Hepatic adropin is regulated by estrogen and contributes to adverse metabolic phenotypes in ovariectomized mice

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ABSTRACT

Objective: Menopause is associated with visceral adiposity, hepatic steatosis and increased risk for cardiovascular disease. As estrogen replacement therapy is not suitable for all postmenopausal women, a need for alternative therapeutics and biomarkers has emerged.

Methods: 9-week-old C57BL/6 J female mice were subjected to ovariectomy (OVX) or SHAM surgery (n = 10 per group), fed a standard diet and sacrificed 6- & 12 weeks post-surgery.

Results: Increased weight gain, hepatic triglyceride content and changes in hepatic gene expression of Cyp17a1, Rgs16, Fitm1 as well as Il18, Rares2, Retn, Rbp4 in mesenteric visceral adipose tissue (VAT) were observed in OVX vs. SHAM. Liver RNA-sequencing 6-weeks post-surgery revealed changes in genes and microRNAs involved in fat metabolism in OVX vs. SHAM mice. Energy Homeostasis Associated gene (Enho) coding for the hepatokine adropin was significantly reduced in OVX mice livers and strongly inversely correlated with weight gain (r = −0.7 p < 0.001) and liver triglyceride content (r = −0.4, p = 0.04), with a similar trend for serum adropin. In vitro, Enho expression was tripled by 17β-estradiol in BNL 1 ME liver cells with increased adropin in supernatant. Analysis of open-access datasets revealed increased hepatic Enho expression in estrogen treated OVX mice and estrogen dependent ERα binding to Enho. Treatment of 5-month-old OVX mice with Adropin (i.p. 450 nmol/kg/ twice daily, n = 4,5 per group) for 6-weeks reversed adverse adipokine gene expression signature in VAT, with a trended increase in lean body mass and decreased liver TG content with upregulation of Rgs16.

Conclusions: OVX is sufficient to induce deranged metabolism in adult female mice. Hepatic adropin is regulated by estrogen, negatively correlated with adverse OVX-induced metabolic phenotypes, which were partially reversed with adropin treatment. Adropin should be further explored as a potential therapeutic target and biomarker for menopause-related metabolic derangement.

Keywords: Menopause; OVX; Fatty Liver; Transcriptome; Estrogen; Adropin

1. INTRODUCTION

Menopause defined as the permanent cessation of menstruation resulting from the loss of ovarian follicular activity [1], is associated with a deranged metabolic phenotype including weight gain, visceral adiposity, liver steatosis and insulin resistance resulting in a significantly increased risk of developing type 2 diabetes (T2D) and cardiovascular disease (CVD) in post-versus pre-menopausal women [2]. Weight gain results primarily from decreased energy expenditure rather than increased food intake [3,4]. There is a shift in adipose tissue distribution from a lower body predominant subcutaneous fat depot towards a more visceral abdominal distribution pattern [5,6]. Increased visceral adiposity is associated with adipocyte hypertrophy and secretion of adipokines and pro-inflammatory cytokines implicated in insulin resistance and CVD pathogenesis [7–9]. Fat oxidative capacity in the liver and skeletal muscle is reduced resulting in excessive lipid accumulation and increased insulin resistance [10]. As a major metabolic organ as well as a target of the metabolic syndrome, the liver appears to play an important role in mediation of metabolic changes associated with menopause. Known direct liver involvement

Abbreviation: OVX, ovariectomy

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includes hepatic steatosis [11], inflammation [12], decreased fatty acid oxidation [13], and increased hepatic glucose production [14]. Studies in women with natural or iatrogenic premature cessation of ovarian function have shown that estrogen depletion and the disruption of estrogen signaling are key factors in menopause-related metabolic dysfunction [15]. However, the benefits and harms of estrogen replacement therapy (ERT) are still a matter of controversy and ERT is not suitable for all women [16].

Historically, recommendations for prevention and treatment of cardiometabolic derangement in postmenopausal women were derived from studies conducted primarily in men. Moreover, pre-clinical biomedical research in animals did not have equal representation of females, reportedly due to the higher susceptibility of males to high fat diet (HFD)-induced metabolic dysfunction [17]. However, a growing body of evidence shows clear sexual dimorphism in metabolic homeostasis with distinct sex- and gender-specific cardio-metabolic phenotypes [6,18,19]. In addition to sex-hormones, there are sex differences in gene expression and networks that impact sex-specific metabolic phenotypes [20,21]. With a mean age at menopause of 50 and life expectancy over 90 years, women in developed countries are expected to spend nearly half their lives in the post-menopausal state [22]. There is an urgent need to discover new biomarkers and therapeutic targets to alleviate metabolic derangement in this rapidly growing population.

Using RNA-sequencing from liver tissue of a mouse model for human menopause (ovariectomy, OVX), we revealed new players in OVX-induced metabolic abnormalities. To separate the effects of sex hormone deficiency from those of aging and obesity, studies were conducted in young adult female mice fed a standard diet.

2. RESULTS

2.1. OVX induced weight gain, pro-inflammatory and adverse adipokine gene expression in VAT and increased hepatic TG content and related genes

The study design is depicted in Figure 1A. Baseline body weight did not differ between SHAM and OVX mice (Figure 1B), and uterine weight was significantly lower in OVX mice, confirming successful OVX-induced suppression of endogenous estrogen production (Figure 1C). Weight gain in OVX mice was significantly higher than in SHAM mice over the course of 12 weeks post-surgery and approached forty percent from baseline (Figure 1D). A significant increase in gene expression of pro-inflammatory cytokines and adipokines was found in VAT of OVX compared to SHAM mice 6-weeks post-surgery.
2.3. OVX repressed hepatic expression of Enho coding for adropin
We were particularly intrigued by the significant reduction in hepatic expression of the Energy Homeostasis Associated (Enho) gene, observed in OVX compared to SHAM mice 6- and 12-weeks post-surgery (Figure 3A). Enho relative expression was significantly inversely correlated with hepatic triglyceride content 12-weeks post-surgery (Figure 3B) with a similar trend within the OVX group alone (r = −0.58, P = 0.07). Interestingly, we found a positive correlation between uterine weight, a reflection of the level of estrogen and hepatic Enho expression (Figure 3C). Relative weight gain in OVX and SHAM mice was observed in 6 weeks post-surgery also showed a strong inverse correlation with Enho liver mRNA expression (Figure 3D), though driven only by the inter-group differences. Enho codes for the peptide adropin, a highly conserved hepatokine shown to regulate whole body energy homeostasis [26]. Despite the change in mRNA levels, no differences in hepatic adropin protein level in T6/OVX vs. T6/SHAM or T12/OVX vs. T12/SHAM mice could be detected using ELISA (Figure 3E). This pattern is consistent with previous reports [27], and may be explained by adropin being a secreted peptide. Indeed, we found a trend for decreased plasma adropin 28 days after OVX which was negatively correlated with weight gain (Figure 3F). Reduced liver Enho expression and low serum adropin have been previously reported in the context of obesity [28], but have never been investigated in the context of estrogen deficiency, OVX, menopause or disruption of estrogen signaling.

To understand if Enho expression is directly regulated by E2 and not merely a byproduct of changes in body weight or liver fat, we conducted in vitro studies in BNL 1 ME murine hepatocytes. We found a 3-fold increase in Enho expression in response to E2 administration with an increase in adropin levels in the supernatant (Figure 4A,B). To further validate our findings of direct regulation of hepatic Enho expression by E2, we analyzed relevant publicly available datasets from the Gene Expression Omnibus (GEO). In human cells, E2-treated MCF-7 breast cancer cells showed a significant increase in ENHO (Figure 4C). In primary human hepatocytes higher ENHO expression was found in cells obtained from females compared to males (Figure 4D), compatible with the sexually dimorphic expression pattern observed in mice [28]. Consistent with our own experimental data, we found decreased liver Enho expression in OVX vs SHAM mice both on chow or HFD (Figure 4E) and a marked increase in Enho expression following E2 treatment in OVX mice on HFD (Figure 4F).

To gain insight into the potential mechanisms of estrogen regulation of Enho we analyzed data from liver ERz knockout (KO) and ERz ChIP-Seq experiments. In mouse livers, Enho expression was reduced in ERz KO vs. WT mice (Figure 5A) with a significant peak for ERz binding at Enho following E2 treatment (Figure 5B). We found a significant ERz binding peak in the promotor region of ENHO in pooled human livers (three males and three female, age 44–73 years, Figure 5C). We could not detect an estrogen responsive element (ERE) near the Enho promoter. Enrichment analysis on Enho co-expressed genes in our RNA-seq data using CheA [29] revealed enrichment for targets of retinoid X receptor (RXR) (data not shown), known to form a heterodimer with liver X receptor α (LXR). Notably, LXR was previously reported to negatively regulate adropin in vitro and in vivo [26].

2.4. Adropin treatment partially reversed OVX induced adverse metabolic changes
To test the hypothesis that reduced liver adropin expression plays a causative role in OVX induced metabolic derangement, we performed a small proof of concept intervention study using previously reported therapeutic doses of adropin, administered for 6-weeks following OVX. As expected, adropin treated mice showed a marked increase in adropin serum level (Figure 6A). Surprisingly, neither vehicle nor adropin treated OVX mice gained total body weight relative to baseline (Figure 6B), possibly due to the significant stress of a prolonged twice daily injection regimen. Nonetheless, the adropin treated mice exhibited a trend towards increased lean mass (Figure 6C) and reduced liver fat (Figure 6D–F) with increased expression of Rgs16 and reduced expression of the fat transporter cluster of differentiation 36 (CD36) (Figure 6G). Importantly, the adipose adipokine pattern in VAT seen with OVX vs SHAM was partially reversed with a significant reduction in gene expression of Il1b and Retn and trend for Il18, Rars2, Rbp4 in adropin vs. vehicle treated OVX mice (Figure 6H). Steraryl-CoA desaturase-1 (Scd1), a key fat storage gene was downregulated in both visceral and subcutaneous adipose tissues in adropin treated mice (Figure 6G,H). Scd1 was previously shown to be downregulated by adropin in the liver [26]. We found no significant change in blood glucose levels including following a glucose tolerance test (GTT) in adropin vs vehicle treated mice (Figure 5J).

2.5. OVX-induced changes in miRNAs involved in fatty acid metabolism
As miRNAs play a role in post-transcriptional regulation of mRNA, we performed miRNA-seq analysis on the same samples used for the RNA-seq. The relative abundance of specific miRNAs is presented in Figure 5A. We found 48 miRNAs to be differentially expressed in OVX vs. SHAM mice, with 24-up and 24-down regulated (Figure 7A). Samples clustered by gonadal status (Figure 7B). Pathway enrichment analysis of the predicted targets of the differently expressed
Figure 2: OVX induced liver transcriptional changes 6-weeks post-surgery. (A). Principal component analysis based on top 1000 variably expressed genes. (B) Volcano plot: DEGs (differentially expressed genes) marked in green (fold change $\geq 12$, FDR $<0.05$). (C) Heatmap with hierarchical clustering of DEGs. (D) Top upregulated gene sets by Gene Set Enrichment Analysis by normalized enrichment score (NES) and FDR. (E) Gene expression of key DEGs in liver in SHAM and OVX mice 6-and-12 weeks post-surgery determined by qPCR normalized to geometric mean of Gapdh, actB and Polr2a. Results are Mean $\pm$ SEM. Data analyzed by unpaired two-sided t-test; *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001 vs. SHAM. (A–E) n = 10 mice/group.
miRNAs revealed enrichment for the KEGG pathways fatty acid biosynthesis and fatty acid metabolism (Figure 7C). We used negative co-expression analysis between previously described miRNA-mRNA pairs to create a miRNA-mRNA network (Figure S4B). 3 of the miRNAs with the most statistically significant differential expression were chosen for validation using RT-PCR (Figure 7D). Liver expression of miRNA-802 and miRNA-455 has been reported to be sexually dimorphic in young adult mice with higher miR-802 and
Figure 4: *Enho* expression is regulated by estrogen status. (A) *Enho* mRNA in vehicle- or E2-treated (10 nM) BNL 1 ME cells at 24 h determined by qPCR normalized to geometric mean of *Gapdh*, *actB* and *Polr2a* expression; Adropin level measured in cell supernatant (n = 9 replicates/group) (B) *ENHO* mRNA in MCF7 cell line with or without E2 (GPL4133 at GSE42619, n = 3 replicates/group). (C) *ENHO* mRNA in primary human hepatocytes (GSE17251; data obtained from 2 female and 4 males at 2 time points). (D) *Enho* mRNA in liver from OVX vs SHAM mice (RNA-Seq, GSE12947 n = 4 replicates per group) (E) *Enho* mRNA in liver from OVX mice on HFD treated with VEH (43% DMSO, 15% ethanol, and 42% saline) vs E2, (RNA-seq, GSE92968 n = 3 replicates/group). Results are Mean ± SEM. Data analyzed by unpaired two-sided t-test; *P < 0.05; **P < 0.01; vs. vehicle (A-B, E); vs. male (C); vs. SHAM (D).
lower miR-455 expression in females vs. males [30]. It seems plausible that the opposite expression pattern we observed in OVX vs. SHAM results from the loss of female sex hormones. miRNA-200a is known to be over-expressed in nonalcoholic fatty liver disease and was previously shown to be up-regulated in OVX mice [31]. Of note, miRNA-29e3p family has been previously reported to regulate hepatic Enho expression in-vivo in insulin resistant male mice [32], but we found no significant correlation between the two transcripts in OVX and/or SHAM mice (Figure S4A).

3. DISCUSSION

This study demonstrates that OVX without additional insults is sufficient to induce a deranged metabolic phenotype, characteristic of the post-menopausal state including weight gain, increased liver fat and increased gene expression of pro-inflammatory cytokines and unfavorable adipokines in visceral abdominal adipose tissue. While OVX in rodents is often used to study human menopause-related metabolic dysfunction [33], most studies have used older-age, high-fat diet or genetic interventions in addition to OVX to accentuate the deranged phenotype. Our experimental design highlights the net effects of sex-hormone deficiency.

OVX led to increased gene expression of Il18, Rares, Retn and RBP4 in mesenteric VAT. Il18 is a pro-inflammatory cytokine from the Il1 family that is increased in visceral adipocytes of obese humans [34] and involved in accelerated atherosclerosis [35]. Increased serum IL18 level was previously reported in post-compared to pre-menopausal women [36]. Il18 has been shown to be down-regulated by estrogen in multiple tissues [37]. Rares2 codes for the adipokine chimerin, a regulator of adipocyte development [38] specific to mesenteric fat [39]. Lower serum chimerin level was reported in pre-compared to post-menopausal women [40]. Retn codes for the adipokine resistin, a driver of insulin resistance in adipose tissue [41] by inducing pro-inflammatory cytokines production from
Figure 6: Adropin treatment partially reversed OVX induced metabolic derangement. (A) Serum adropin determined upon sacrifice in adropin and vehicle treated OVX mice 2 h after the last dose. (B) Percent body weight change. (C) Body composition at 4 weeks, p-value for change in lean mass from ANCOVA with total body weight. (D) Representative slides from liver stained with oil-red-o and quantification of lipid droplet area. (E) Liver triglyceride (TG) content. (F) Gene expression of fat metabolism genes in liver. (G) Gene expression of pro-inflammatory cytokines and adipokines in VAT and SCAT; (F,G,H) determined by qPCR normalized to geometric mean of Gapdh, actB and Polr2a. n = 4 mice/group; Results are Mean ± SEM; Data analyzed by unpaired one-sided t-test; *P < 0.05; **P < 0.01 vs. vehicle.
Macrophages residing in adipose tissue [42]. Resistin expression in VAT was shown to be downregulated by estrogen in-vitro and in-vivo [43], and increased serum resistin was observed in post-menopausal women with the metabolic syndrome compared to those without [44]. Rbp4 induces inflammation and increased insulin resistance in mice [45]. In humans RBP4 expression is downregulated by estrogen in-vivo [46] and exhibits a sexually dimorphic expression pattern in adipose tissue [47].

The liver is a target of the metabolic syndrome but also may play a causative role in menopause-related metabolic derangement. Thus, we focused on transcriptional changes in the liver induced by OVX that occurred prior to the changes observed in liver fat. To our knowledge,
This is the first report of the OVX-induced transcriptional landscape, integrating data from same-sample RNA-seq and miRNA-seq, thus minimizing the effects of sample handling on the data and allowing for establishing co-expression networks. As expected, we discovered significant changes, primarily in gene sets and miRNAs involved in lipid and fatty acid metabolism.

In the liver, OVX induced expression of several genes known to regulate liver fat content: Fasn1 is a target for Peroxisome proliferator-activated receptor gamma (Pparg), involved in lipid droplet formation and steatohepatitis [48]; Gds2 is a master regulator of lipid and energy metabolism [49] and Rgs16 provides a signaling mechanism for glucose production to inhibit G-protein-coupled receptor-stimulated fatty acid oxidation in hepatocytes [50], with a sexually dimorphic expression pattern [28]. Recently, Rgs16 and Gds2 have been shown to regulate hepatic substrate oxidation as well as inflammation and fibrosis in livers of obese male mice [51].

Of particular interest is our discovery of reduced liver Enho expression in OVX compared to SHAM mice, and its strong inverse correlation with adverse menopause-related clinically important phenotypes, namely weight gain and hepatic steatosis. Enho codes for adropin, a conserved 76 amino-acid peptide, highly expressed in the brain. In the liver, adropin is considered a hepatokine, with amino acids 1–33 serving as signal peptide and Adropin34–76 as the secreted portion [26]. Adropin has both paracrine effects on hepatocytes as well as endocrine effects on muscle and adipocytes. The cell surface receptor for adropin has not yet been characterized, but some studies suggest its effects are mediated via GPR19 [52,53]. Adropin has been implicated in the regulation of glucose and lipid homeostasis mostly in studies conducted in male mice. Adropin-over-expressing male and female mice display milder hepatosteatosis and insulin resistance compared to WT mice when fed HFD [26]. Consistently, male and female adropin knockout mice exhibit hepatic steatosis, increased serum TG, insulin resistance and increased relative fat mass [54]. Importantly, numerous studies in male mice have investigated the therapeutic potential of Adropin in ameliorating HFD-induced metabolic derangement and have demonstrated beneficial effects in key metabolic organs. Systemic administration of Adropin34–76 to male WT mice on HFD attenuated liver steatosis and hyperinsulinemia [26], suppressed hepatic gluconeogenesis in vivo and in vitro [55] and decreased liver fatty acid uptake via downregulation of Cd36 [56], consistent with our results. In another study [57], adropin treatment ameliorated NASH progression by up regulating the expression of Glutamate-Cysteine Ligase Catalytic and Modifier Subunits (Gclc, Gclm) as well as Glutathione Peroxidase 1 (Gpx1) via the factor Nuclear factor erythroid 2-related factor 2 (Nrf2). Other metabolically beneficial effects of adropin have been reported in a wide range of conditions of cardiometabolic dysregulation [61]. Specific polymorphisms in the Enho gene in humans have been described leading to lower serum adropin levels and an increased severity of glucose homeostasis impairment and fat metabolism disorders [62]. In women, lower levels of serum adropin were observed in women with polycystic ovary syndrome, gestational diabetes and endometrial cancer [63–65]. Interestingly, unlike in men, pre-menopausal women exhibited no association between serum adropin and BMI [54] or LDL cholesterol [66]; Though serum adropin was reported to be affected by changes in dietary fat content in young women [67]. To our knowledge, thus far, only one brief related serum adropin to menopausal status, where a differential response to meal composition on serum adropin was observed in pre but not post-menopausal women [68].

Known regulators of hepatic adropin are feeding state, nutrients and clock signals [60]. In mice, a short term (2 day) high-fat-diet leads to increased hepatic Enho expression, whereas a prolonged HFD (1–3 months) as well as in genetic mouse models of obesity caused by melanocortin receptor or leptin deficiency, liver Enho is downregulated [26,66]. In HepG2 cells, adding cholesterol to the culture was shown to downregulate ENHO. Treatment with an LXR agonist (GW3965) downregulated ENHO in HepG2 cells, while antisense RNA targeting LXRa blocked this effect [26,65]. Moreover, lean, chow-fed, male B6 mice treated with an i.v. GW3965 injection exhibited a rapid reduction in hepatic Enho expression 4 h after injection [26,66]. Some evidence also points to Enho regulation by the circadian clock; Treatment with a Rev-erb agonist was shown to suppress hepatic Enho in vitro and in vivo while the RORα/γ inverse agonist (SR1001) suppressed Enho in vitro [26].

To our knowledge this is the first report of Enho direct regulation by sex-hormones and its association with OVX-related metabolic derangement. Our findings of Enho regulation by estrogen are supported by data we extracted and analyzed from available in vivo and in vitro studies in mice and humans. Results from these analyses showed increased hepatic Enho expression in estrogen-treated vs. untreated OVX HFD mice, upregulation of ENHO in response to 17β-estradiol in a human breast cancer cell line, higher expression in hepatocytes from women vs. men and a genomic ERα binding site by ENHO in liver. The mechanisms of hepatic Enho regulation by estrogen are elusive and remain to be investigated. LXRα has been shown to directly downregulate hepatic Enho. As ERα is known to interact with liver LXRα and inhibit its function [69], a plausible mechanism for Enho regulation by estrogen involves ERα repression of LXRα-mediated inhibition of Enho transcription. In a recently available dataset, we found reduced hepatic Enho expression in OVX vs SHAM both in ERα[71] as well as in liver specific ERα knockout (LERKO) mice [70], suggesting additional non-ERα or extra hepatic regulation by estrogen. This is further supported by the reported increase in hepatic Enho expression following E2 treatment to LERKO mice [71].

Importantly, using a small proof-of-concept intervention study, we were able to show that adropin reversed some of the metabolic derangement induced by OVX including a trended increase in lean body, a decrease in VAT gene expression of I118, I11b, Rares, Retn and reduced liver fat with marked upregulation of Rgs16 gene expression. Larger studies with varying doses and regimens of adropin treatment as well as additional metabolic insults such as HFD seem warranted.

This study has several strengths. By using young adult female OVX mice on a normal diet, we avoided the confounding effects of aging and obesity. Our initial unbiased approach using NGS of the hepatic transcriptome enabled the discovery of a new and potentially druggable player in OVX-induced metabolic abnormalities. Same-sample miRNA and miRNA profiling allowed for direct interaction analysis. We employed a broad-spectrum of experimental methodologies spanning from studies in-vivo in the OVX mouse model, in-vitro in a murine hepatocyte cell line model, as well as bioinformatic analyses of our own data and in-silico analysis of relevant open-access data. Importantly, our results may have therapeutic implications, as administration of synthetic adropin[72–74] has been shown to improve multiple metabolic parameters, albeit in mostly in male mice. Indeed, in proof-of-concept intervention study we showed that adropin treatment partially reversed deranged phenotypes induced by OVX.
This study is not without limitations. In vivo studies were performed after an overnight fast. As Enho liver expression is suppressed by fasting, our results may have underestimated the magnitude of difference in Enho and adiponectin between OVX and SHAM mice. Additionally, the intervention study may have been too small and short to show more significant effects; conversely, prolonged twice daily injections may have put too much stress on the mice.

In conclusion, this study indicates that changes in adiponectin liver expression may contribute to OVX-induced metabolic derangement and provides a basis for further intervention studies in OVX animals. In a search for new therapies and biomarkers beyond estrogens to improve women’s health in the postmenopausal phase, the association between perimenopausal serum adiponectin level and the changes in metabolic parameters over time in post-menopausal women warrants further evaluation. Finally, although our study was aimed at searching for new targets to alleviate metabolic dysfunction associated with menopause, estrogens and ERx play a role in hepatic steatosis and glucose metabolism also in males and thus our results may be relevant to both sexes [71].

4. METHODS

4.1. Animals & experimental protocol

Forty 8-week-old C57BL/6 J female mice were purchased from Harlan (Rehovot, Israel), housed in constant temperature rooms with 12-hour light/dark cycles and allowed free access to Harlan Teklad chow (2018S) and water. 9-week-old mice were randomly subdivided into 2 groups of bilateral ovariectomy (OVX) and 2 groups of sham surgery (SHAM), (10 animals/group). To allow metabolic changes to occur a group of each OVX and SHAM animals was sacrificed 6- and 12-weeks post-surgery (T6/OVX, T6/SHAM, T12/OVX, T12/SHAM, respectively, Figure 1A). Body weight was determined weekly and uterine weight was determined upon sacrifice. The results may have underestimated the magnitude of differences; conversely, prolonged twice daily injections may have put too much stress on the mice.

In conclusion, this study indicates that changes in adiponectin liver expression may contribute to OVX-induced metabolic derangement and provides a basis for further intervention studies in OVX animals. In a search for new therapies and biomarkers beyond estrogens to improve women’s health in the postmenopausal phase, the association between perimenopausal serum adiponectin level and the changes in metabolic parameters over time in post-menopausal women warrants further evaluation. Finally, although our study was aimed at searching for new targets to alleviate metabolic dysfunction associated with menopause, estrogens and ERx play a role in hepatic steatosis and glucose metabolism also in males and thus our results may be relevant to both sexes [71].

4.2. Gene and mRNA expression analysis

RNA was extracted from liver and mesenteric VAT using BIO-TRI RNA Bio-Lab) reverse transcribed into cDNA (qScript cDNA Synthesis Kit, Quanta Biosciences) and analyzed with SYBR Green-based quantitative RT-PCR. Relative mRNA expression was determined by the comparative CT method. For each sample, the mean cycle threshold (CT) for each gene (run in triplicate) was normalized to the geometric mean of the mean CT of the 3 reference genes (Gapdh, actB and Polr2a) using the formula: 2^-ΔΔCT for each gene was used to calculate relative gene expression changes between samples. Genes with CT values > 35 were considered not expressed. Supplementary Table 1A provides all the primer sequences used in this study. miRNA-specific cDNA was generated with TaqManTM microRNA Reverse Transcription Kit (4366596, Applied Biosystem). Relative mRNA expression was determined by the comparative CT method and normalized to the geometric mean expression of U6 (TaqMan™ microRNA assay; ID 001973) and Snord1202 (TaqMan™ microRNA).

4.3. RNA-sequencing, gene expression profiling and pathway enrichment analysis

RNA was extracted from liver tissue samples of T6/OVX and T6/SHAM mice with Qiagen miRNeasy Mini Kit (cat. 217004, Qiagen). All RNA samples had an RNA Integrity Number (RIN) of more than 8.1 confirmed by Agilent TapeStation system. RNA libraries were generated with 1ug of RNA as input using the TruSeq RNA Sample Prep kit (Illumina) and poly(A)-enriched according to the TrueSeq protocol. Single-Read 75 reads were sequenced on 4 lanes of an Illumina NextSeq 500. Output was ~ 29 million reads per sample. Poly-A/T stretches, and Illumina adapters were trimmed using cutadapt [73]. Resulting reads shorter than 30bp were discarded. Reads were mapped to the Mus musculus GRCm38 reference genome using STAR [74] supplied with gene annotations downloaded from Ensembl release 88 with EndToEnd option.

In conclusion, this study indicates that changes in adiponectin liver expression may contribute to OVX-induced metabolic derangement and provides a basis for further intervention studies in OVX animals. In a search for new therapies and biomarkers beyond estrogens to improve women’s health in the postmenopausal phase, the association between perimenopausal serum adiponectin level and the changes in metabolic parameters over time in post-menopausal women warrants further evaluation. Finally, although our study was aimed at searching for new targets to alleviate metabolic dysfunction associated with menopause, estrogens and ERx play a role in hepatic steatosis and glucose metabolism also in males and thus our results may be relevant to both sexes [71].

4.4. Micro-RNA sequencing

micro-RNA (miRNA) sequencing was performed on same liver samples used for RNA-Seq. Libraries were prepared using TrueSeq Small RNA. SR60 reads were sequenced on 1 lane of an Illumina HiSeq 2500 (v4). Output was ~ 9 million reads per sample. Low quality bases were trimmed from the reads using Burrows-Wheeler Alignment [78], adapters were trimmed using cutadapt [73]. Reads were mapped to the mouse genome using Bowtie [79]. Resulting reads shorter than 17bp and longer than 25bp were discarded. Reads were collapsed into tags for quantification. BLAT was run on tags against 3 blast databases (hairpin, mature and nt). Tags that were not mature miRNA were filtered out, and remained reads were merged by miRna id. Normalization and differential expression analysis were performed by DESeq2 with the betaPrior, cooksCutoff and independentFiltering parameters set to False. Principle component analysis (PCA) was performed with DESeq2 package. Heatmaps were created with Morpheus (https://software.broadinstitute.org/morpheus). Gene set enrichment analysis (GSEA) was performed using GSEA and MSigDB [77]. Raw P values were adjusted for multiple testing using the procedure of Benjamini and Hochberg’s false discovery rate (FDR q < 0.05).
4.5. **In vitro studies in the BNL 1 ME murine liver epithelial cell line model**

Murine BNL 1 ME liver cells [83] (ATCC©TIB-75) were cultured in Dulbecco’s modified eagle medium (DMEM/10% FBS) with a medium change every 3 days. All experiments were conducted in hormone-free medium (charcoal stripped fetal bovine serum; Biological industries, 04-201-1 A). Cells were plated at density of 9x10⁵ cells/well and treated medium (charcoal stripped fetal bovine serum; Biological industries, 04-201-1 A). Cells were plated at density of 9x10⁵ cells/well and treated with 17β-estradiol (E2) (Calbiochem, #3301) at 10 nM for 24 h. Experiments were performed in triplicates and repeated 3 times. Adropin in cell supernatant was measured using ELISA kit (Phoenix Pharmaceuticals, EK-032-35) according to manufacturer’s instructions.

4.6. **Liver triglyceride content**

Liver triglyceride content was determined by a Triglyceride colorimetric assay (No. 10010303, Cayman Chemical). Triglyceride content was normalized to protein content determined by Bio-Rad Protein Assay (Cat. #500-0006, Bio-Rad).

Liver slides (3 per mouse) were stained using oil-red-o (ab150678, abcam) per manufacturer’s instructions. The entire slides were photographed and scanned on Olympus VS200 and automatically analyzed using ImageJ (1.8.0_172) as previously described for quantification of lipid droplet area [84].

4.7. **Adropin serum measurement**

Serum adropin was measured in duplicates using an EIA kit (Phoenix Pharmaceuticals) as per manufacturer’s instructions.

4.8. **Analysis of open access datasets**

Expression data for OXV vs SHAM mouse liver on chow and HFD were downloaded from GEO GSE112947. Specific experimental methods were previously published [85]. Briefly, sixteen female C57BL/6 J were maintained on a chow (Ralston Purina Company) or HF/HS diet (Research Diets D12266B) at 8 weeks of age until 16 weeks of age. At 6 weeks of age the mice were ovarioctomized or sham operated. Frozen liver tissues were RNA-Seq expression profiling using Illumina HiSeq 2000. We used DESeq2 for differential expression analysis. Expression data for OXV HFD mouse liver with E2 or vehicle treatment were downloaded from GEO GSE92966. The specific experimental methods were previously published [86]. Briefly, female C57BL/6 J mice were ovarioctomized at 8 weeks of age, switched to a high-fat diet (Harlan TD.88137) and divided randomly to vehicle or E2 (0.72 mg, 60-day release E2 pellets, Innovative Research of America) for six weeks till sacrifice. Three liver RNA samples from each treatment group were used for RNA-Seq. Gene expression values quantified from BAM files were calculated using StrandNGS and DESeq normalization. Normalized data was directly available to us for download. We assessed the data for quality control using DESeq package for R. We used unpaired t-test for Enho expression analysis in E2-vs. vehicle-treated mice.

Expression data for mouse ERζKO liver were downloaded from GEO GSE95283. The specific experimental methods were previously published [57]. Briefly, WT and global ERζKO female mice (lacking exon 3 of Esr1) were placed on regular ad-libitum diet (NIH-31, Harlan Laboratory) and sacrificed at age 20 weeks. Liver tissue was harvested for RNA extraction and hybridization on GPL1261 Affymetrix Mouse Genome 430 2.0 Array. We assessed raw data for quality control and log2 transformed using the R packages GEOquery, limma and umap. Probe ID 9947 was used to obtain normalized Enho expression. Unpaired t-test was used to compare Enho expression in between each two pairs of groups.

Expression data in primary human hepatocytes were downloaded from GEO GSE17251. The specific experimental methods were previously published [88]. Briefly, primary hepatocytes were isolated from surgical liver biopsies obtained from six individual donors, 2 females and 4 males. Cells were incubated in fresh medium dissolved in DMSO for 6 and 24 h, followed by RNA isolation and gene expression profiling using Affymetrix Human Genome U133 Plus 2.0 Array. We assessed raw data for quality control and log2 transformed using the R packages GEOquery, limma and umap. Probe ID 228403_at was used to obtain normalized Enho expression. Enho expression in male vs female samples at both time points was compared using an unpaired t-test.

Data for mouse liver ERα ChIP-Seq were downloaded from GEO GSE52351. The specific experimental methods were previously published [90]. Briefly, liver fragments were harvested from 8-week-old female C57/BL6 OXV mice and treated ex vivo with 10 nM E2 or ethanol as vehicle for 45 min in DMEM (n = 5 mice/condition). ChIP-Seq performed with anti-ERα HC-20 antibody (Santa Cruz Biotechnology) and 10 μL of anti-ERα Ab-10 (NeoMarkers). ERα binding peaks were called using Model-based Analysis of ChIP-Seq (MACS) [91]. Data for human liver ERα ChIP-Seq were downloaded from GEO GSE158856. The specific experimental methods were previously published [92]. Briefly, six human liver tissues (three males and three female, age 44–73 yrs) were pooled. ChIP-Seq was performed using a commercial service (Active Motif, Carlsbad, CA, USA) with an anti-ESR1 antibody (sc-543, Santa Cruz Biotech, Dallas, TX, USA). Peaks were called using Homer v4.10 [93]. We used the resulting bedgraph files from both data sets to display significant peaks using the IGV browser for windows (http://software.broadinstitute.org/software/igv/) on top of the mouse (mm9) and human (hg38) genomes, respectively.

4.9. **Statistical analysis**

Statistical significance was calculated using GraphPad Prism for Windows (GraphPad Software, La Jolla California USA). Outliers were automatically removed using the ROUT method with Q = 1%. Unpaired Student’s t-test was used to compare means of 2 groups. To assess differences between multiple means one-way ANOVA followed by Sidak post-hoc correction was performed. Correlation coefficients were calculated with Pearson’s correlation analysis. Differences of P < 0.05 were considered significant.

**AUTHOR CONTRIBUTIONS**

JS & IG – planned and conducted experiments, acquired, and analyzed data, wrote and prepared manuscript; ECK, OY, & NH - conducted experiments; HB - analyzed data; NL, NO & JT conducted experiments and analyzed data. RDP – designed research, wrote and prepared manuscript.
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CONFLICTS OF INTEREST

The authors have declared no conflicts of interest exist.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data to this article can be found online at https://doi.org/10.1016/j.molmet.2022.101482.

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