Sex differences underlying pancreatic islet biology and its dysfunction

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ABSTRACT

Background: The sex of an individual affects glucose homeostasis and the pathophysiology, incidence, and prevalence of diabetes as well as the response to therapy.

Scope of the review: This review focuses on clinical and experimental sex differences in islet cell biology and dysfunction during development and in adulthood in human and animal models. We discuss sex differences in β-cell and α-cell function, heterogeneity, and dysfunction. We cover sex differences in communication between gonads and islets and islet-cell immune interactions. Finally, we discuss sex differences in β-cell programming by nutrition and other environmental factors during pregnancy.

Major conclusions: Important sex differences exist in islet cell function and susceptibility to failure. These differences represent sex-related biological factors that can be harnessed for gender-based prevention of and therapy for diabetes.

Keywords Sex differences; Gender differences; Islet; β-cell; α-cell; Diabetes; Immune cells

1. INTRODUCTION

Sex differences in physiology begin early in development from the combination of genetic and hormonal cues and they continue after puberty [1]. These differences result from the combination of three major events: 1. The differences in the number and type of sex chromosomes; 2. The perinatal testosterone surges that masculinize the reproductive tract and the organization of neural circuits; and 3. The activity of gonadal hormones after puberty. The combination of these factors produces distinct male and female biological systems for islet cells in vivo.

Increasing evidence suggests that sex affects glucose homeostasis and the pathophysiology, incidence, and prevalence of diabetes, as well as response to therapy. Sex differences in glucose homeostasis and diabetes have been recently reviewed elsewhere [2–4]. Normoglycemic women have lower fasting plasma glucose concentrations than men and higher 2-h plasma glucose concentrations following an oral glucose tolerance test (OGTT) [5], slower gastric emptying [6], and higher insulin sensitivity than men [7]. In addition, in individuals with prediabetes, women tend to exhibit glucose intolerance, whereas men exhibit impaired fasting glucose [8]. This review will focus on clinical and experimental sex differences in islet biology and dysfunction during development and in adulthood. The study of sex differences in islet function and failure is of fundamental importance, because it will generate knowledge on sex-related biological factors that can be harnessed for better options for prevention of and therapy for diabetes.

2. SEX DIFFERENCES IN β-CELL FUNCTION UNDER NORMAL AND STRESS CONDITIONS

2.1. Clinical studies

Sex differences in β-cell function are apparent in clinical studies. Compared to healthy men and despite similar plasma glucose levels, healthy women exhibit enhanced postprandial plasma insulin and C-peptide concentrations after a meal [9], suggesting that women have increased insulin secretion for a given glucose load. In addition, the disposition index, which reflects insulin secretion for a given level of insulin action, is higher in women than in men, supporting greater insulin secretion in women [9]. In a recent study of 63 healthy Japanese men and women, the insulin responses to an oral glucose load were higher in females than males, with no differences in insulin sensitivity between the sexes [10]. Likewise, studies in older men and women suggest that for a given level of insulin action, women have higher levels of insulin secretion [11]. It has been hypothesized that the increased glucose-stimulated insulin secretion (GSIS) in females was due to sex differences in glucose-stimulated GLP-1 production. In the ADDITION-PRO study, a large study population of 1,462 Danish adults, normoglycemic women had a greater increase in serum GLP-1 concentrations following an OGTT, than normoglycemic men, even after adjusting for body weight, height or BMI [12]. This sex difference could be explained by the fact that the female hormone 17β-estradiol (E2) stimulates glucose-induced GLP-1 secretion as will be discussed below. However, the relationship between the increased serum GLP-1

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following OGTT and the increased first phase insulin secretion was similar in men and women [12]. Therefore, sex differences in serum GLP-1 concentrations following OGTT do not seem to explain sex differences in β-cell function in humans. In the same cohort, individuals with prediabetes and type 2 diabetes (T2D) exhibited less of a GLP-1 increase than normoglycemic controls following an OGTT. This impaired serum GLP-1 increase in response to OGTT was most pronounced in women compared with men [12]. Thus, healthy women exhibit a greater increase in GLP-1 following an oral glucose challenge, but, as glucose tolerance deteriorates, this sex difference is no longer apparent. Interestingly, in individuals with prediabetes, women tend to be predisposed to impaired glucose tolerance, compared to men [8]. However, it is unknown if these phenotypic differences reflect differences in β-cell function.

Gender differences in β-cell failure are also observed in insulin-deficient forms of diabetes. For example, type 1 diabetes (T1D) is the only common autoimmune disease characterized by a male predominance [2–4,13]. Girls presenting with T1D in puberty have higher residual β-cell function than boys at diagnosis [14]. Similarly, ketosis-prone diabetes (KPD) is a phenotypically-defined form of T2D characterized by a strong male predominance (75%) and rapid β-cell failure leading to severe insulin deficiency [15]. In subjects with KPD, the male gender is associated with a more pronounced decrease in β-cell insulin secretory reserve, assessed by fasting and glucagon-stimulated C-peptide [16]. The rare women developing KPD were in an anovulatory state. Finally, a mutation in the β-cell transcription factor V-Maf avian musculoaponeurotic fibrosarcoma oncogene homolog A (MAFA), causes autosomal dominant inheritance of diabetes or insulinomatosis (insulin-secreting tumors of the pancreas causing adult-onset hyperinsulinemic hypoglycemia) with a sex dimorphism. Women with the MAFA mutation are more likely to develop insulinomatosis while men are more likely to develop diabetes [17]. The reason for this divergent phenotype has not yet been determined, but, interestingly, the women who developed insulinomatosis had a prior pregnancy, suggesting that the hormonal milieu of pregnancy influences the effect of this mutation on the β-cell.

2.2. Studies using human islets and β-cells
The Integrated Islet Distribution Program (IIDP; https://iidp.coh.org/) has greatly facilitated the use of cadaver donor human islets in research. Thus, we can now more easily assess whether intrinsic differences exist between islets and β cells obtained from male and female individuals. In one study, 53 male and 34 female human pancreatic islet donors were compared [18]. There were no significant differences in age, body mass index (BMI), hemoglobin A1c (HbA1c), islet purity or β-cell content between males and females. However, ex vivo GSIS was slightly higher in islets from females compared with males.

Information generated from studies of pancreas development and endocrine differentiation from multiple organisms has resulted in multistep directed-differentiation protocols that can convert human embryonic stem cells (hESCs) into pancreatic progenitor cells, which, several months following implantation into immunocompromised mice, develop into mature glucose-responsive insulin-secreting cells that are capable of reversing experimentally-induced diabetes [19,20]. To date, clinical trials assessing the utility of encapsulated hESC-derived pancreatic progenitors in patients with poorly controlled T1D have not been successful in regulating glycemia, likely due to fibrosis in the transplanted cells. Interestingly, recent studies suggest that the sex of the host into which hESC-derived pancreatic progenitors are transplanted may affect their ability to mature into optimally functional β-cells. Male and female mice were transplanted with two different stages of hESC-derived pancreatic cells: endocrine progenitors or insulin-positive cells. In vivo maturation of both cell populations into glucose-responsive insulin-secreting cells (as measured by circulating human C-peptide) was accelerated in female recipients (12 weeks compared to 16 weeks) compared with male hosts [21]. The authors concluded that E2 in female recipients promoted more rapid β-cell maturation. Indeed, a large body of evidence demonstrates that E2 protects rodent and human islets from multiple metabolic injuries [Reviewed in [22,23]], and E2 promotes human islet engraftment and revascularization in diabetic mice [24]. Interestingly, long-term (35 weeks) graft function was higher in male hosts compared to females, potentially due to increased adipose tissue associated with the grafts in females [21].

2.3. Studies using animal models
Most rodent studies examining effects of gene and/or environmental manipulations on β-cell mass and function have traditionally focused only on male adults given the stronger diabetic phenotype compared to females. This biased approach already reflects differences between the two sexes and has led to a paucity of data on sex differences in islet gene expression or β-cell function or whether islets from males and females react differently to stressors. However, recent evidence suggests that GSIS is modulated in a sex-specific manner by gonadal hormones. For example, testosterone enhances GSIS in vivo in male mice via action on the androgen receptor (AR) in β cells [25]. In cultured male mouse and human islets, testosterone binding an extranuclear AR enhances cAMP production and the insulinotropic effect of GLP-1. In contrast, in females, E2 increases glucose-induced GLP-1 secretion in vivo and GLP-1 secretion from primary cultures of mouse and human β cells and intestinal explants through the activation of estrogen receptors (ERs) [26]. Thus, although male and female mammals exhibit the same overall mechanism of nutrient-induced insulin secretion, the fine-tuning of insulin secretion is regulated in a sex-specific manner by sex hormones. GSIS in rodents declines significantly with age in both sexes [27,28], however, isolated islets from female rats at 18 months of age still show higher glucose-stimulated insulin secretion (GSIS) ex vivo than those of males [27], confirming results obtained in humans [18]. Islets from elderly female rats showed elevated mitochondrial function (ATP content and oxygen consumption) compared with males when exposed to high glucose ex vivo. Mitochondrial biogenesis was also significantly higher in elderly female rats compared with males.

In multiple rodent models of diabetes in which β-cell failure is observed, there is also sex dimorphism. For instance, female animals are usually protected from development of the disease, with the exception of the NOD mouse as discussed in Section 4. Sexually dimorphic models including mice with streptozotocin-induced diabetes, the transgenic mouse overexpressing human islet amyloid polypeptide (hAPP) in β cells, or the Zucker diabetic fatty (ZDF) rat have been reviewed elsewhere and will not be discussed in detail here [22,23]. These models, however, have been instrumental in revealing the role of sex in islet pathophysiology and have contributed to identifying the main female ovarian hormone 17β-estradiol (E2) as a critical factor in protecting human islets from metabolic injuries including oxidative stress, gluco-lipotoxicity, and apoptosis.

3. ISLET AND β-CELL HETEROGENEITY
Islet endocrine cell composition is known to vary according to the anatomical location within the pancreas and among different species
and

In summary, in humans, female islets secrete more insulin in culture.

MicroRNAs were also differentially methylated.

BCL11A, HNF4a, CDKN2B, ATP11A, ADCY5

uncovered differential methylation in other diabetes and metabolism.

Over the past several years it has become apparent that individual β-cells from humans and animal models exhibit substantial heterogeneity in gene expression, electrical activity, secretory capacity, proliferation, and antigenicity [35–37]. These analyses have been greatly aided by the onset of single-cell ‘omics’ methodologies. Most animal studies have been undertaken with islets isolated from male mice, although in many publications, the sex from which the beta cells were obtained is not mentioned. In humans, four distinct subtypes of β-cells have been identified based on differential expression of cell surface markers [38]. In this study, islets from 16 non-diabetic donors (10M, 6F) and eight donors with T2D (2M, 6F) were assessed for the proportion of the four different β-cell subtypes. Although there is variability in the proportion of the four subtypes among individuals, and in general a decrease in the most commonly found β-cell subtype in individuals with T2D, there is no difference in subtype distribution between males and females. Studies using human cadaver donor islets also reveal differences in gene expression and epigenetic marks between males and females. For example, a study of 53 male and 34 female donors identified 2,853 sites on the X chromosome that were differentially methylated between males and females with >5% higher methylation in males [18]. These sites correspond to 757 individual genes out of a total of 890 genes on the X chromosome. There were no significant differences in age, body mass index (BMI), or hemoglobin A1c (HbA1c) between the two sexes. One of the genes that shows an increase in DNA methylation in male islets is DUSP9, a gene associated with insulin resistance and type 2 diabetes [39]. In addition, these analyses uncovered differential methylation in other diabetes and metabolism candidate genes located on autosomal chromosomes including BCL11A, HNF4a, CDKN2B, ATP11A, ADCY5, and IRS1. Several microRNAs were also differentially methylated.

In summary, in humans, female islets secrete more insulin in culture and in vivo, harbor more β-cells, and exhibit a lower degree of methylation of the X chromosome than male islets.

4. ISLET AND IMMUNE CELL INTERACTIONS

Studies in humans also show variable results with regards to sexual dimorphism in the presence of autoantigens associated with T1D progression. In one study of individuals at high risk for developing T1D, the frequency of autoantibodies against glutamic acid decarboxylase (GAD) was higher in females (88%) than males (71%), while insulin autoantibodies (IAA) were found at a higher prevalence in males (82% versus 60%) [40]. However, another study saw no difference in autoantibody prevalence in male versus female offspring of parents with T1D, although there was an increased prevalence of islet autoantibodies if the paternal parent had T1D versus the maternal parent [41]. Thus, T1D incidence is predominated by males and the onset of puberty is often associated with a decreased incidence of T1D in females suggesting that the female hormone E2 is protective [2–4,13]. Surprisingly, this sex dimorphism is the opposite in the NOD mouse model of T1D, which is characterized by a female predominance as will be discussed below. Several possibilities can explain this relative female protection from T1D in humans. T1D progression results from immunological defects in central and peripheral immune tolerance that culminates in the destruction of β-cells [42,43]. The immune system is one of the most sexually dimorphic. For example, females show heightened immunity to pathogens and are predisposed to autoimmune disorders compared to males [44,45]. This female predominance, however, is not observed for T1D. One possibility is that E2 protects female islets from apoptosis in vivo but also promotes immune tolerance. Consistent with the first possibility, E2 is known to protect human islets from multiple pro-apoptotic stimuli in vitro and in vivo (Reviewed in [22,23]). It is also plausible that in some women, decreased circulating levels of E2 contribute to defects in peripheral immune tolerance and progression to autoimmune responses in T1D. In support of this hypothesis, serum E2 levels and estrogenic activity are decreased in adolescents with T1D, and the potential protective effects of E2 are lost [46]. The immunological effects of E2 on the innate and adaptive immune arms of the immune system are being increasingly appreciated. The immunopathogenesis of T1D involves innate immune activation comprising of islet-resident macrophages and dendritic cells to engulf β cell autoantigens that can be ferried to the pancreatic lymph node to promote the activation of naïve autoreactive CD4 and CD8 T cells [47]. The activation of islet-resident macrophages and dendritic cells is a key initiating event since they can directly facilitate β cell destruction by generating pro-inflammatory cytokines, chemokines, and reactive oxygen species. Notably, E2 is protective against innate immune pro-inflammatory responses and prevents apoptosis of cultured human islets exposed to IFN-γ, IL-1β, and TNF-α [48]. Importantly, following transplantation in diabetic mice, islets treated with E2 ex vivo showed improved insulin secretion and were more efficient in restoring euglycemia than non-treated islets. Other beneficial effects of E2 on innate immunity were observed with Coxsackievirus-infected human islets in which E2 produced a profound decrease in the synthesis of the chemokine CXCL10 [49]. There is a wealth of evidence demonstrating an association of T1D onset with enteroviral infections [50,51]. The ability of E2 to decrease CXCL10 synthesis is significant since this pro-inflammatory chemokine can recruit autoreactive T cells into the islets to propagate β cell destruction [52]. E2 can also suppress pro-inflammatory cytokine synthesis from neutrophils, macrophages, and dendritic cells including TNF-α, IL-1β, IL-12, and IL-23 [53]. Finally, E2 can decrease the synthesis of pro-inflammatory cytokines including IL-6 from stimulated macrophages. Interestingly, genetic polymorphisms within the IL-6 promoter element result in an inability of E2 to promote resistance and downregulate IL-6 synthesis and may confer an early risk of T1D onset in females [54]. Therefore, in females, E2 could delay islet destruction in T1D by suppressing the innate immune response. However, E2 can also protect islets from the adaptive immune response. One of the best examples demonstrating that E2 promotes peripheral immune tolerance is the immunological effects of E2 on regulatory T (Treg) cell and invariant natural killer T (iNKT) cell differentiation and effector responses. Treg cells provide self-tolerance and are essential for suppressing autoimmune responses in individuals with T1D [55,56]. Previous studies have shown that E2 treatment promotes the expansion of immunosuppressive Treg cells including Foxp3 mRNA accumulation and protein expression to protect
female mice from experimental autoimmune encephalomyelitis, an autoimmune mouse model of multiple sclerosis [57]. Recent molecular mechanistic studies demonstrated that E2 treatment of Treg cells facilitates FOXP3 expression via estrogen receptor-α (ERα) binding eight estrogen response elements in the FOXP3 promoter [58]. E2 can also influence the expression of inhibitory receptors and anti-inflammatory cytokines expressed by Treg cells to facilitate peripheral tolerance including PD-1 and IL-10, respectively [59,60]. In addition to promoting the differentiation of immunosuppressive Treg cells, E2 also enhances the immunomodulatory function of iNKT cells. Although female NOD mice are predisposed to T1D, treatment of female NOD mice with E2 delays spontaneous autoimmune T1D partly due to the activation of iNKT cells [61]. Stimulation of iNKT cells with the glycolipid alpha-galactosylceramide (α-GalCer) was effective in delaying T1D in NOD mice by skewing T cell cytokine responses to a less inflammatory Th2 phenotype and also by recruiting tolerogenic dendritic cells to promote Treg cell differentiation [62,63]. Notably, the immunomodulatory effects of iNKT cells to delay autoimmune diabetes could be enhanced with combinatorial E2 and α-GalCer treatment [61,64]. Interestingly, the NOD mouse model of spontaneous autoimmune diabetes, when housed in specific pathogen free (SPF) conditions, displays a female sex bias in disease incidence (F:M, 2:1). Under germ-free (GF) conditions, this sex dimorphism is not observed suggesting that the microbiome is instrumental in the female bias [65]. Transfer of adult NOD male intestinal microbiota into female NOD mice increased serum testosterone levels and protected from T1D [65]. Another study also reported that colonization of GF NOD females with bacterial taxa from SPF males also increased serum testosterone levels [66]. Notably, the protection against T1D induced by male microbiome transfer was abolished in female recipients treated with an androgen receptor antagonist, again implicating testosterone in the microbiome transfer was abolished in female recipients treated with an androgen receptor antagonist, again implicating testosterone in the progression to autoimmunity in the NOD mouse.

5. α-CELL FUNCTION AND DYSFUNCTION

5.1. Counter-regulatory responses to hypoglycemia

Insulin-induced hypoglycemia is counterregulated by glucagon, epinephrine (adrenaline), norepinephrine (noradrenaline), cortisol, and growth hormone [67]. Sex differences in counter regulation and especially α-cell function has been a matter of debate for years. Historically, premenopausal women were reported to exhibit lower blood glucose than men during a prolonged 72 h fasting providing the first indication of gender difference in the counterregulation to hypoglycemia [68]. However, glucagon concentrations were not assessed and the role of the α-cell vs catecholamines in counter-regulation is unknown. Subsequent studies assessing the effect of gender on catecholamine responses to hypoglycemia during euglycemic-hypoglycemic clamps showed that catecholamines (epinephrine, norepinephrine) and/or cortisol/GH responses to hypoglycemia, but not to glucagon, were diminished in healthy women compared to men [69,70]. Since the autonomic nervous system (ANS) contributes to increased glucagon secretion during insulin-induced hypoglycemia in rodents and primates [71], the diminished catecholamine responses to hypoglycemia would be predicted to reflect lower glucagon responses to hypoglycemia in women. Indeed, a study reported that the glucagon response to insulin-induced hypoglycemia was blunted in nondiabetic women in the presence of the ganglion nicotinic receptor trimethaplan [72], suggesting that the ANS mediates the majority of the glucagon response to insulin-induced hypoglycemia. Further, Davis et al. performed glucose clamps on healthy human subjects and reported that females showed lower glucagon response to equivalent fixed hypoglycemia and hyperinsulinemia compared to age-and-BMI-matched males [73]. However, during low-dose insulin-induced hypoglycemia, suppression of endogenous glucose output was more prolonged in healthy women compared to men but catecholamine and glucagon concentrations did not differ between the two sexes during this challenge [69]. In mice, neuroglucopenia is associated with a stronger glucagon response in females [74]. In studies targeting healthy young Japanese adults, females exhibited a stronger suppression of glucagon secretion after an oral glucose load compared with males [10]. These data are consistent with female mice exhibiting an increased sensitivity of α-cells to cholinergic activation of glucagon secretion that in turn led to larger glucagon response to hypoglycemia [74].

In summary, it appears that women exhibit reductions in the counterregulatory hormones, glucagon, and epinephrine, together with blunted rates of endogenous glucose production compared to men. These differences could explain why blood glucose concentrations fall to lower levels during fasting in women. It is unclear, however, if this sex difference in the glucagon response to insulin-induced hypoglycemia reflects differences in autonomic activation or in direct pancreatic α-cell regulation.

5.2. Sex hormones and glucagon secretion

Early studies reported an increase in glucagon content in pancreases from female mice [75,76]. The glucagon level in the portal vein was reduced relative to insulin in estrogen-treated ovariectomized rats [77]. In rodent models, ovariectomy increased circulating glucagon, and this effect was reversed by E2 administration [78]. Similarly, following islet transplantation, E2 suppresses hyperglucagonemia in hyperglycemic mice [24]. The effect of E2 is direct since the estrogen receptor-α (ERα) and the G protein-coupled ER (GPER) are expressed in islet cells [79], and E2 decreases glucagon secretion from islets isolated from males and females [80]. In addition, the GPER agonist G-1 inhibited glucagon secretion in cultured islets from female wild type mice to an extent similar to E2 [81]. However, it is unknown if these effects are mediated via a direct glucagonostatic action on ERs in α-cells or an indirect insulinotropic effect on ERs in β cells. Further, the above studies were undertaken in animal models, and the direct effects of estrogens on glucagon secretion in humans have not been studied. Mechanistically, endocrine disruptors such as bisphenol A (BPA) and diethylstilbestrol (DES), which mimics E2, were shown to suppress low glucose-induced intracellular calcium ion ([Ca2+]i) oscillations, which triggers glucagon secretion from α-cells [82]. However, a detailed morphological analysis including single cell approaches in different genders has not been reported.

6. CROSS-TALK BETWEEN ORGANS AND ISLET CELLS

Recent reviews have discussed the importance of organ cross-talk with regard to the regulation of islet biology [83]. We will focus on the effect of male and female gonadal hormones on islet cell biology.

6.1. Ovarian—islet axis in female

In women, the ovarian—islet axis influences β-cell biology during reproductive years and at menopause. As discussed earlier in Section 5, estrogens increase GLP-1 production in mice. Sex-specific effects of estrogens and progesterone on β-cell survival, function, and
proliferation during pregnancy have been reviewed elsewhere [22].

Prolactin secreted from the pituitary and placental lactogen contribute to the expansion of β-cell mass during pregnancy [84]. Prolactin and lactogen mediate their actions on β-cell proliferation through HGF, Menin, serotonin, and/or osteoprotegerin pathways [85–89]. However, the factors that promote maternal β-cell adaptation during pregnancy in humans are still unclear and warrant additional studies. Studies of the effect of menopause on islet function in women suggest that it alters insulin secretion [90]. However, menopausal changes in insulin clearance and metabolism make this assessment particularly difficult to interpret.

6.2. Testicular-islet axis in males

In men, the testicular-islet axis also impacts islet biology via production of testosterone. Men with primary testosterone deficiency due to androgen-depletion therapy (ADT) for prostate cancer exhibit an increased diabetic risk. In two, large, population-based studies of men with prostate cancer, ADT with gonadotropin releasing hormone (GnRH) analogs was associated with a 28–44% increased risk of incident diabetes compared to controls [91,92]. Thus, severe testosterone deficiency is instrumental in predisposing to hyperglycemia and diabetes in these patients. Obviously, the diabetogenic effect of androgen depletion in men involves the combination of insulin resistance and visceral adiposity [93]. However, evidence suggests that androgen action in β-cells enhances β-cell function in men [94]. First, testosterone increases GSIS via AR in human islets from male donors [25]. Second, in hypogonadal men with T2D, testosterone therapy improved β-cell function (measured by HOMA %B) [95]. In fact, women with ovarian hyperandrogenism also exhibit β-cell dysfunction. Some show higher basal insulin secretory rates and attenuated secretory responses to meals [96] or exhibit exaggerated acute insulin response to glucose [97]. These abnormalities are closely associated with testosterone levels [98] suggesting that excess testosterone leads to insulin hypersecretion in women [99]. Further, testosterone produces insulin hypersecretion from cultured islets of human female donors and chronic testosterone administration to female mice promotes insulin hypersecretion, islet oxidative injury and secondary β-cell failure as a result of AR activation in β-cells [100].

7. IN UTERO β-CELL PROGRAMMING

The Developmental Origins of Health and Disease (DOHaD) hypothesis states that the in utero environment impacts postnatal susceptibility to disease. Exposure to either under-nutrition or over-nutrition in utero has been shown to lead to increased susceptibility to obesity and diabetes in the offspring in humans and in animal models (reviewed in [101]). Taken together, data reveal sexual dimorphism in susceptibility to environmental manipulations during critical periods of pancreas development that impact β-cell differentiation.

7.1. Effect of maternal obesity

Offspring born to overweight or obese mothers are more likely to be obese as children and into early adulthood [102]. Additionally, offspring of mothers who were obese before and during pregnancy are more likely to develop insulin resistance and impaired glucose tolerance as adults independent of birth weight [103]. Given the fact that nearly 50% of women of child-bearing years are currently overweight or obese [104], understanding how the in utero environment affects islet development, mature function, and adaptability to stress has become increasingly important. Molecular mechanisms underlying these phenomena are likely to include epigenetic modifications. Several different animal models have been used to study the effects of in utero over-nutrition on islet structure and function (reviewed in [101]). Three common threads emerge when considering the body of animal studies as a whole. First, overwhelming evidence of maternal HFD results in significant changes in islet architecture and/or function, many of which would be predicted to increase susceptibility to Type 2 diabetes later in life. Second, although changes in α- or β-cell mass may vary, there is usually evidence of impaired islet function. Finally, in the minority of studies that focused on sex differences, there were observable differences in islet function between males and females exposed to HFD in utero, although which sex was more severely affected depended on the design of the study.

The non-human primate represents a model in which islet architecture, composition, and gene expression more closely resembles humans [31]. Studies in Japanese macaques reveal that in utero HFD exposure results in increased fasting blood glucose and decreased first phase insulin secretion in both males and females at 13 months, with males specifically showing an increased in total pancreas IL1-β (post-weaning) [105]. In one study using C57Bl/6J dams placed on a HFD during pregnancy and lactation, male and female offspring were compared after weaning onto a control Chow diet (CD) or maintenance on the maternal HFD [106]. Adult female (but not male) HFD-exposed offspring that were maintained on the HFD post-weaning showed a significant increase in body weight, plasma insulin, and blood glucose compared with CD-exposed females weaned on HFD. However, β-cell compensation and adaptability seem to be preserved in females exposed to HFD in utero, since islet size and insulin content were increased when either CD- or HFD-exposed females were placed on HFD post-weaning. In contrast, males exposed to HFD in utero showed higher blood glucose levels but lower plasma insulin and decreased islet insulin content as adults regardless of post-weaning diet. HFD-exposed males also showed reduced islet area and, unlike female offspring, were unable to expand islet mass when maintained on HFD post-weaning. Defects in β-cell compensation in response to post-weaning HFD were also observed in a rat model of maternal over-nutrition. Expression of the β-cell transcription factor Pdx1 was decreased in islets from males exposed to maternal HFD regardless of post-weaning diet, while female offspring exposed to maternal HFD had increased Pdx1 expression regardless of post-weaning diet. Islets from HFD-exposed males showed increased expression of NADPH oxidase and superoxide production. Other measures of oxidative stress were also higher specifically in HFD-exposed males regardless of the post-weaning diet. Thus, despite HFD-exposed female offspring having impaired glucose tolerance, they seem to be protected from oxidative stress and β-cell dysfunction. In rats, female offspring were more affected by maternal HFD than were male offspring [107]. Females had increased β-cell mass, but reduced expression of GLUT2 and decreased GSIS. In contrast, paternal high fat diet consumption and insulin resistance in rats is associated with impaired insulin secretion in female but not male offspring [108]. In this study, female offspring of obese fathers had reduced β-cell mass due to smaller islets, suggestive of decreased replication. Gene expression analyses of islets from female offspring revealed decreased expression of genes involved in cell proliferation, metabolism, and granule exocytosis.

7.2. Effect of maternal undernutrition

There is some evidence in humans that adverse in utero exposures have differential effects on male and female offspring that can increase risk for metabolic disease. For example, changes in DNA methylation at
metabolic gene loci differed between male and female offspring born during the Dutch Hunger Winter in 1944–45 [109]. At age 50, women born during the Dutch Hunger Winter had increased BMI and waist circumference compared with men [110]. However, most of our current understanding of DOHaD comes from studies in animal models. In animal models of intrauterine growth restriction (IUGR), male offspring tend to show impaired insulin sensitivity as adults, while females tend to have reduced insulin secretion [111]. In rats, either IUGR or maternal under-nutrition results in decreased β-cell in both sexes [107,112,113]; however, males had a higher pancreatic insulin content than females [107] and females showed reduced expression of the critical β-cell transcription factor Pdx1 as well as insulin [113]. In mice, male offspring whose mothers were exposed to a low protein diet during gestation had decreased β-cell mass and a lower mitochondrial-nuclear DNA ratio compared with controls [112]. Expression levels of genes involved in the TCA cycle were reduced in islets of both males and females, while higher ROS production was observed specifically in islets from male offspring. The defective mitochondrial gene expression and increased ROS likely contributes to β-cell dysfunction and in the longer-term, β-cell demise, potentially leading to increased susceptibility to T2D later in life.

7.3. Effect of maternal hyperglycemia

There is evidence that human exposure to hyperglycemia in utero has differential effects on male and female offspring that can increase risk for diabetes. For example, human fetal exposure to maternal T1D is associated with altered GSIS in adult offspring of both sexes independent of adiposity and insulin resistance [114]. Notably, this β-cell defect is characterized by a sexual dimorphism [115]. In response to oral glucose stimulation, male and female offspring of diabetic mothers displayed reduced insulin secretion. In contrast, in response to I.V. glucose stimulation, only female offspring of diabetic mothers exhibited decreased insulin secretion. This suggests that males are more protected against or can compensate for the deleterious effect of maternal hyperglycemia on fetal β-cells. In a mouse model of hyperglycemia during pregnancy [116], both male and female offspring showed impaired glucose tolerance (although this was more pronounced in males). Interestingly, the F2 generation from these hyperglycemia-exposed offspring also showed impaired glucose tolerance regardless of whether it was the male or female parent who had been exposed to in utero hyperglycemia (again, male F2 offspring were more severely affected). Expression of two differentially imprinted genes, insulin-like growth factor 2 (Igf2) and the long non-coding RNA H19, was decreased in islets from both males and females, but again, the magnitude of the decrease was greater in male offspring than female offspring.

7.4. Effect of in utero environmental exposures

Studies in mice also suggest that male and female offspring respond differently to in utero environmental exposures. For example, exposure of dams to bisphenol A (BPA), a xenoestrogen used in the manufacturing of some plastics, causes a reduction in β-cell mass and impairs GSIS in male offspring at the F1 and F2 generation, but has no effect on these parameters in female offspring [117]. BPA influences on pancreatic islet mitochondrial function could play a role since abnormal mitochondrial number and function generates reactive oxygen species (ROS) and affects GSIS. Indeed, in rodent islets, BPA exposure is known to alter expression of key mitochondrial genes such as oxoglutarate dehydrogenase (Ogdh) and uncoupling protein 2 (Ucp2) [118,119]. Thus, insulin secretory defects in males exposed to BPA may be due to greater susceptibility of mitochondria in males to environmental insults. Although there were no differences in total oxygen consumption in islets isolated from males exposed in utero to BPA, there were reductions in basal and maximal oxygen consumption. Ogdh expression was mildly increased and Ucp2 levels were significantly increased in BPA-exposed males compared with control males consistent with mitochondrial impairment. In BPA-exposed mouse dams, metabolic phenotypes in the F1 and F2 male offspring were linked to fetal over-expression of Igf2 [120]. Both Igf2 and H19 are involved in growth, proliferation and weight gain.

8. CONCLUSION AND FUTURE PERSPECTIVES

Evidence presented in this review underscores the influence of sex and gender that contribute to differences in islet cell function and susceptibility to failure (Figure 1). These differences represent sex-related
biological factors that can be harnessed for developing better approaches to prevent and/or treat diabetes. However, major obstacles that currently prevent scientific progress in sex differences in islet biology should be addressed. For one, there are no suitable male and female islet cell lines available to study sex differences in culture. Establishing genetically suitable immortalized human islet lines of both sexes seems unrealistic. The utility of primary islet cells from inbred rodents of the same genetic background is limited due to their short lifespan in culture and the inability to expand and passage these cells. The development of clonal, immortalized cell lines from littermate rodents of the same genetic background is limited due to their short lifespan in culture and the inability to expand and passage these cells. The development of clonal, immortalized cell lines from littermate inbred animals seems a logical first step in creating suitable male and female islet cell lines. While the comparison of cultured male and female islet cells oversimplifies the biology of sex because of the limitations of the in vitro environment, their availability as in vitro ‘tools’ will be useful for certain experiments that are difficult to undertake using primary islet tissue. Because of the sex differences described above, the in vivo environment of male and female islets differs in multiple factors including hormones, metabolites, and neural inputs. These defined phenotypic sex differences cannot be fully reproduced in culture. Finally, studies on sex differences will inform islet transplant procedures in male and female recipients and will generate for comparison of embryonic or adult stem cell derived islet cells for research and therapy that continues to be an important aim in diabetes research. In conclusion, the current research tools are not fully adapted to the study of sex differences in islet cell biology and dysfunction. The recent advances in technology should allow the development of modern research tools for experimental purposes as well as to contribute to precision and personalized medicine in the context of gender-specific therapies for diabetes.

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MG edited the review and contributed to sections on β-cell function, heterogeneity, and in utero programming. RNK edited the review and contributed to sections on organ crosstalk and alpha cells and acknowledges assistance from J. Shirakawa MD PhD (Joslin Diabetes Center, Boston and Yokohoma University, Japan) and K. Shibue MD PhD (Joslin Diabetes Center). HT contributed to section on islet cell and immune interactions. FMJ contributed to all sections and edited the manuscript.

CONFLICT OF INTEREST

None declared.

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