (B) Spearman correlation  
989 depicting the variance for the first 5 principal components. (C) Top significantly enriched  
990 Reactome pathways from the top 1000 significantly differentially expressed genes ( $p$ -  
991 adj<0.01) for females and males that were upregulated or downregulated between 6-12 hours  
992 of Tg treatment. Gene ratio is calculated as k/n, where k is the number of genes identified in  
993 each Reactome pathway, and n is the number of genes from the submitted gene list  
994 participating in any Reactome pathway. (D) Protein abundance from proteomics data of  
995 female and male mouse islets treated with DMSO or 1  $\mu$ M Tg for 6 hours. Top 45 differentially  
996 expressed proteins are shown ( $p$ < 0.05).  
997

998 **Table 1 – Human  $\beta$  cell pathway gene numbers.** The number of genes corresponding to  
999 each T2D upregulated pathway in males, females or both sexes.

1000

1001 **SUPPLEMENTAL FIGURE LEGENDS**

1002 **Figure S1. Sex-specific and non-sex-specific differentially expressed genes in T2D.**

1003 scRNAseq data from male and female human  $\beta$  cells. (A-C) Top 60 significantly differentially  
1004 expressed genes ( $p$ -adj < 0.05). Non-sex-specific (A), female-specific (B), or male-specific  
1005 (C). For complete gene lists see Supplementary file 1.

1006

1007 **Figure S2. Gene expression changes in T2D.** scRNAseq data from male and female  
1008 human  $\beta$  cells. (A, B) Top 60 differentially expressed genes ( $p$ -adj < 0.05) in females (A) and  
1009 males (B). Sex-specific genes are indicated in red text. For complete gene lists see  
1010 Supplementary file 1.

1011

1012 **Figure S3. Correlations between donor attributes and insulin secretion.** Pearson  
1013 correlation of all human donors (A), or all donors with Type 2 Diabetes. Significant  
1014 correlations are denoted with a star (\*). For donor metadata see Supplementary file 6.

1015

1016 **Figure S4. Equivalent insulin sensitivity in male and female mice.** (A) Insulin tolerance  
1017 test (ITT). 20-week-old female and male B6 mice were fasted for 6 hours. Glucose levels (%  
1018 baseline) from insulin tolerance tests (ITT) following a single insulin injection (0.75U insulin/kg  
1019 body weight). AUC calculations (n=11 females, n=11 males). ns indicates not significant; error  
1020 bars indicate SEM.

1021

1022 **Figure S5. Mouse islet gene expression clusters by sex.** (A) Unsupervised hierarchical  
1023 clustering of RNAseq data from female and male mouse islets. Sorting was based on all  
1024 genes where the total count was >10 across all samples.

1025

1026 **Figure S6. ER stress-induced protein synthesis repression persists in male mouse islet**  
1027 **cells.** (A) Representative images of dispersed islets stained with nuclear mask and OPP  
1028 labeled with Alexa Fluor 594. (B, C) Integrated staining intensity of Alexa Fluor 594 in nuclear  
1029 mask positive islet cells in control media (B, FBS+) or after treatment with DMSO control or 1  
1030  $\mu$ M Tg for 2- or 24-hours (C, FBS-). Protein synthesis is displayed on a per cell basis from

1031 data shown in Figure 3. n=4-5 mice, >1000 cells per group. Mean values are indicated under  
1032 each group. (B) Protein synthesis was significantly higher in male islet cells than female islet  
1033 cells in control media, 3.9% (p=0.027; unpaired Student's *t*-test). (C) In female islet cells,  
1034 protein synthesis was significantly repressed from control-2 hour treatments (p=<0.0001;  
1035 unpaired Student's *t*-test) and significantly increased from both control-24 hour treatments  
1036 and 2-24 hour treatments (p=<0.0001; unpaired Student's *t*-test). In male islet cells, protein  
1037 synthesis was significantly repressed from control-2 hour treatments (p=<0.0001; unpaired  
1038 Student's *t*-test) and control-24 hour treatments (p=<0.0001; unpaired Student's *t*-test);  
1039 however, was not significantly different between 2-24 hour treatments (p=0.07; unpaired  
1040 Student's *t*-test). \* indicates p<0.05, \*\*\* indicates p<0.001; ns indicates not significant; error  
1041 bars indicate SEM.

1042

1043 **Figure S7. *Ins2* gene activity is repressed by ER stress induction.** *Ins2* gene activity in  $\beta$   
1044 cells from 20-week-old male and female B6 mice treated with Tg (0.1  $\mu$ M or 1  $\mu$ M Tg) or  
1045 DMSO for 60 hours (n=6 mice per sex, > 1000 cells per group). (A, B) Average change in  
1046 fluorescence intensity from all GFP expressing female (A) and male (B)  $\beta$  cells over time.  
1047 Data was normalized to the first 2 hours to examine relative change in *Ins2* gene activity. (C,  
1048 D) Density plot of *Ins2*<sup>GFP/WT</sup>  $\beta$  cell GFP fluorescence intensity, log transformed. Data is  
1049 shown for each run for females (C) and males (D). (E-H) Average change in high (E, F) and  
1050 low (G,H) GFP *Ins2*<sup>GFP/WT</sup>  $\beta$  cells fluorescence over time from females (E, G) and males (F,  
1051 H). Data was normalized to the first two hours to examine relative change in *Ins2* gene  
1052 activity.

1053

1054 **Figure S8. Representative western blot images of UPR protein markers.** (A-D) Levels of  
1055 ER stress proteins were quantified in isolated islets from 20-week-old male and female B6  
1056 mice cultured in DMSO or 1  $\mu$ M Tg for 24 hours. (A) BiP levels were significantly upregulated  
1057 in female Tg vs DMSO (p=0.0011; paired Student's *t*-test) but not male Tg vs DMSO  
1058 (p=0.1187; paired Student's *t*-test). (B) pIRE1 $\alpha$  levels were significantly upregulated in female  
1059 Tg vs DMSO (p=0.0001; paired Student's *t*-test) and in male Tg vs DMSO (p=0.0148; paired  
1060 Student's *t*-test). (C) CHOP levels were significantly upregulated in female Tg vs DMSO  
1061 (p=0.0333; paired Student's *t*-test) and in male Tg vs DMSO (p=0.0164; paired Student's *t*-  
1062 test). (D) p-eIF2 $\alpha$  levels were not significantly upregulated in either sex. (E-G) Levels of ER  
1063 stress proteins were quantified in isolated islets from 60-week-old male and female B6 mice

1064 cultured in DMSO or 1  $\mu$ M Tg for 24 hours. (E) BiP levels were significantly upregulated in  
1065 male Tg vs DMSO ( $p=0.0048$ ; paired Student's  $t$ -test) but not in female Tg vs DMSO  
1066 ( $p=0.3319$ ; paired Student's  $t$ -test). (F) p-IRE1 $\alpha$  levels were not significantly upregulated in  
1067 either sex ( $p=0.9257$  [female] and  $p=0.8273$  [male]; paired Student's  $t$ -test). (G) p-eIF2 $\alpha$   
1068 levels were not significantly upregulated in either sex ( $p=0.8451$  [female] and  $p=0.3076$   
1069 [male]; paired Student's  $t$ -test). (H) Representative western blot images of 20-week Tg treated  
1070 mouse islets. (I) Representative western blot images of 60-week Tg treated mouse islets. \*  
1071 indicates  $p<0.05$ , \*\* indicates  $p<0.01$ ; ns indicates not significant.

1072

1073 **Figure S9. Female mouse islets retain greater insulin secretion during ER stress.** (A)  
1074 Insulin secretion at basal (3 mM; low glucose, LG) and stimulatory (20 mM; high glucose, HG)  
1075 glucose. Female islet LG secretion was significantly higher compared with control after 2- and  
1076 4-hour Tg pre-treatments ( $p=0.0047$  [2-hour] and  $p=0.0003$  [4-hour]; Mann Whitney test).  
1077 Female islet HG secretion was significantly higher compared with control after 0- and 2-hour  
1078 Tg pre-treatments ( $p=0.0012$  [0-hour] and  $p=0.0061$  [2-hour]; Mann Whitney test). Male islet  
1079 LG secretion was significantly higher compared with control after a 0-hour Tg pre-treatment  
1080 ( $p=0.0371$ ; Mann Whitney test). Male islet HG secretion was significantly lower compared with  
1081 control after a 4-hour Tg pre-treatment ( $p=0.0012$ ; Mann Whitney test). (B) Proinsulin  
1082 secretion at basal (3 mM) and stimulatory (20 mM) glucose. Female islet HG secretion was  
1083 significantly higher compared with control after 0- and 2-hour Tg pre-treatments ( $p=0.0075$  [0-  
1084 hour] and  $p=0.0437$  [2-hour],; Mann Whitney test). Male islet HG secretion was significantly  
1085 lower compared with control after a 4-hour Tg pre-treatment ( $p=0.0025$ ; Mann Whitney test). \*  
1086 indicates  $p<0.05$ , \*\* indicates  $p<0.01$ ; ns indicates not significant; error bars indicate SEM.

1087

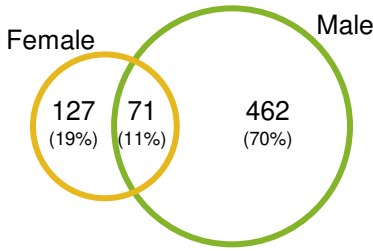
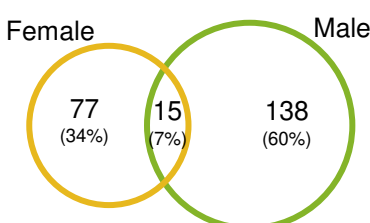
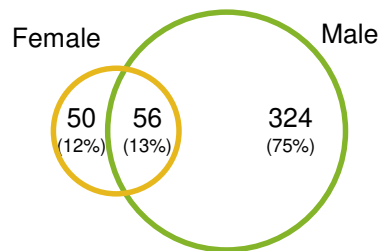
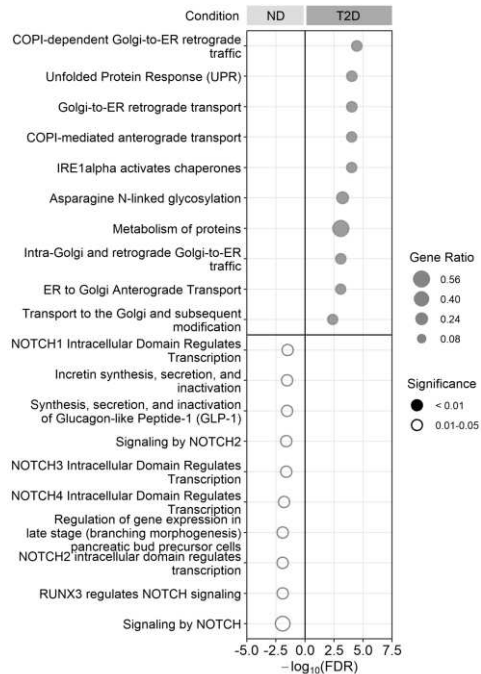
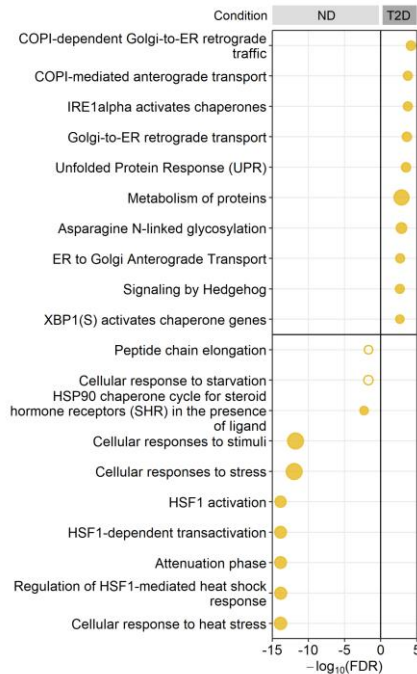
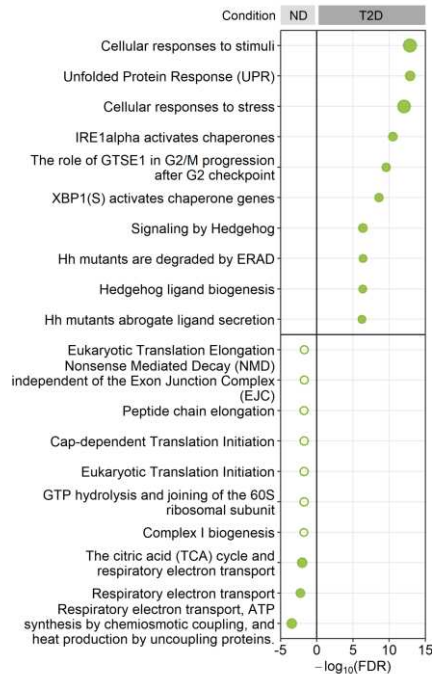
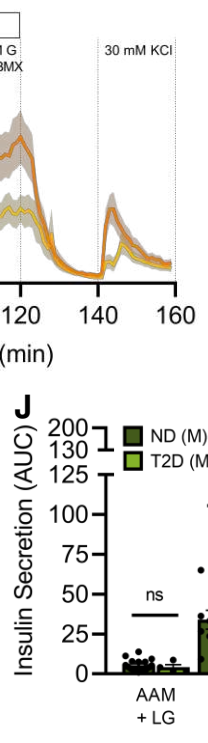
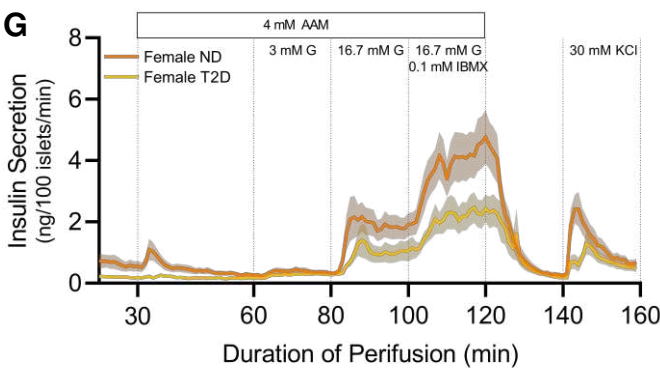
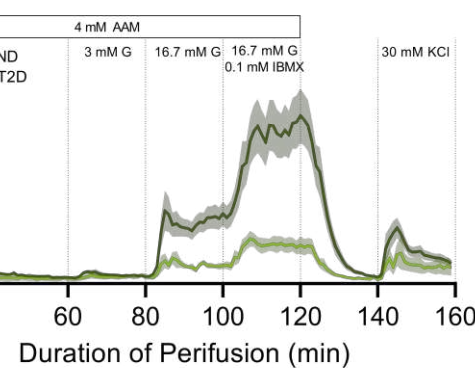
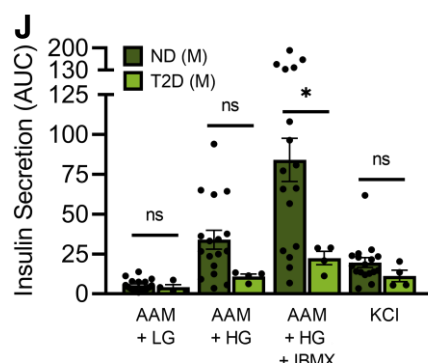
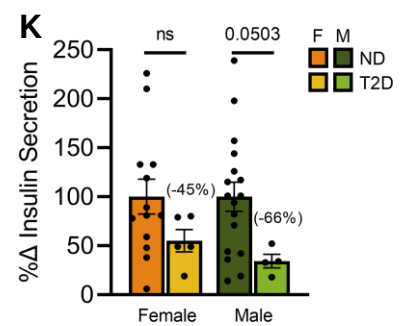
1088 **Figure S10. Mouse islet gene expression clusters by sex, treatment and time.** (A)  
1089 Unsupervised hierarchical clustering of RNAseq data from female and male DMSO or Tg  
1090 treated mouse islets. Sorting was based on all genes where the total count  $>10$  across all  
1091 samples.

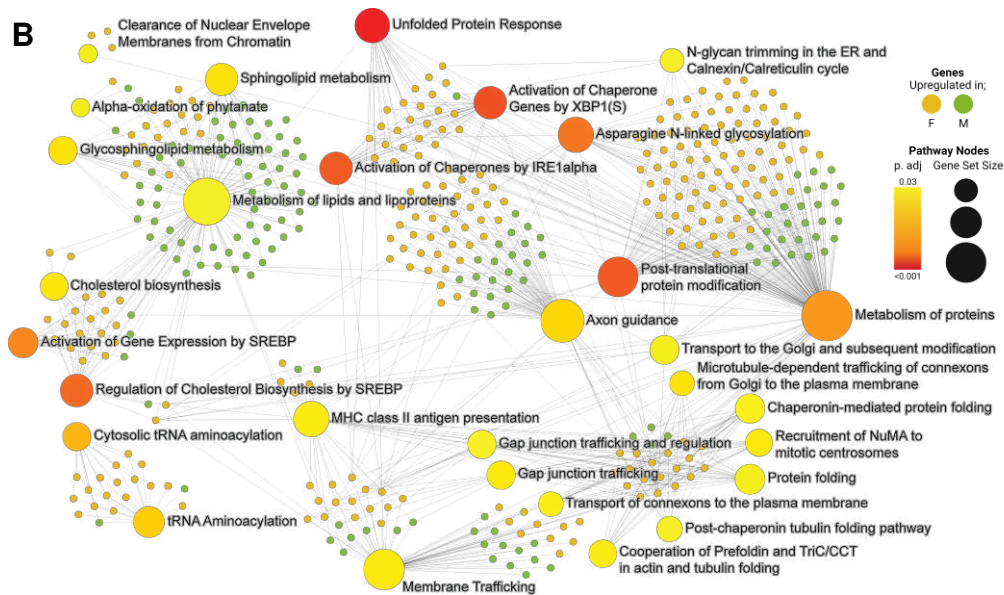
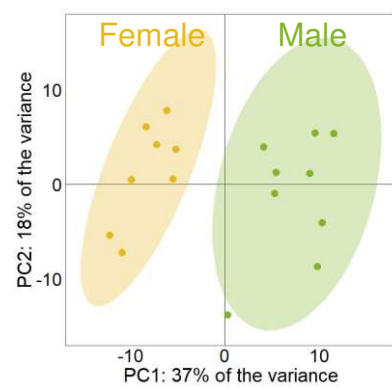
1092

1093 **Figure S11. Female and Male mouse islets are enriched in similar pathways following**  
1094 **6- and 12-hour Tg treatments.** (A, B) Most significantly enriched Reactome pathways from  
1095 the top 1000 significantly differentially expressed genes. ( $p$ -adj  $< 0.01$ ) for females and males  
1096 between DMSO vs Tg after 6 hours (A) or 12 hours (B) of Tg treatment. Gene ratio is

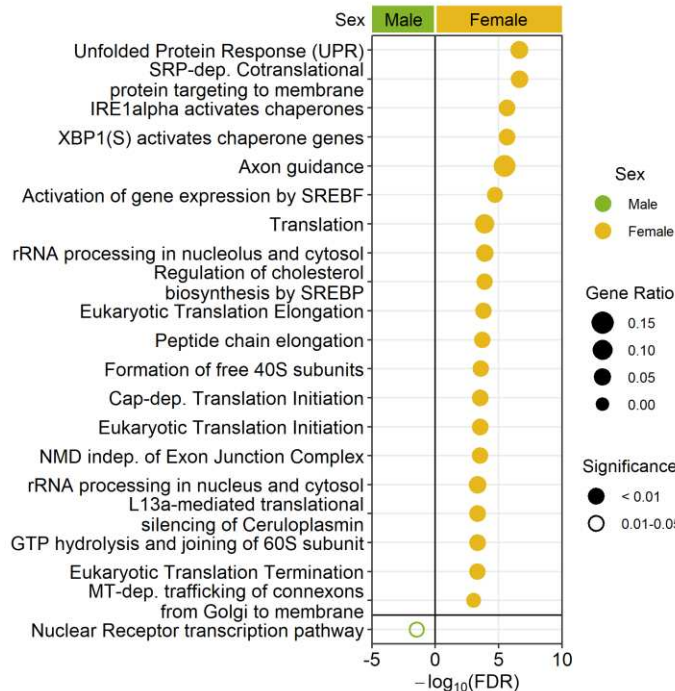
1097 calculated as k/n, where k is the number of genes identified in each Reactome pathway, and  
1098 n is the number of genes from the submitted gene list participating in any Reactome pathway.  
1099

1100 **Figure S12. A greater number of  $\beta$  cell identity genes are downregulated between 6-  
1101 and 12- hour Tg treatment times in female mouse islets.** (A, B) Treatment:Time interaction  
1102 plots of female islet (A) and male islet (B)  $\beta$  cell identity genes in Reactome pathway  
1103 “Regulation of gene expression in  $\beta$  cells”. The fold change (FC) for DMSO vs Tg was  
1104 calculated for each sex and time point (Female 6-hour, Female 12-hour, Male 6-hour, Male  
1105 12-hour). The change in FC values (12-hour FC – 6-hour FC) were plotted according to  $p$ -adj  
1106 values. In females, FC values between 6- and 12-hours are represented by orange and  
1107 purple dots, respectively. In males, FC values at 6- and 12-hours are represented by green  
1108 and blue dots, respectively. A solid black line connecting the dots indicates genes with a  
1109 significant treatment:time interaction.

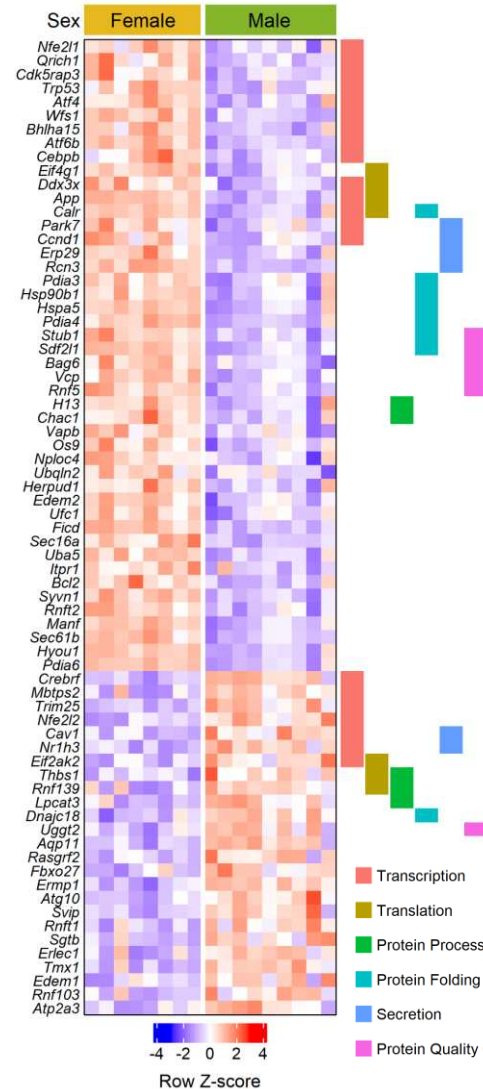
**A Differentially Expressed****B T2D Downregulated****C T2D Upregulated****D Non-Sex-Specific  $\beta$  Cell Enriched****E Female  $\beta$  Cell Enriched****F Male  $\beta$  Cell Enriched****G****H****J****K****Figure 1**



## C Enriched Pathways



## D ER Stress Genes



## Ribosome Genes

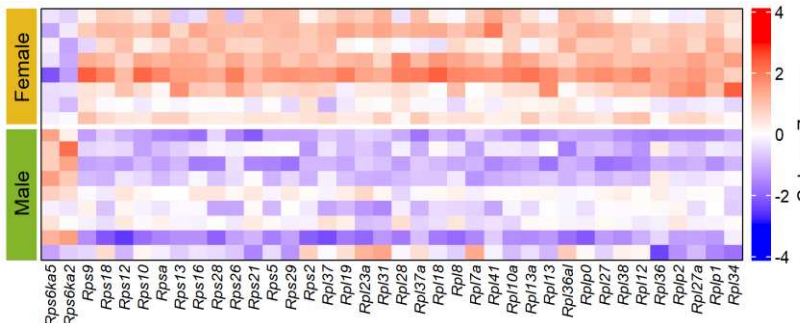


Figure 2

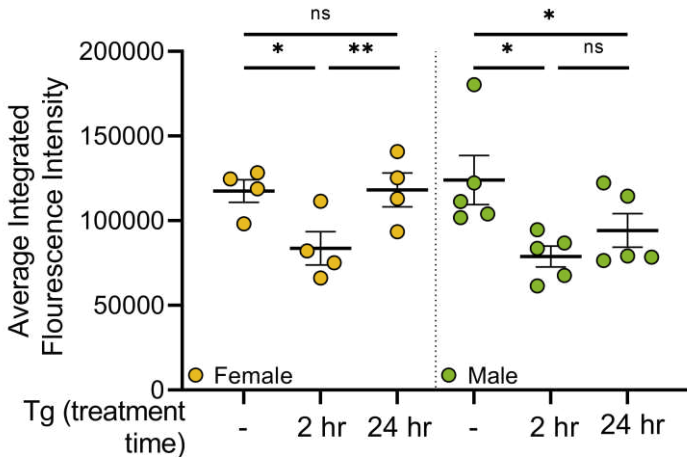
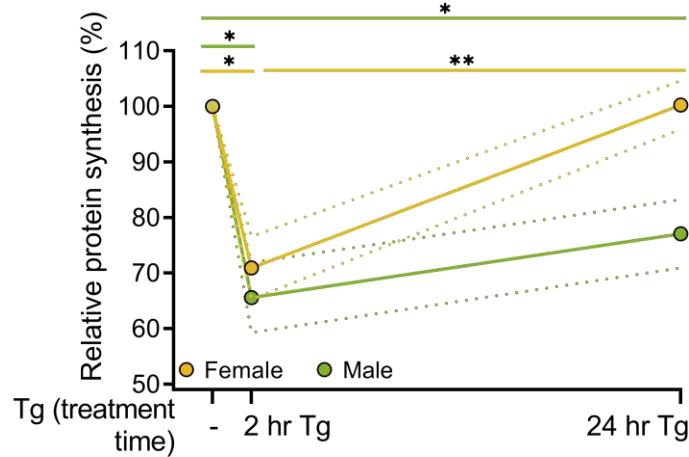
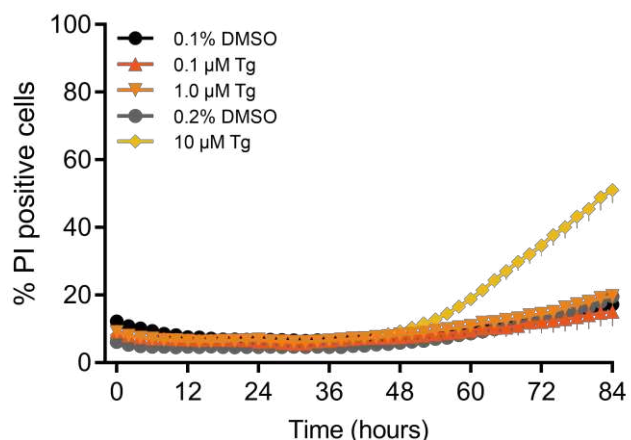
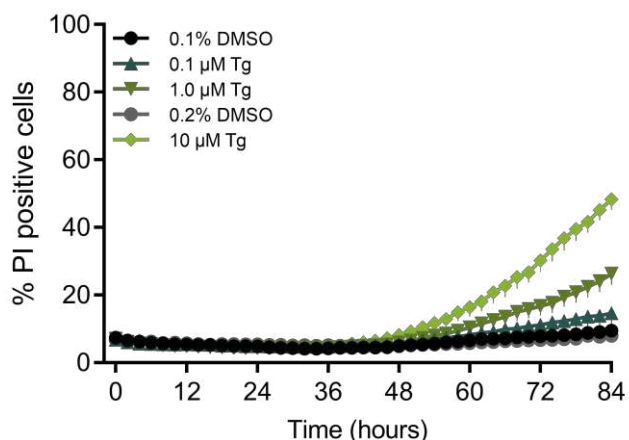
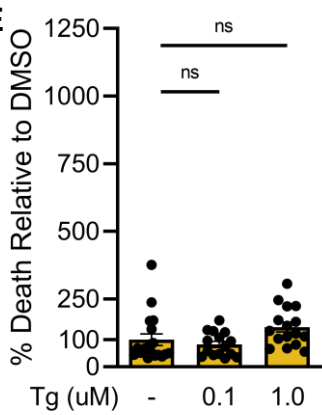
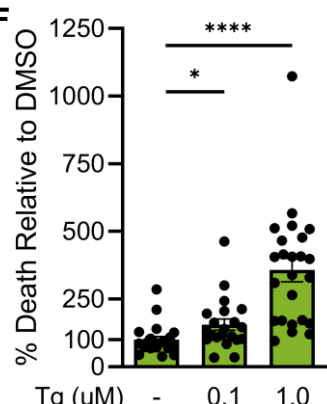
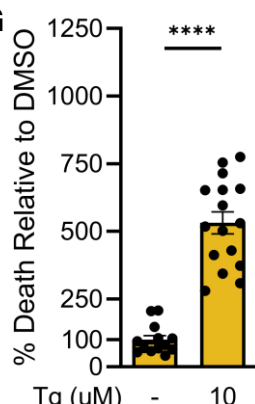
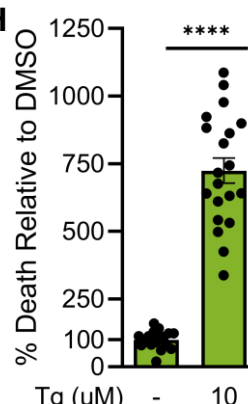
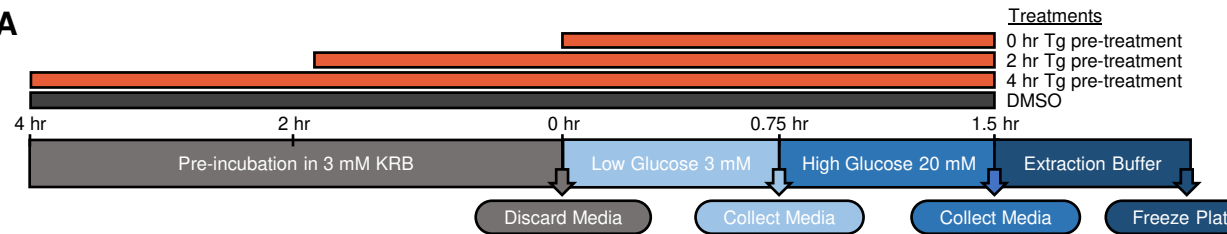
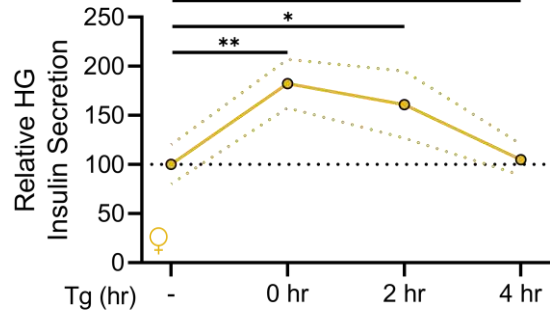
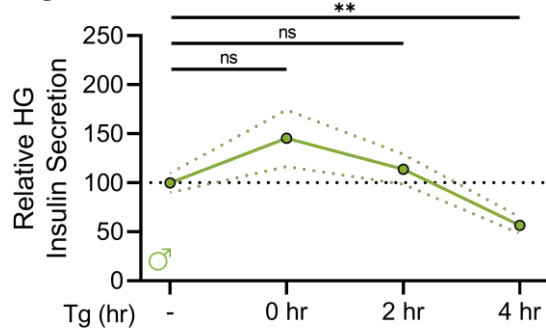
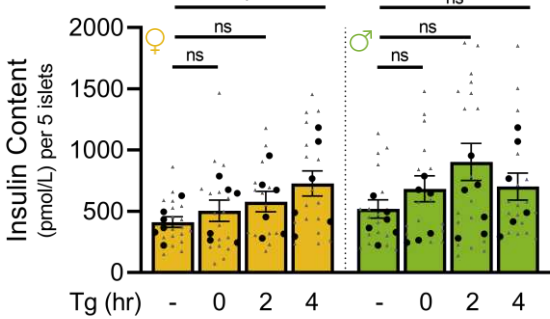
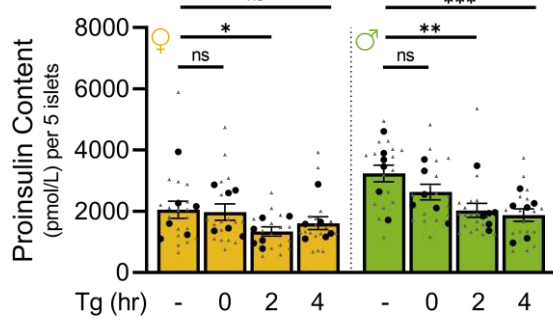
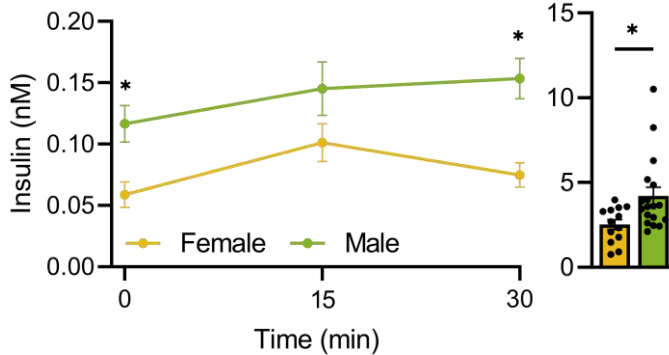
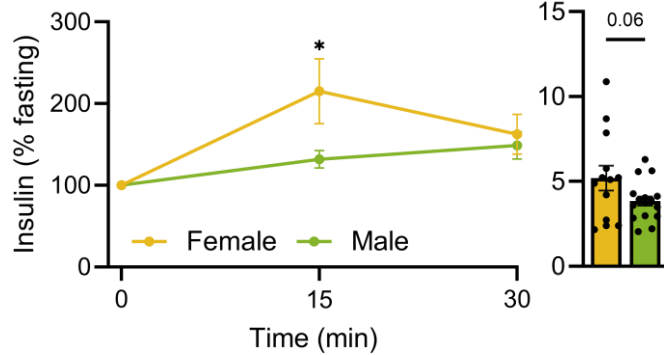
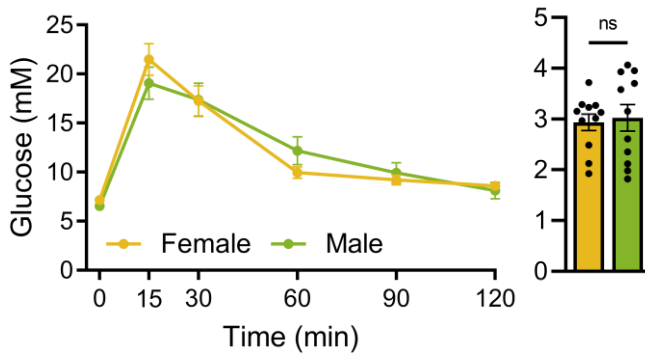
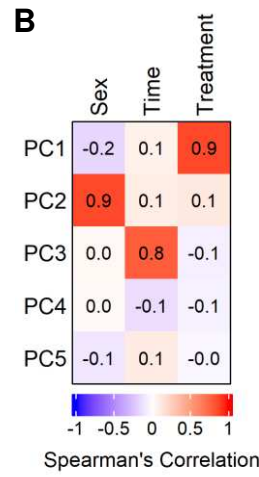
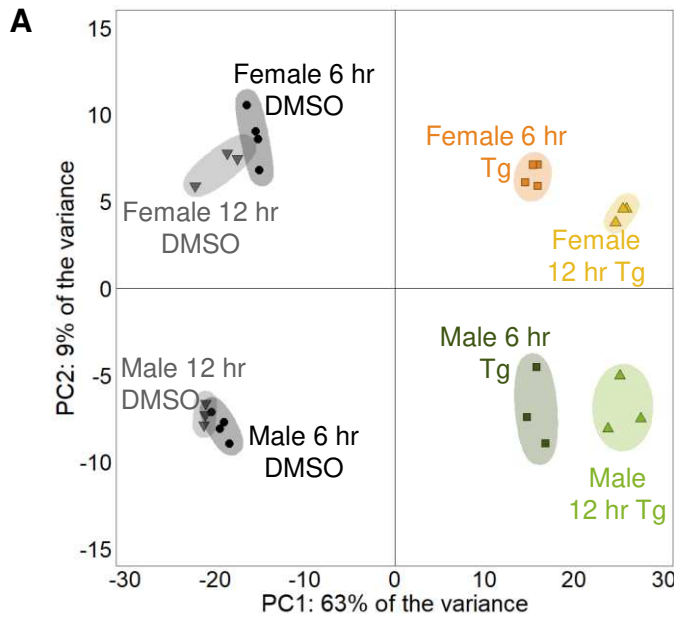
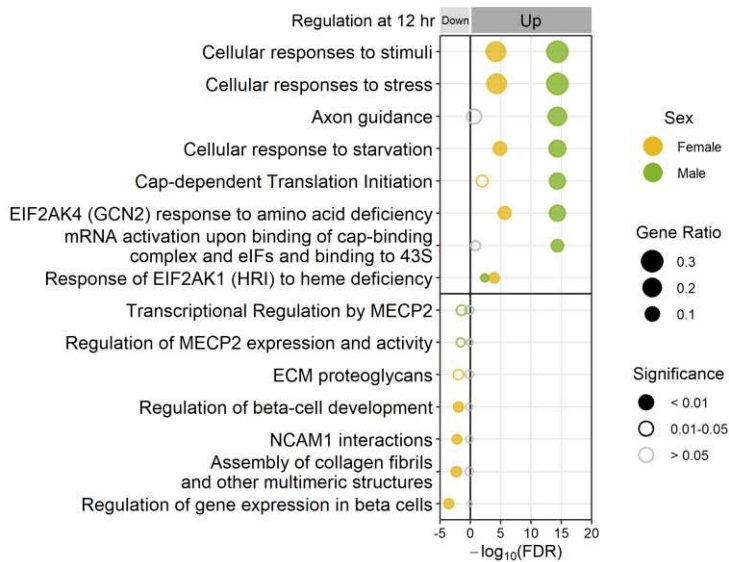
**A** *Islet Protein Synthesis***B** *Relative Islet Protein Synthesis***C** *Time Course Cell Death - Female***D** *Time Course Cell Death - Male***E****F****G****H**

Figure 3

**A****B****C****D****E****F****G****H****Figure 4**



**C Islet Pathway Enrichment [up/downregulated genes 6-12 hr]**



**D Differentially Expressed Islet Proteins**

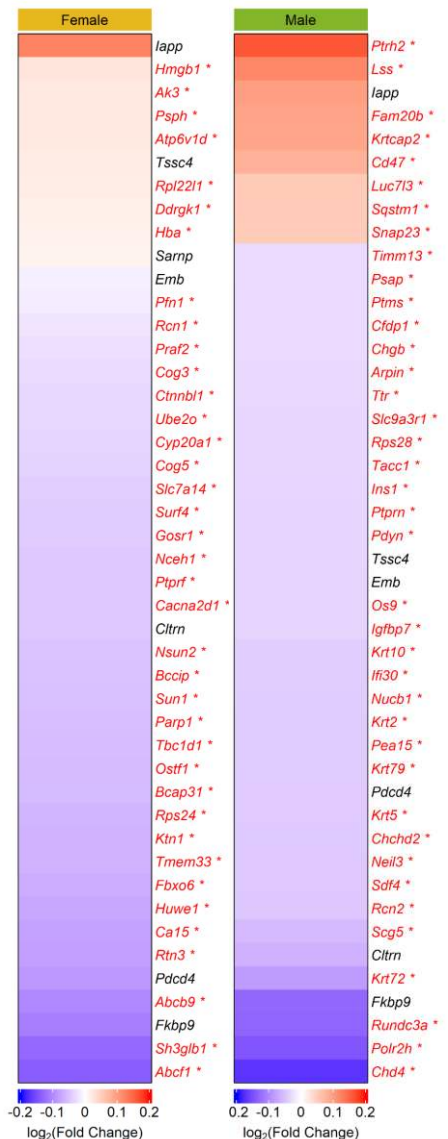
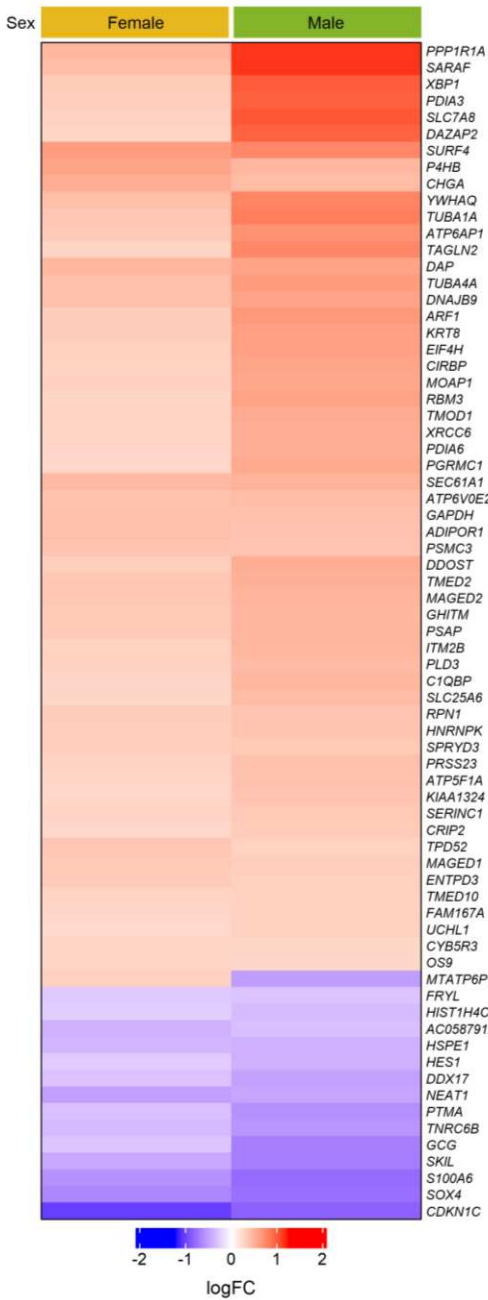
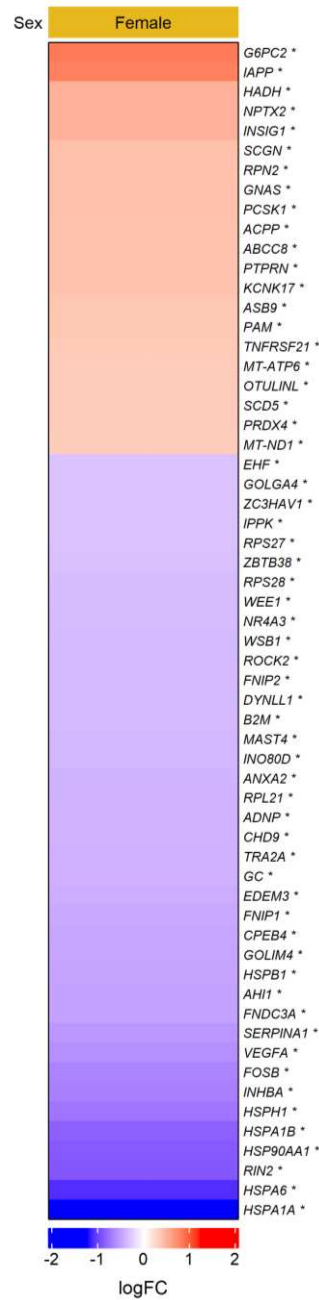
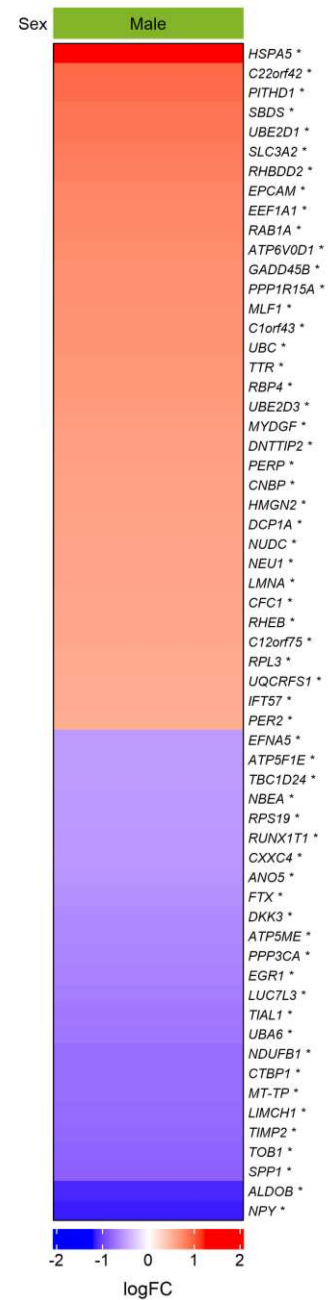
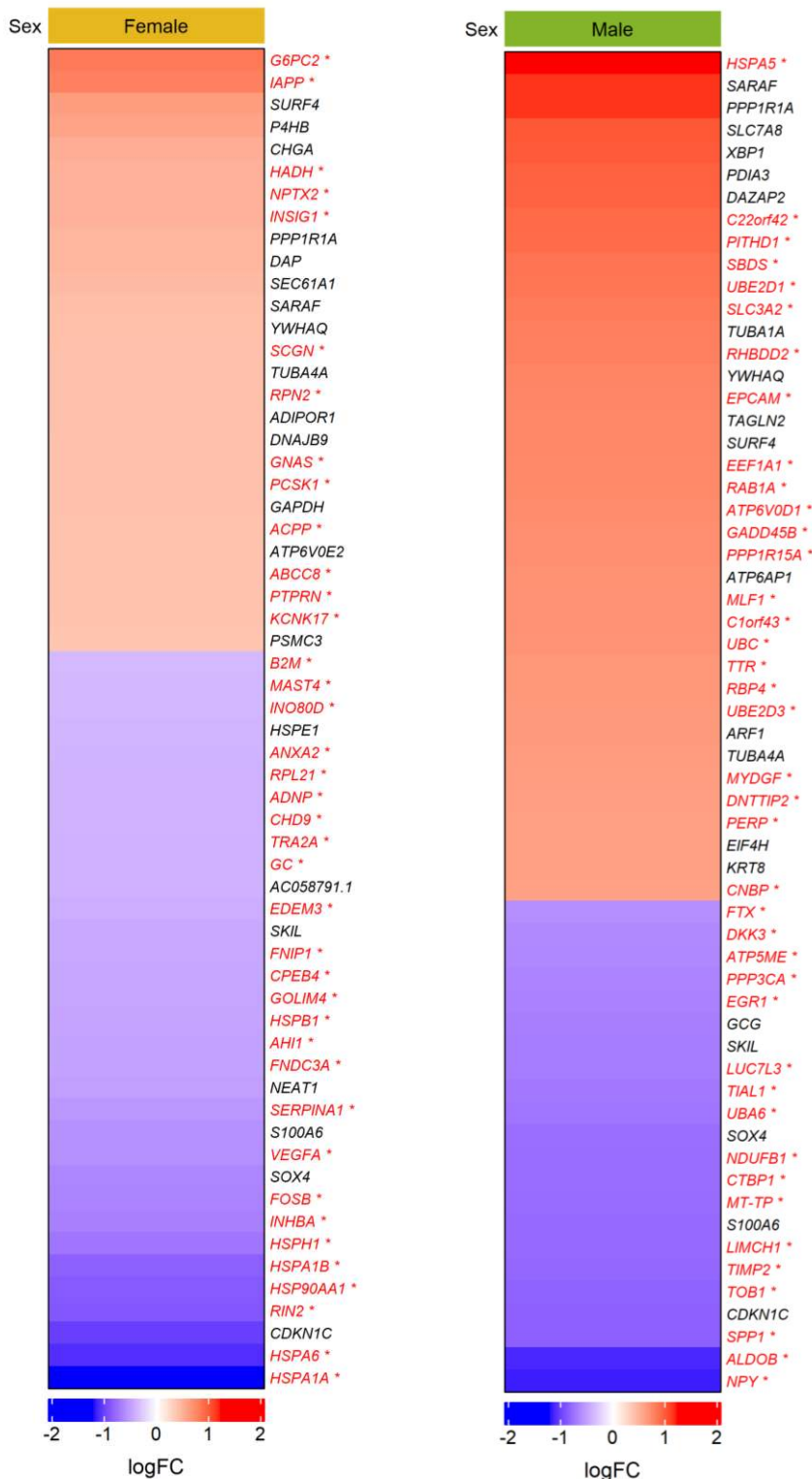


Figure 5

Pathway Name	Number of Pathway Genes		
	Unique Male	Common	Unique Female
Asparagine N-linked glycosylation	16	9	4
Cellular responses to stimuli	49	9	6
Cellular responses to stress	47	9	6
COPI-dependent Golgi-to-ER retrograde traffic	7	6	2
COPI-mediated anterograde transport	7	5	2
ER to Golgi Anterograde Transport	10	5	2
Golgi-to-ER retrograde transport	7	6	2
Hedgehog ligand biogenesis	12	4	1
Hh mutants abrogate ligand secretion	12	3	1
Hh mutants are degraded by ERAD	12	3	1
IRE1alpha activates chaperones	9	3	1
Metabolism of proteins	69	20	13
Signaling by Hedgehog	16	6	2
The role of GTSE1 in G2/M progression after G2 checkpoint	14	4	1
Unfolded Protein Response (UPR)	13	3	1
XBP1(S) activates chaperone genes	8	3	1

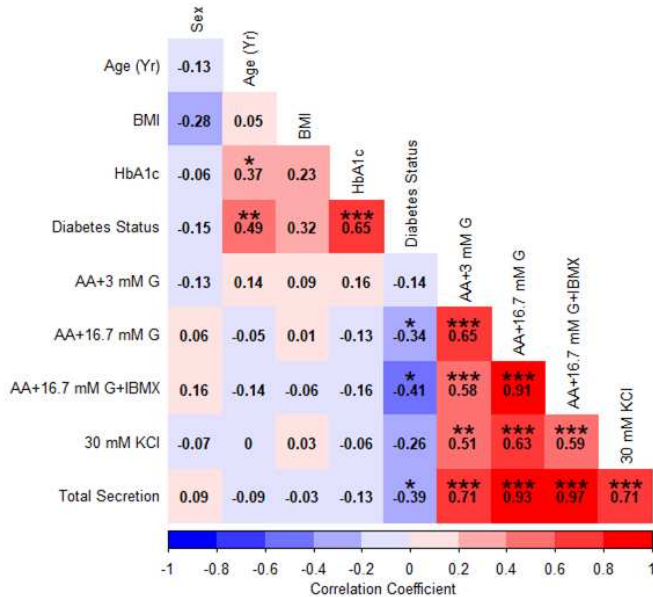
**A** Non-Sex-Specific Differentially Expressed Genes in  $\beta$  Cells**B** Female-Specific Differentially Expressed Genes in  $\beta$  Cells**C** Male-Specific Differentially Expressed Genes in  $\beta$  Cells

**A** Top 60 Differentially Expressed Genes in Female  $\beta$  Cells**B** Top 60 Differentially Expressed Genes in Male  $\beta$  Cells

\*Indicates genes unique to females

\*Indicates genes unique to males

### A All Donors



### B Donors with Type 2 Diabetes

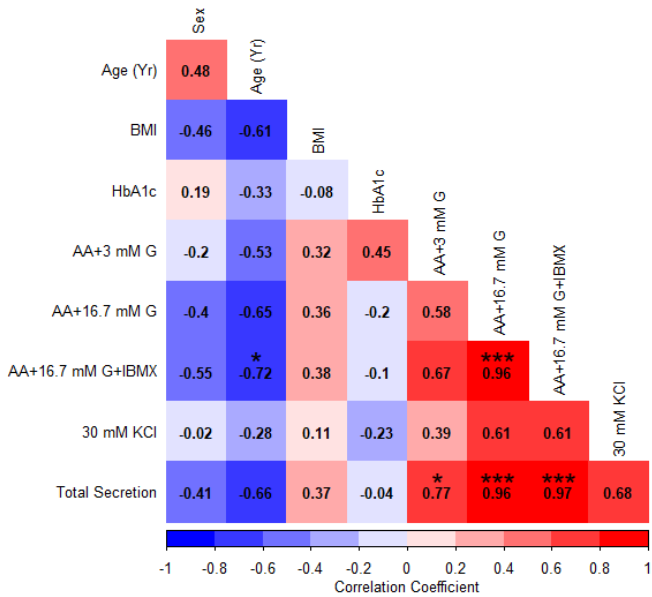


Figure S3

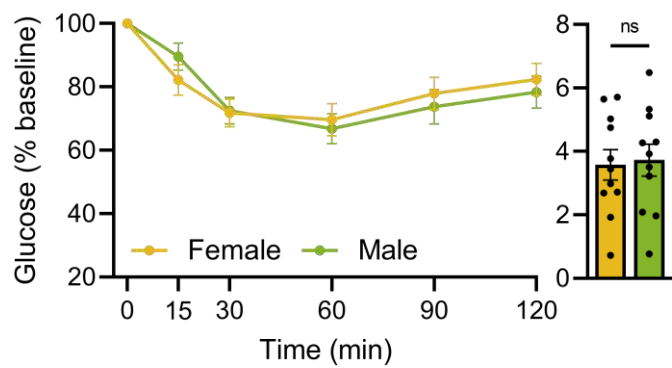
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Figure S4

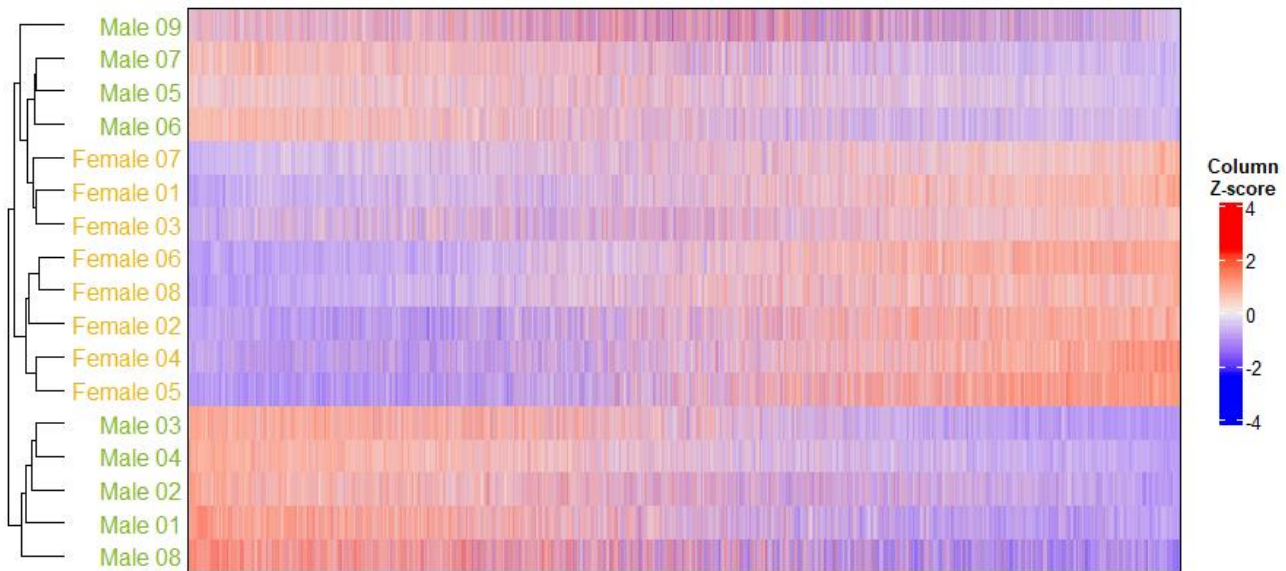
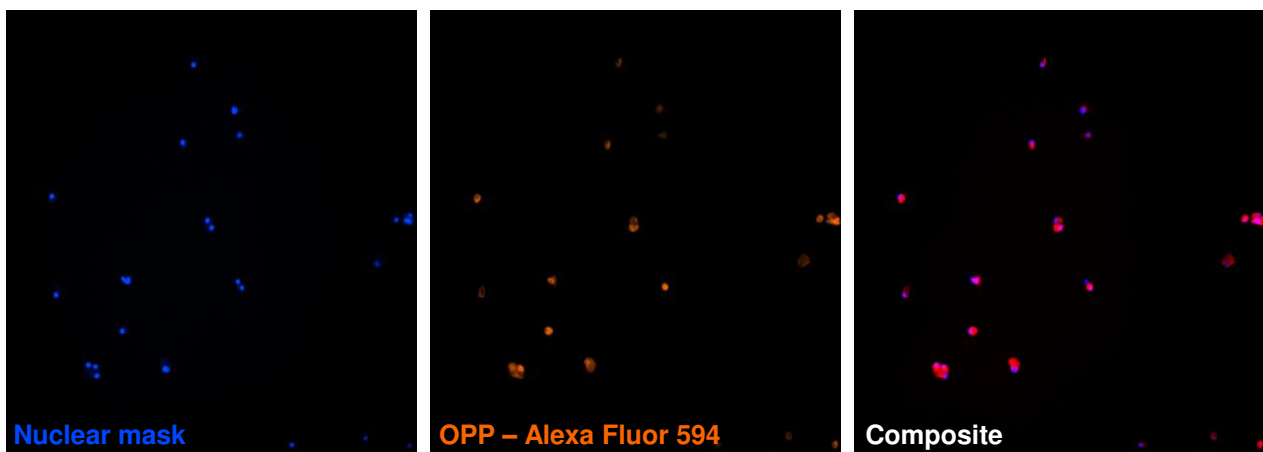
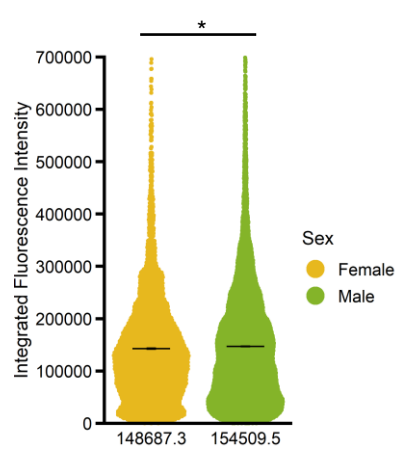
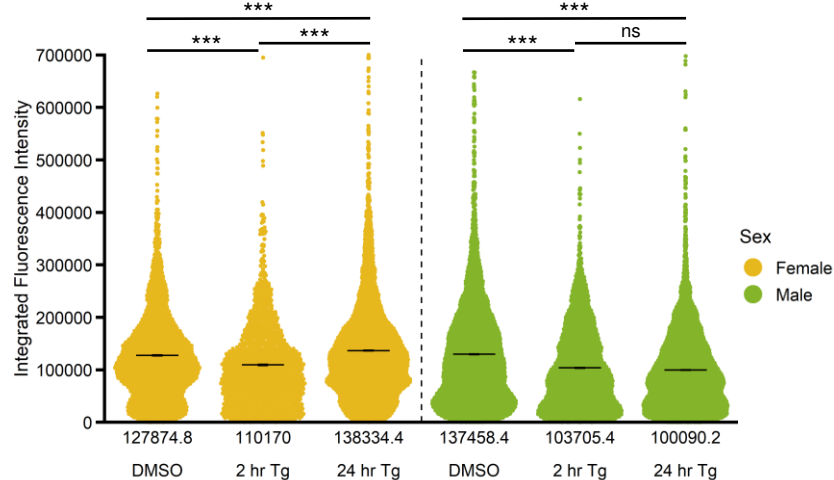
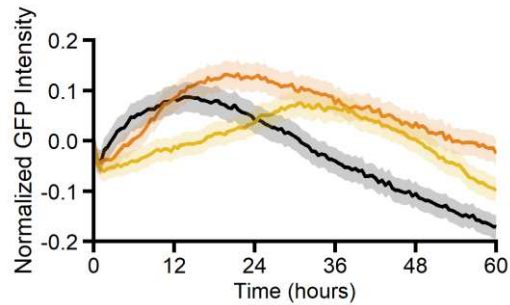
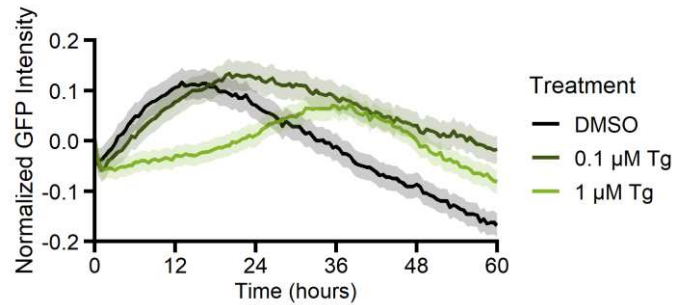
**A***Unsupervised Hierarchical Clustering of Mouse Islet Genes*

Figure S5

**A** Representative Protein Synthesis Staining Images**B** Protein Synthesis per Cell**C** Protein Synthesis per Cell

**A***Ins2 Gene Activity - Female***B***Ins2 Gene Activity - Male*

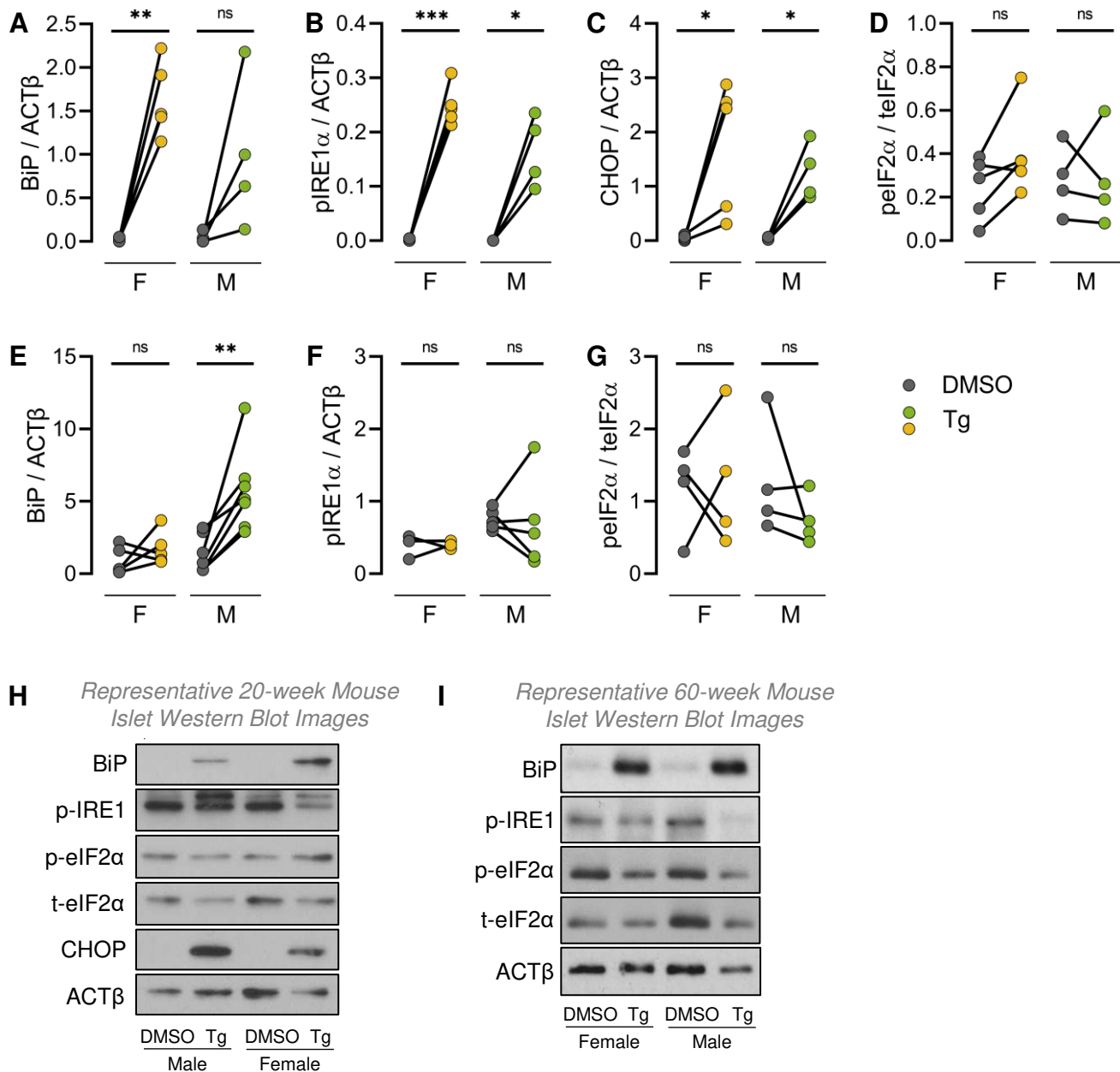
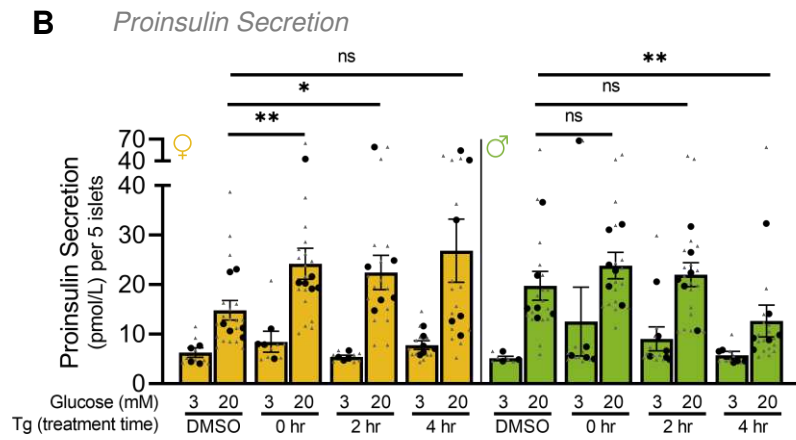
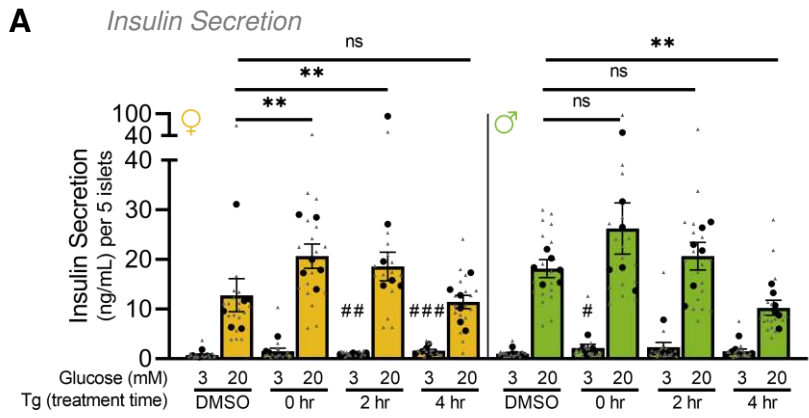
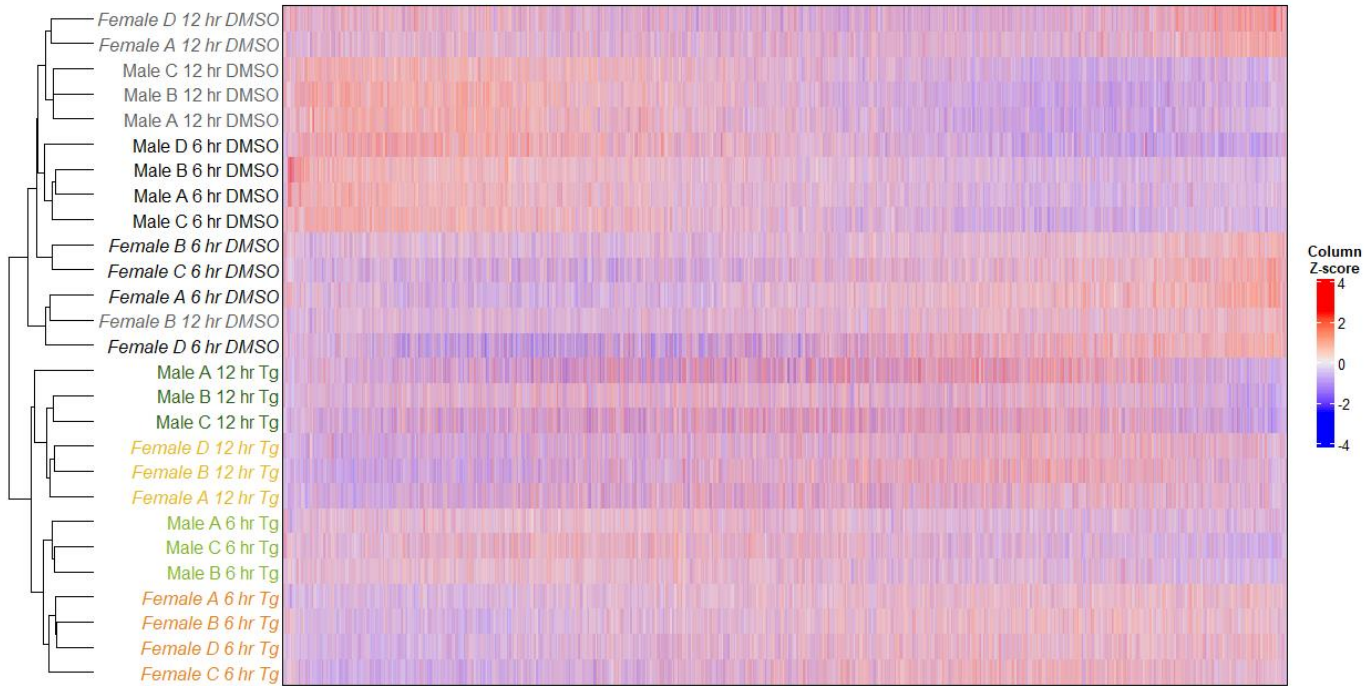


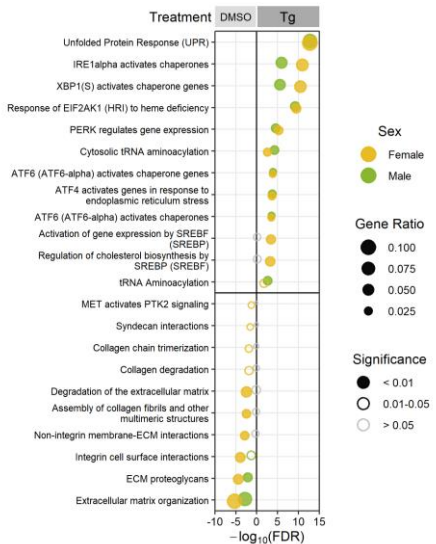
Figure S8



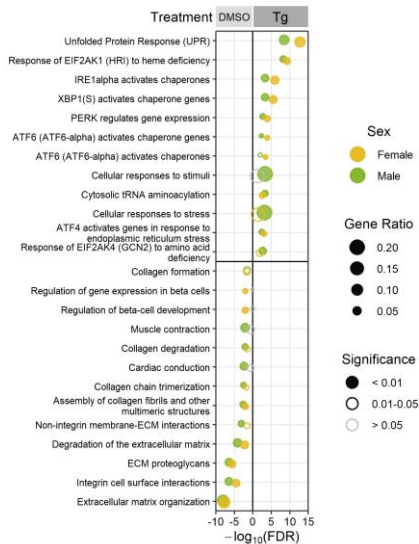
**A***Unsupervised Hierarchical Clustering of Mouse Islet Genes*

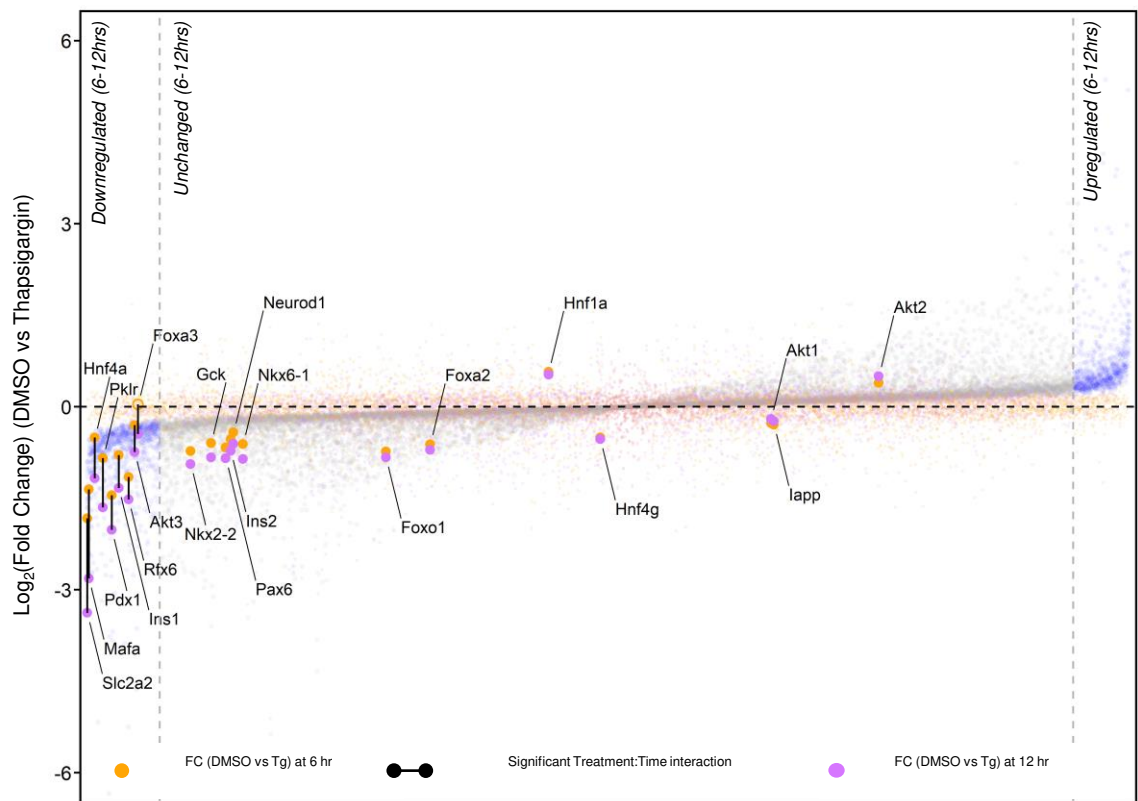
**A**

*Islet Pathway Enrichment*  
[6 hr, DMSO vs Tg]

**B**

*Islet Pathway Enrichment*  
[12 hr, DMSO vs Tg]



**A**  $\beta$  cell identity genes - Female**B**  $\beta$  cell identity genes - Male