Sex dependent impact of gestational stress on predisposition to eating disorders and metabolic disease

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ABSTRACT

Objective: Vulnerability to eating disorders (EDs) is broadly assumed to be associated with early life stress. However, a careful examination of the literature shows that susceptibility to EDs may depend on the type, severity and timing of the stressor and the sex of the individual. We aimed at exploring the link between chronic prenatal stress and predisposition to EDs and metabolic disease.

Methods: We used a chronic variable stress protocol during gestation to explore the metabolic response of male and female offspring to food restriction (FR), activity-based anorexia (ABA), binge eating (BE) and exposure to high fat (HF) diet.

Results: Contrary to controls, prenatally stressed (PNS) female offspring showed resistance to ABA and BE and displayed a lower metabolic rate leading to hyperadiposity and obesity on HF diet. Male PNS offspring showed healthy responses to FR and ABA, increased propensity to binge and improved coping with HF compared to controls. We found that long-lasting abnormal responses to metabolic challenge are linked to fetal programming and adult hypothalamic dysregulation in PNS females, resulting from sexually dimorphic adaptations in placental methylation and gene expression.

Conclusions: Our results show that maternal stress may have variable and even opposing effects on ED risk, depending on the ED and the sex of the offspring.

1. INTRODUCTION

Eating disorders (ED) are damaging mental and metabolic illnesses that dramatically diminish the quality of life and induce life-threatening side effects [1]. EDs predominantly affect women [2,3] and have an adolescent/young adulthood onset [4]. The main EDs include anorexia nervosa (AN, largely characterized by self-starvation), bulimia nervosa (BN, compulsive/binge eating followed by compensatory behaviors) and binge eating disorder (BED, compulsive/binge eating without compensatory behaviors). While the early origins of these disorders remain largely unknown, recent evidence suggests that epigenetic mechanisms may be involved in initiating and maintaining them [5,6]. Epigenetic changes involved in complex adult psychiatric and neurodegenerative disorders include DNA methylation/hydroxymethylation, histone acetylation/deacetylation, noncoding RNAs and microRNAs. While epigenetic mechanisms are known to modulate behavior and health at various stages during the life cycle, it is in the womb where these processes begin to shape the exposed offspring, with a potentially adverse impact on later physical health, emotional adjustment, and stress reactivity. This maladaptation is particularly frequent in response to intrauterine stress, with the general consensus anticipating that early life stress can dramatically increase the probability of developing an ED later in life [1,7—9]. However, the frequent positive association between early life stress and EDs reported in epidemiological studies may be incomplete. There is an imperative need to make appropriate distinctions between different types of early life stress (e.g., anxiety/depression, trauma, nutrition or obstetric complications) and the different developmental windows at which these stressors occur (e.g., gestation/whole/early/late), lactation, childhood or adolescence.

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Abbreviations: ABA, activity based anorexia; BE, binge eating; EDs, eating disorders; FR, food restriction; HF, high fat; PNS, prenatal stress

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to truly understand the link between early life factors and subsequent predisposition to disease. It is also necessary to make a clear distinction between the different EDs, which may have very different developmental origins.

This distinction must also be applied to the overweight extreme of the eating disorder spectrum (obesity with metabolic syndrome). In utero stress is broadly considered a contributor to subsequent risk of obesity and metabolic dysfunction [10,11]. However, when males are exposed

![Figure 1: Prenatal stress (PNS) causes basal metabolic abnormalities specifically in female offspring.](image)

(A) Experimental design. (B) PNS dams weighed less than controls (CTRLs) during gestation \( F_{(1,18)} = 7.01, p = 0.016 \). (C) Corticosterone levels were higher in PNS dams \( t_{(18)} = 5.15, p < 0.0001 \) and fetuses \( t_{(18)} = 5.79, p < 0.0001 \) on GD17.5 compared to CTRLs. (D) Maternal behavior in the first and third postpartum weeks (PPW) was similar between the groups. (F–J) PNS female offspring displayed similar BW (F) and heat production (H) but tended to high adiposity (J) and displayed higher food consumption \( (F_{(1,14)} = 6.19, p = 0.026) \) (G) and activity levels \( (F_{(1,14)} = 7.68, p = 0.015) \) (I) than CTRLs. (K–Q) Male PNS offspring displayed a normal metabolic profile. (P–Q) Glucose tolerance was normal but insulin tolerance was affected by PNS in both sexes \( (F_{(1,15)} = 5.60, p = 0.033 \) for females (P) and \( F_{(1,15)} = 7.39, p = 0.017 \) for males (Q). Glucose tolerance test (GTT) is displayed on the left y-axis and insulin tolerance test (ITT) on the right y-axis. (R) Plasma corticosterone levels were lower in PNS than in CTRL females \( (t_{(10)} = 2.55, p = 0.029) \). Data presented as mean and S.E.M. N = 6–10.
to prenatal stress in early gestation, they have a reduced body weight and adiposity when exposed to a high fat (HF) diet [12]. Sex-specific differences in the response to ED modeling challenges may be linked to placental expression of O-linked bN-acetyl glucosamine (O-GlcNAc) transferase (OGT). This enzyme, which senses changes in maternal energy homeostasis and regulates epigenetic marks on chromatin, is reduced by early gestational stress in males only and therefore may regulate sex-specific epigenetic modifications of genes important for adult metabolism [13]. In contrast, the impact of late-gestation prenatal stress on adult responsivity and metabolism appears to be more profound in females [14—16].

According to clinical and preclinical studies, intrauterine fetal stress that occurs during sexual differentiation induces sex-dependent effects on fetal brain development. These effects occur within highly sexually dimorphic regions that regulate mood, the stress response, metabolic function, and the autonomic nervous system. Thus, maternal stress has been shown to have a sex-dependent impact that is timing specific during gestation, affecting different developing brain-areas and, consequently, resulting in multiple systemic sex-dependent effects on metabolic functions, ultimately predisposing male and female offspring to different diseases [17—19]. Although fetal programming, mediated by the placenta, may be intended to benefit the health and survival of the offspring, it can lead to a variety of complications when the developmentally adapted organism is exposed to unexpected challenges throughout its life. Furthermore, compromised placental function can have both short- and long-lasting consequences for the developing fetus [17], as is the case for maternal hypoxia [20], preeclampsia [21], or placental insufficiency [22].

Here, we explored the effects of chronic variable stress during the whole period of gestation on basal metabolism, the response to food restriction (FR), and susceptibility to develop activity-based-anorexia (ABA), BE and metabolic syndrome in male and female offspring. We show that fetal programming by PNS is mediated by global and robust sex-specific alterations in placental DNA/RNA methylation and gene expression. Finally, we focused on hypothalamic programming by PNS specifically in females and detected a variety of abnormalities that underlie the response to the different ED-inducing protocols. Altogether, we report that chronic PNS induces sexually dimorphic effects on placental function, affecting fetal hypothalamic programming and subsequent basal metabolism and the response to a variety of different metabolic challenges.

2. RESULTS

2.1. Prenatal stress induces basal metabolic abnormalities in female offspring

To examine the potential link between gestational stress and a later predisposition to eating and metabolic disorders, we examined the metabolic profile of prenatally stressed males and females under basal conditions. To create these groups, we exposed pregnant ICR/CD1 dams to a chronic variable stress (prenatal stress - PNS) protocol for most of the pregnancy period, beginning a day after the detection of the copulation plug (gestation day (GD)1.5) and finishing on GD16.5 (Figure 1A). PNS dams gained less weight and had higher corticosterone (CORT) levels towards the end of the protocol (Figure 1B,C), which was already reported in a previous study [23]. PNS fetuses also displayed higher CORT levels but showed similar body weight at birth compared to controls (CTRLs; Figure 1C,D). Maternal behavior did not differ between the groups, excluding this environmental factor as a potential confounding variable (Figure 1E, [23]). Body weight follow-up of the offspring showed no differences between the groups in either sex (Figure 1F,K). At around 10 weeks of age, the offspring’s metabolic profile was examined using metabolic cages. PNS females showed higher basal food intake (Figure 1G), similar heat production (Figure 1H), hyperactivity (Figure 1I) and a trend to increased body adiposity (Figure 1J) compared to CTRLs. PNS males showed a similar basal metabolic profile (Figure 1L—N) and adiposity (Figure 1O) compared to CTRLs. Glucose sensitivity was not affected by PNS in either sex, but insulin sensitivity decreased by PNS in both sexes (Figure 1P,Q), a finding previously reported in humans [10,11]. Finally, CORT levels were lower in adult PNS offspring (Figure 1R) [10]. Thus, PNS appears to have stronger basal metabolic effects in female than in male offspring.

2.2. Prenatally stressed animals cope better with short-term food restriction

As a first step to examine the response of prenatally stressed offspring to later life metabolic challenge, we exposed the offspring to 4 days of moderate FR, starting on postnatal day 40 (Figure 2A). While CTRL females reacted to the challenge with moderate weight loss from the first day, PNS females did not show the same weight drop but rather maintained their weight throughout the protocol (Figure 2B). During the challenge, PNS females consumed fewer calories than CTRLs (Figure 2C). Examination of their metabolic profile post-recovery exposed hypophagia during the dark phase, decreased heat production and similar activity compared to CTRLs (Figure 2D). Male offspring showed a similar but more moderate response (Figure 2E—G). The 4-day FR manipulation resulted in increased post-recovery adiposity in females (Figure 2H), but not in males (Figure 2I) and did not affect glucose or insulin tolerance. In fact, the moderate insulin intolerance observed under basal conditions in PNS mice disappeared in both sexes (Figure 1N—O).

2.3. Prenatal stress abolishes the innate predisposition to activity based-anorexia in adolescent females and causes metabolic adaptations in males

To assess the susceptibility and metabolic response of PNS offspring to ABA, we exposed the adolescent offspring (age 30 days) to a modified version of the ABA protocol [23,24]. The ABA model mimics the choice of exercise over eating, relying on limited food intake while allowing unlimited access to running wheels (RW) [25,26]. The limited period of food access allows them to retain the majority of their BW as long as they engage in compensatory food intake when accessible, instead of excessive wheel running. The ABA protocol is characterized by adolescent vulnerability and sex specificity to females [23,27,28], hyperactivity [23,25,29], dramatic weight loss as a consequence of decreased food intake [23,24,30], and circadian disruption in the running pattern [23,28]. Briefly, we habituated the animals to voluntary wheel running for one week with free access to food, followed by four days of free access to wheels combined with gradual FR in the form of 3—4 h of food access a day during the dark cycle (Figure 3A). As expected, CTRL females split into ABA-prone and resilient, with ABA-prone females showing dramatic weight loss, decreased food intake, increased running, and circadian disruption (Figure 3B), all of which was reported in a previous study [23]. ABA-prone females showed increased food intake and activity in the light phase and higher heat production post-recovery compared to CTRL resilient females, highlighting the long-lasting circadian disruption in this group (Figure 3C). As previously reported, none of the PNS females developed ABA [23] and showed a similar profile to resilient CTRL females (Figure 3B,C).

In an extended cohort, we found here that the ABA challenge did not affect body adiposity in CTRL females, but did result in higher body
adiposity in (ABA resilient) PNS females compared to resilient CTRLs (Figure 3F). As expected, males were largely resilient to the protocol regardless of prenatal treatment and beside one male in the CTRL group, all others coped remarkably well with the ABA manipulations (Figure 3D). However, coping with ABA resulted in decreased heat production and hyperactivity in PNS males (Figure 3E), without affecting adiposity (Figure 3G). Glucose tolerance was normal in all groups and the moderate insulin intolerance observed under basal conditions in PNS animals disappeared in both sexes (Figure 3H,I), as seen after FR (Figure 2J,K). Altogether, the female PNS group showed a remarkable improvement in their capacity to cope with the ABA-inducing protocol compared to CTRL females, an adaptation that may grant them increased survival chances in a food deprived environment.

2.4. Prenatal stress abolished the propensity of females to binge on palatable food, but induced metabolic abnormalities in response to intermittent access to Western diet in both sexes

Since PNS females appear to be more efficient than CTRLs at coping with FR and ABA, gaining body fat after both manipulations, we next explored their vulnerability to binge on HF/high palatable food when exposed to a BE-inducing manipulation. Using the intermittent access protocol [23,31,32], we habituated the offspring to Western diet (WD) for 5 days, switched them back to chow, and then allowed them access to WD for 2 h a day, 3 days a week for a period of 4 weeks (a total of 12 BE sessions) (Figure 4A). Contrary to our expectations, PNS females gained similar weight (Figure 4B), consumed less WD during the habituation period (Figure 4C) when compared to CTRL group and overall did not develop BE (Figure 4D). Thus, the PNS females’ response appears to reflect anhedonia rather than the expected compulsivity observed in a different model of late gestational stress [15]. From a metabolic perspective, PNS females exposed to this protocol showed decreased food intake post-recovery, lowered metabolic rate, and similar activity in comparison to CTRLs (Figure 4E). In contrast to females, PNS males showed a significantly different pattern of BW progression throughout the BE protocol compared to CTRLs (Figure 4F). While they showed anhedonia (or maybe hypophagia due to increased anxiety/neophobia) to WD during the habituation period similarly to PNS females (Figure 4G), they displayed a moderate and gradual increase in WD consumption compared to CTRLs.
males (Figure 4H). In contrast to the PNS females, the BE protocol induced an increase in metabolic rate in males (Figure 4I) without affecting any other metabolic parameters. The BE protocol did not affect body adiposity in males but tended to increase it in females (Figure 4J,K). Glucose tolerance was not affected in either sex (Figure 4L,M). The insulin tolerance test revealed a different pattern of response in PNS females compared to CTRLs (Figure 4L) and a tendency to improvement in PNS males compared to CTRLs (Figure 4M). Taken together, the BE protocol did not induce the expected compulsive consumption of palatable food in PNS females, but exposed metabolic abnormalities leading to decreased heat production, a potential mechanism of exaggerated long-term weight/adiposity gain. In males, the pattern was the opposite, and the increased heat production and improved insulin tolerance may paradoxically hint at a better coping mechanism when exposed to high caloric diets.

2.5. Prenatal stress programs females to diet induced obesity and males to leanness when exposed to high fat diet

In order to explore the possibility that females may be programmed to gain adiposity/weight, while males may be programmed to lose adiposity/weight as a result to PNS, we exposed a further set of PNS offspring to 3 weeks of 60% HF diet (Figure 5A). After a period of recovery, we found that PNS HF females gained significant amounts of weight compared to CTRLs (Figure 5B), while PNS males gained less weight than CTRLs (Figure 5C). Female PNS HF offspring consumed similar amounts of food post-recovery but showed dramatically lower heat production (Figure 5D). Male PNS HF offspring consumed similar amounts of food post-recovery but showed dramatically higher heat production and hyperactivity (Figure 5E). These findings led to higher adiposity in PNS females (Figure 5F) and lower adiposity in PNS males compared to CTRLs (Figure 5G). Exposure to HF diet had no evident effects on glucose and insulin sensitivity in PNS females (Figure 5H), but dramatically affected PNS males. Remarkably, PNS males showed improved glucose tolerance and insulin sensitivity after HF (Figure 5I).

2.6. Prenatal stress programs the female fetuses to abnormal metabolic responses through robust placental adaptations

In order to explore the origins of the metabolic responses of PNS offspring to the different protocols, we next focused on the placenta. We first explored the effects of PNS on global DNA (5-mc) methylation (Figure 6A), DNA (5-hmc) hydroxymethylation (Figure 6B), and RNA (m6A) methylation (Figure 6C) of placentas of both sexes. These...
two processes are known to be important regulators of gene expression in the placenta [33,34] and to be affected by stress [35]. N6-Methyladenosine (m6A) RNA methylation, not studied in the placenta until now, is an abundant modification in mRNA preferentially enriched within 3′UTRs and around stop codons, allowing for potential alternative polyadenylation and miRNA binding regulation[36]. We found that PNS increased DNA methylation (Figure 6A) and reduced RNA methylation (Figure 6C), without affecting DNA hydroxymethylation in females (Figure 6B), hinting at major global effects of PNS on placental function. In males, the results were not significant. Accordingly, enzymes responsible for these processes, such as DNA methyltransferase (Dnmt)3a (Figure 6D), the Ten-eleven translocation (Tet) demethylation enzymes [22] (Figure 6E), the RNA methylation regulatory subunits WT1 associated protein (Wtap) and KIAA1429 and the demethylases Fat mass and obesity-associated protein (Fto) and Alkylated DNA repair protein alkB homolog (Ailkbh)5 (Figure 6F,G) were all increased by PNS specifically in females. Lastly, to gain insight into the potential effects of PNS on gene expression and function of the uterinal environment that the fetuses were exposed to, we performed RNA-Seq comparing CTRLs and PNS female placentas. Pathway enrichment analysis revealed significant enrichment for 15 KEGG signaling pathways (Table S1). Among these, strong changes were observed for the ABC transporters (q-value = 0.073), (Figure 6H, Table S2), circadian entrainment (q-value = 0.060) (Figure 6I, Table S2), cytokine cytokine receptor interaction (q-value = 0.004) (Figure 6J, Table S2) and the oxytocin pathway (q-value = 0.073) (Figure 6K, Table S2). Finally, we focused on the solute carrier (SLC) group of membrane transport proteins and the insulin-like growth factor (IGF) family, which play a key role in placental structure and function[23,37]. Interestingly, PNS placentas express lower expression levels of the amino acid transporters Slc6a15, Slc1a4, Slc6a17, and Slc43a1, the choline transporters Slc44a5 and Slc22a4 and the B Vitamin transporters Slc5a6 (B5, B7) and Slc19a2 (B1) potentially affecting methylation processes in the fetus. In contrast, several other
far more abundant amino acid transporters were upregulated, such as Slc38a6, Slc36a1, Slc43a2, and, in particular, the Slc7 family (Slc7a1 and Slc7a4 (CATs) and Slc7a5, Slc7a7, Slc7a8 (LATs)). In addition, PNS placentas further displayed higher expression levels of the main glucose transporter Slc2a1 and Igf2 (Figure 6L). All significant genes are shown in Table S3.

2.7. Prenatal stress affects life-long hypothalamic gene expression

To explore the mechanisms underlying our metabolic findings (summarized in Table 1) and the impact of differential placental methylation caused by PNS, we examined gene expression in the adult hypothalamus using a whole-genome gene expression array comparing CTRL and PNS females. Gene ontology analysis revealed abnormalities in a variety of biological processes (Figure S1), including upregulation of the innate immune response and the response to stress (Figure 7A). Long-lasting inflammation in different areas of the brain has previously been reported in the context of PNS [38,39] and likely partially underlies the observed long-term metabolic abnormalities and response to challenge found here and by others [40]. Our findings further include upregulation of 17 long intergenic non-coding RNAs (lincRNAs) (Table S4). LincRNAs functions are largely unclear, but they have recently been implicated in the cellular stress response [41] and in anxiety-like behavior in mice [42]. But more remarkably, the long-term effects of PNS on the adult female hypothalamus are linked to long-lasting alterations in genes linked to neuronal processes, more notably in the vesicle,
synapse, axon, dendrite, presynapse and neuronal cell body (Figures 7A, S1 and S2), suggesting major structural and/or functional remodeling in this group (all predictions based on STRING database [43]).

Finally, we focused on a few individual genes affected by PNS that are heavily involved in the maintenance of energy homeostasis and the response to stress [44–47] (Table S4). Among them, we validated oxytocin (Oxt), arginine vasopressin (Avp), adrenergic receptor beta 3 (Adrb3), neurenomed S (Nms), and neuronatin (Nnat), most of which are involved in many of the processes revealed by GO analysis (Figure 7A,C). Oxt, Avp, Adrb3, and Nnat were downregulated, and Nms was upregulated by PNS (Figure 7B). Finally, in order to test the fetal origins of the observed phenotypes, we examined the expression levels of these genes in the fetal hypothalamus. We found significant alterations in all five genes in response to PNS, suggesting these abnormalities are programmed in utero and remain into adulthood, producing the abnormal responses of PNS females to metabolic challenge.
Table 1 – Effects of the different ED protocols on metabolic parameters. BW was assessed during the ED protocols and all the rest of the parameters were assessed post recovery. For the ABA group, PNS animals were compared to resilient CTRLs.

| Experiments | BW | Food intake | Metabolic rate | Activity | Adiposity | Glucose tolerance | Insulin tolerance |
|-------------|----|-------------|----------------|----------|-----------|------------------|------------------|
| PNS vs. CTRLs | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |
| Basal response to PNS | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |
| Food restriction | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |
| Activity based anorexia | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |
| Binge eating | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |
| High fat diet | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] | ![arrows] |

Figure 7: Prenatal stress (PNS) induced abnormalities in hypothalamic gene expression in adult females and fetuses. (A) Gene ontology (GO) enrichment analysis based on a hypothalamic gene expression array of the adult female hypothalamus of control (CTRL) and PNS animals. (B) RT-PCR validation shows different gene expression between the groups ($F_{(4,69)} = 9.48, p = 0.001$). (C) STRING pathway analysis linking the affected genes (https://string-db.org). (D) Relative expression of the selected genes in the female fetal hypothalamus shows abnormalities resulting from PNS ($F_{(4,80)} = 17.74, p = 0.001$). Data presented as mean and S.E.M. N = 6–12.
3. DISCUSSION

In the present set of studies, we aimed to address a common confounding variable encountered in studies exploring the effects of early life stress and predisposition to eating and metabolic disorders by using the same stress protocol to test vulnerability to a variety of different EDs and challenges. It is well known that the effect of PNS on the offspring depends on the severity, timing, sex of the offspring and the variables being tested. Thus, early or “first trimester” PNS reduces BW and adiposity in males exposed to HF diet (females not tested) [12], but late gestation stress increases BW, glycemia, and food intake in male mice (females not tested) [48,49] and insulin resistance in rats of both sexes when exposed to HF diet [16,50,51]. Interestingly, late PNS and HF diet increases adiposity in female, but not male rats [52]. Late PNS further increases susceptibility to BE in female mice [15] and to ABA in a subset of passively stress-coping female rats [52]. In contrast, when PNS takes place during the whole gestational period, the females’ innate susceptibility to ABA disappears [23]. In girls, prenatal stress during the whole pregnancy period is associated with an increased overall risk of BN but not AN [4], and this is particularly true for late gestation stress [53]. Chronic stress and psychiatric comorbidity are also associated with the onset of EDs when the stressor takes place in the year preceding or generally close to the onset of the ED [54]. Again, this effect was more robust for BN and BED than for AN, which showed mixed results [55]. In addition, stress alone does not always trigger EDs/metabolic disorders, and it is likely that rather a combination of an innate predisposition pooled with the additive effect of early life trauma is more likely to lead to disease [8,56]. Thus, different underlying processes appear to be involved in the different EDs, and these processes may be differentially affected by early life alterations depending on the developmental window at which they occur. The heterogeneity of the protocols and time windows considered, combined with the fact that the majority of the still limited amount of studies in the field examined only one sex or one ED with the same protocol, makes drawing conclusions about PNS-linked predisposition to EDs rather difficult. However, these findings highlight that maternal stress is associated with risk of EDs in the offspring with a different time window for each particular ED and sex.

The stress protocol used here, unlike the ones cited above, included the whole period of pregnancy in an attempt to mimic a real-life situation in which the mother experiences chronic anxiety. The resulting responses to the ED protocols and metabolic challenges according to the sex were remarkable, with females showing clear programming to diet induced obesity through fat accumulation and decreased metabolic rate, regardless of the metabolic challenges. This was especially visible in the FR experiment, in which the sudden food limitation led to a transitory but sharp weight loss in CTRLs but not in PNS animals, suggesting body weight changes are more tightly regulated in this group (Figure 2B). In contrast to the females, the males’ metabolic rate was adapted to the type of manipulation (decreased when FR and increased when overfed), so they maintained the appropriate levels of fat and even coped more effectively than CTRLs with HF diet. This was interesting given the usual consensus that PNS combined with HF diet increases insulin resistance, overeating and obesity as suggested in some studies [16,57]. This may be again related to the window of intervention, since other studies focusing on different time-windows reported the opposite effect [12,51,58]. Altogether, while late gestation adversities appear to more profoundly affect females [14–16] and early gestation adversities appear to have more profound effects on males, whole pregnancy adversities appear to have strong and opposite effects for each sex, at least regarding metabolic responses to challenge.

The dramatically different coping style PNS females had compared to CTRLs in response to the different ED protocols is likely mediated by hypothalamic adaptations resulting from PNS. These include abnormal levels of Avp, Oxt, Adrb3, Nms, and Nnat; all deeply involved in several aspects of metabolic regulation, combined with a lifelong inflammatory state and structural and/or functional remodeling of the hypothalamic neurons. Avp is robustly expressed in parvocellular cells of the paraventricular nucleus of the hypothalamus (PVN) and has been implicated in the regulation of blood glucose, locomotor activity and food intake through melanocortin dependent feeding pathways [44,59–61]. There is also substantial evidence supporting the role of Oxt, a stress-buffering hormone, in energy homeostasis since Oxt neuron ablation makes mice more sensitive to HF diet-induced obesity due solely to reduced energy expenditure [62]. Oxt circulating levels are also decreased in obese diabetic patients, potentially worsening their phenotype [46]. Nnat, an endoplasmic reticulum proteolipid implicated in intracellular signaling, is primarily concerned with antenatal brain growth and development. In addition, Nnat expression in the hypothalamus is responsive to acute nutrient and leptin signaling, is decreased by fasting/increased in obesity [63] and is associated with acute appetite and energy homeostasis and in metabolic-inflammation [64]. Nms, specifically expressed in the suprachiasmatic nucleus of the hypothalamus, is involved in the regulation of circadian rhythms [65] and feeding behavior. Intracerebroventricular administration of Nms induced phase shifts in the circadian rhythm of locomotor activity and had an anorexigenic effect [66] through increases in proopiomelanocortin mRNA expression in the arcuate nucleus and corticotropin releasing factor in the PVN. Nms is also involved in thermoregulation via Adrb3 [47]. In turn, polymorphisms in the adrenergic receptor genes have been extensively studied for association with metabolic syndrome and diabetes [67] and appear to have a sex-specific effect [68]. In addition, beta-adrenergic receptor stimulation can prevent androgenization of the neonatal brain [69]. This is particularly relevant given that PNS can, through prenatal androgen exposure, masculinize the food intake pattern of female offspring [70]. Thus, Adrb3 down-regulation can enable masculinization, leading to the (CTRL) male-like response that PNS females show in response to the different ED protocols. Finally, the adipogenic response of PNS females to energy deficit (Figure 1P) in the FR experiment may be linked to CORT rises, similarly to those found among obese women. FR (and probably BE and HF) stress-related CORT rises can potentially facilitate fat deposition [71]. Mechanisms underlying this process may include direct central nervous system action on peripheral tissues with high glucocorticoid receptor concentration in regions that are sexually dimorphic, producing multiple systemic endocrine and metabolic effects [19]. Altogether, a combination of all these defects can induce lipid accumulation through lower metabolic rate and CORT, selective masculinization and metabolic inflammation despite similar food consumption, in response to any metabolic challenge.

Finally, it is well known that when it comes to gestational programming, the placenta plays a critical role in fetal development, by carrying out the feto-maternal exchange, barrier, and endocrine functions during pregnancy. Placentas exhibit pronounced sexual dimorphism in response to variations in the maternal environment both in humans [72] and mice [12,73–75]. The sexual dimorphism found here in placental DNA and RNA methylation in response to PNS can induce dramatic differences in morphology and functioning and consequently in fetal the environment. Specifically, dysregulation of the expression of the transmembrane ATP-binding cassette (ABC) transporters in PNS
females can affect the distribution of nutrients and exchange of waste metabolites across the placenta, consequently playing a critical role in regulating immunological responses and lipid trafficking [76]. Abnormal placental cytokine—cytokine receptor interaction further hints at heightened immunological response resulting from PNS. Next, abnormalities in the placental Oxt signaling pathway, which is thought to play a critical role in placental function [77], may in addition be linked to the low levels in Oxt gene expression observed in the stressed fetal and adult hypothalamus. The observed circadian entrainment in PNS placentas, including abnormally high levels of the Period (Per) canonical clock genes, hints at alterations in placental rhythmicity, which is independent from the mother and the fetus [78]. The fetus in turn may be influenced by dynamic circadian signals originating both in the mother and the placenta, adding a further layer of fetal programming resulting from PNS in females, through mediators of maternal—fetal circadian interactions [79]. Finally, ifg is heavily involved in nutrient transport and can regulate the thickness of the membrane connecting the maternal and fetal circulation [23,80]. Thus, high expression of placental ifg can decrease the thickness of the junction, further enabling an increase in transport of glucose and amino acids by the over-expressed glucose transporter Slc2a1 and the Slc7 family of amino acid transporters to the stressed fetuses, all of which can further alter fetal metabolic programming.

Thus, while it is very well documented that maternal stress impacts the placenta causing varied negative effects on anxiety, depression, cognition, memory [81,82], metabolism [40], and inflammation [38,39] in the offspring, these effects are largely sex-specific [81] arguably originating in evolutionary adaptations aimed at helping each sex better adapt in challenging conditions. A frequent interpretation of these findings proposes that early adversity may threaten the males’ viability, culling the weak and creating a surviving cohort of the fittest. While females may be more successful in adjusting to early adversity with a variety of different strategies, this may come at the high price of increased vulnerability to metabolic disease expressed later in development.

In summary, we found that through global sex-specific placental adaptations, the female hypothalamus is programmed to diet induced obesity and not to develop increased vulnerability to EDs, when exposed to chronic PNS. Moreover, our results highlight the complexity and tortuosity of the interaction between PNS and proneness or resistance to different EDs and the importance of using both sexes and applying comparable manipulations during similar time windows to address the early origin of eating and metabolic disease.

4. MATERIALS AND METHODS

4.1. Animal care

ICR (CD1) mice (Harlan Sprague Dawley Inc., Indianapolis, IN) were maintained in a pathogen-free temperature-controlled (22 ± 1 °C) mouse facility on a reverse 12 h light-dark cycle at the Weizmann Institute of Science, according to institutional guidelines. Food (Teklad global, Harlan Sprague Dawley Inc., Indianapolis, IN) and water were given ad libitum. All experimental protocols were approved by the institutional Animal Care and Use Committee of The Weizmann Institute of Science.

4.2. Breeding

Female ICR mice were mated at 11–13 weeks of age. Two or 3 females were housed with 1 male (minimum age 12 weeks) at the beginning of the dark period and were examined for the presence of a vaginal copulation plug at the end of the dark period. Presence of a copulation plug denoted day 0.5 of gestation. After breeding, the females were individually housed.

4.3. Experimental groups

Females with plugs were randomly assigned to the control or PNS treatment. From day 18.5 of gestation, females were checked twice a day for the presence of a litter (9:00—10:00, 17:00—18:00). Newborn litters were housed with 8:00 were designated as born on that day-postpartum day 0 (PPD 0). On PPD 1, pups were counted, and litters were culled to 10 pups (with sex distribution kept as equal as possible in each litter). Litters with less than 7 pups were excluded.

4.4. Prenatal stress (PNS) protocol

From gestation day (GD)1.5 until GD16.5, pregnant females were exposed to a chronic variable mild stress protocol, including 2 short manipulations during the dark phase and a further overnight (ON) manipulation during the light phase, every day. The chosen stressors did not induce pain and did not directly influence maternal food intake and were repeated on a weekly basis. The short stressors included multiple cage changes (every 15 min for 2 h), cage tilt (15° for 2 h), white noise (1 h), water in cage (250 ml) or no bedding for 2 periods of 2 h, immobilization in a tube or elevated platform (50 cm high in a bucket of cold water) for 30 min and swimming in warm water for 15 min. Overnight manipulations included illumination during the dark phase, saturated bedding (with water), novel object in the cage (10 glass marbles) and overcrowding (with 14 females in the home cage).

Using a randomized complete block design to avoid potential litter effects, the offspring were randomly allocated to the following experimental groups:

1) Food restricted (FR) group. Pups were weaned (PPD 21–22) and group housed until PPD30. They were then singly housed for a week and then exposed to FR for 4 consecutive days. The first day, food access was limited to 4 h and the following 3 days, food access was limited to 3 h in the dark phase. The BW and food intake data were published in a previous study [23] as a control for the ABA/resistant groups for CTRLs and PNS separately. Here, we performed a new analysis for a different comparison, i.e. between CTRLs and PNS. 2) Activity based anorexia (ABA) model. ABA was performed as previously described [23]. Briefly, pups were weaned (PPD 21–22), and offspring were group housed until PPD30. On PPD30, mice were placed in a running Wheel (RW) system in order to record the voluntary running pattern (diameter 115 mm, TSE-Systems, Bad Homburg, Germany). The protocol lasted for 11 days during which food intake and BW were measured daily. The first week was defined as a “training period”, during which food and water were given ad libitum. On the first day of the second week, food access was limited to 4 h, and, on the following days, the food access was limited to 3 h. Animals were defined as anorexic in retrospect by cluster analysis according to: 1) Percentage weight loss; 2) Food intake during FR; 3) Food intake recovery (%) during the FR period; 4) Circadian disruption (km ran during the light phase); 5) Days until collapse (become immobile and hypothermic), and 6) Total running distance (km) when FR. Animals that developed ABA were removed from the running wheel cage once they collapsed and were subsequently given food ad libitum and monitored until fully recovered. 3) Binge eating protocol. BE was performed as previously described [15] and consisted of sporadic and limited access (for 2 h a day, 3 times a week during the dark cycle) to Western diet (D12079Bi, Research diets Inc., New Brunswick, NJ, USA) for a period of four weeks, without FR. The protocol was preceded by a 5-day
habitation to the new diet and started on PPD30, the pre-pubertal period equivalent to adolescence in humans.

4) **High fat group.** Three-month-old mice were fed ad libitum HFD (60% of calories; D12492 Research Diets Inc., New Brunswick, NJ, USA) for 3 weeks.

### 4.5. Glucose/insulin tolerance tests

Glucose and insulin tolerance tests (GTT and ITT respectively) were performed in awake animals during the dark phase of the day cycle. For GTT, glucose (2 g/kg of body weight) was injected i.p., and whole venous blood obtained from the tail vein at 0, 15, 30, 60, 90, and 120 min after the injection was measured for glucose using an automatic glucometer (Accu-Check performa, Roche diagnostics GmbH, Mannheim, Germany), following 6 h of fasting. For the Insulin tolerance test (ITT), 0.75 U/kg insulin (Sigma—Aldrich) was injected after 4 h of fasting. Blood was tested with the glucometer at 0, 15, 30, 60, and 90 min after the injection. All animals underwent both tests (GTT first and ITT second) with 2 days recovery in between.

### 4.6. Body composition

Body composition was assessed using Echo-MRI-100™ (Echo Medical Systems, Houston, TX, USA).

### 4.7. Metabolic assessment

Indirect calorimetry and food and water intake were measured using the LabMaster system (TSE-Systems, Bad Homburg, Germany). The LabMaster instrument consists of a combination of sensitive feeding and drinking sensors for automated online measurement. The calorimetry system is an open-circuit system that determines O2 consumption, CO2 production, and respiratory exchange ratio (RER). Data were collected for five consecutive days after 24 h of adaptation for the apparatus.

### 4.8. Maternal behavior

On days 6/7 and 17/18 postpartum, patterns of undisturbed nocturnal maternal behavior were observed during 160 min sessions. Each mother was observed every 15 min, for 1—3 s. This allowed the identification of the ongoing maternal behavior at the observation time. Various maternal and non-maternal behaviors were recorded in every observation. The score was “1” if the behavior occurred and “0” if it did not occur. Maternal Behavior measures were based on existing literature [83] and included both self-grooming, eating, and pup-directed behaviors (nursing, licking/grooming) and activity measures.

### 4.9. Adult hypothalamic and blood collection

Adult animals were sacrificed at the age of 3 months. After decapitation, the hypothalamus was dissected and immediately frozen. Trunk blood was collected in the PNS and CTRL mice (without any ED protocel) and immediately centrifuged, and plasma samples were stored at −80 °C.

### 4.10. Placenta and fetal tissue collection

On GD17.5, pregnant females were anesthetized with an overdose of Ketamine-Xylazine (1:1, 20% in saline) and through cesarean section both the embryos and placentas were excised. Embryos were decapitated; the brain was removed from the skull and placed with the cortex down onto a flat dish to expose the ventral part. Typically, the hypothalamus is easily recognized by its unique density, the entrance of the optic nerve, and its slightly bulb-like anatomy, distinguishing it from the rest of the brain. The entire hypothalamus was then excised using a pointed scalpel and a thin metal spatula and immediately frozen on dry ice and stored at −80 °C. Trunk blood was collected in EDTA coated tubes (MiniCollect, Greiner bio-one, Austria). Placentas were immediately stored at −80 °C. A portion of the embryos’ tail was removed for later DNA extraction and sex genotyping. For analyses comparing between groups, only one fetus per litter was used. These were chosen from the central uterinal location, with one male and one female as adjacent fetuses.

### 4.11. Genotyping

Sex of the embryo was determined by Sry genotyping (forward 5’TCACTGAGACTGCAACCACAG-3’ and reverse 5’TATGAGACTGCCAACCAG-3’) [84]. Amplification product was detected by 2% gel electrophoresis.

### 4.12. RNA extraction and real-time PCR

For placental tissue, purification of total RNA was done using Tri Reagent (MRC Inc., Cincinnati, OH) according to the manufacturer’s recommendations. gDNA was removed using Turbo DNase digestion in solution, followed by heat deactivation (#AM2238, Thermo Fisher Scientific). Total hypothalamic RNA was extracted using the miRNeasy Mini Kit (#217004, Qiagen) in conjunction with on column digestion of gDNA using Turbo DNase (see above). RNA preparations were reverse transcribed to generate cDNA using miScriptR Reverse transcription kit for miRNAs/mRNA (QIAGEN, Hilden, Germany). Quantitative mRNA expression was done using the miRNeasy Mini Kit (#217004, Qiagen) in conjunction with on column digestion of gDNA using Turbo DNase (see above). mRNA preparations were reverse transcribed to generate cDNA using miScriptR Reverse transcription kit for miRNAs/mRNA (QIAGEN, Hilden, Germany). Quantitative mRNA expression was done using the SYBR®Green PCR kit (QIAGEN, Hilden, Germany) (#204057 QIAGEN, Germany) respectively, according to the manufacturer’s guidelines and a StepOnePlus® Real time PCR system (Applied Biosystems, Waltham, MA), using specific primers. Tbp was used as internal control for mRNA in placental samples and Hprt was used as internal control for mRNA in hypothalamic samples.

### 4.13. Global DNA methylation and hydroxy-methylation

DNA was extracted using Tri Reagent (MRC Inc., Cincinnati, OH) according to the manufacturer’s recommendations. Global methylation of the placenta was measured using the MethylFlash Methylated DNA Quantification ELISA kit (Epigentek Group Inc., Farmingdale, NY) according to the manufacturer’s instructions. Global hydroxymethyl methylation was determined using the MethylFlash Hydroxymethylated DNA Quantification ELISA kit (Epigentek) according to the manufacturer’s instructions.

### 4.14. Global RNA N6-methyladenosine (m6A) methylation

Global RNA methylation of the placenta was assessed using the Epiquik m6A RNA Methylation Quantification ELISA kit (Epigentek Group Inc., Farmingdale, NY) according to the manufacturer’s instructions.

### 4.15. Hypothalamic whole genome array

Whole genome transcriptome of the hypothalamus was analyzed using Agilent’s microarray kit SurePrint G3 (G4859A). The bioinformatic analysis was performed using the Limma (Linear Models for MicroArray Data) package, which is part of the Bioconductor software. We corrected the background with the Normexp method and used the Loess method for the within arrays normalization and the Quantile for between arrays normalization. A linear model was applied in order to find the differentially expressed genes (a simple Bayesian model). Gene ontology analysis was performed using the STRING database [43].
4.16. RT-PCR primers

| Gene   | Forward | Reverse |
|--------|---------|---------|
| Dnmt3a | AGTGACCAACCAAGTCTCACG | GTTCACCTCGGTCTTCTCCAA |
| Dnmt3b | TTCAGTGACCAGGTCTTCGACAGCAGA | TCAGAAAGGTCGGAGACCTTTCTCT |
| Dnmt1  | CCTAGTTCCGGTTGCTAGGAGGAA | TCTCTTCCCTGCTCGAGCCGACTA |
| Tet1   | ATCGAGGGTTAAGGTCTGCTCC | TGCGGTTTTCCTCCTTC |
| Tet2   | GACTGTGGACAAAGAGGAAG | CGGAGGCTTGGTCTGCTAAT |
| Tet3   | CCAATGGTGTTAGTCCGGAAG | ATCGGCGGTCTGAGAACATG |
| Wtop   | AAGCCGACACAGCCAGGAGT | AGTTCTGATTTGAGTGTG |
| KIAA1428 | AGTAACTGGACACCGGCTCAAG | AGTACTGTCAGAGAACCCG |
| FTO    | GCGGGAGAAGCTAGAAACCTAG | AGGGATTTGACCTGCCCT |
| Alkbh5 | TGCTCACTAGTGGTAGCTGCT | CGGGTGACATCAATCTTG |
| Metl3  | GACCTAGGATGTCGGACAGCAGA | CGTTCTTAATCACAAGTGCCC |
| Metl14 | GATGAGATTGCAAGACCTCGAG | CCCATTTGCAAGCATACCT |
| Tbp    | CTACGTTACACGGTGGCAACAG | ACCAACACTACCAACAGCA |
| Avp    | TCAACACTACGCTCCCTGCTT | CCTTGGGCGCGGG |
| Nms    | TCCGGAGCGCTGATAGTTG | TCCTGTTTTCCTAGTGG |
| Oxt    | TCACCTACAGGCTGATCCACAC | AGACACTGCGCATATCCAGG |
| Adrb3  | GCGCAATCGTGGTGTACATCT | GCATTACGAGAGCCTCAC |
| Nnat   | GTCCCGTGTGCTGCTGCTGC | TGTCGGTGCCTGCTTG |
| Hprt   | GACCTGACACCCAAAATGG | GTCCCTTTCACCGAAGCCT |

4.17. Libraries and RNA-Seq

Libraries were prepared with the Illumina TruSeq Stranded Total RNA Library Preparation kit with Ribo Zero Gold (Illumina, #RS-122-2301) according to the instructions, using 1000 ng total mouse placenta RNA as starting material. Libraries were quantified on a Qubit fluorometer and by qPCR with a KAPA Library Quantification Kit for Illumina libraries (# KK4828). Size distribution was checked using the Agilent High Sensitivity DNA Assay (#5067-4626) on an Agilent Bioanalyzer. Samples were denatured using 1N NaOH, diluted to a concentration of 3 nM with ExAMP mastermix and loaded onto a HiSeq 4000 machine (Illumina, San Diego, CA; #SY-401-4001) with 1% PhiX control (Illumina, #FC-110-3001) spiked in. HiSeq 3000/4000 flow cells and HiSeq 3000/4000 SBS sequencing chemistry were used for paired-end sequencing with a read length of 100 bp for each direction. Sequencing was performed at the Helmholtz Center (Munich, Germany).

4.18. RNA-seq analysis

The quality of sequencing reads was verified using FastQC 0.11.5 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc). Adaptors were trimmed using cutadapt v.1.9.1 [85] in paired-end mode. For quantification of gene expression, kallisto 0.43.1 [86] was employed using the mouse Ensembl annotation v79 (downloaded from http://bio.math.berkeley.edu/kallisto/transcriptomes/). Normalization and differential analysis on gene level was performed using DESeq2 version 1.18.1 [87]. RIN value was corrected for. Only genes with minimum of 5 counts in at least 3 samples were included in the analysis; rRNA and genes located on chrM were filtered out. For pathway analysis, the results from the differential analysis were ranked and inputted into FGSEA [88], using KEGG database [89], 100000 permutations and a q-value cutoff of 0.1 for significance. For visualization purpose, R packages were used (edgeR [90], gage [91]).

4.19. Statistical approach

Statistical analyses were performed using Statistical Package for the Social Sciences (SPSS) software, Version 20.0 (SPSS Inc., Chicago, IL) and GraphPad Prism 6 (GraphPad software, Inc., La Jolla, CA). Tests included repeated measures ANOVA, t tests or one-way ANOVA when relevant. Differences between the groups were assessed using Tukey’s multiple comparisons post hocs. When appropriate, non-parametric tests such as Kruskal–Wallis were used. Specific details of N and tests used are provided in the figure legends.

AUTHOR CONTRIBUTIONS

MS designed the experiments, analyzed the data and wrote the manuscript. AC supervised the project. MS, TP, and YD performed the manipulations, RNA and DNA extractions. MS, TP, and YD sacrificed the animals. MJ constructed the libraries and performed NGS. SR and SB performed the bioinformatic predictions and analysis.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data related to this article can be found at https://doi.org/10.1016/j.molmet.2018.08.005.

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