Maternal high-fat diet induces long-term obesity with sex-dependent metabolic programming of adipocyte differentiation, hypertrophy and dysfunction in the offspring

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Maternal obesity determines obesity and metabolic diseases in the offspring. The white adipose tissue (WAT) orchestrates metabolic pathways, and its dysfunction contributes to metabolic disorders in a sex-dependent manner. Here, we tested if sex differences influence the molecular mechanisms of metabolic programming of WAT in offspring of obese dams. To this end, maternal obesity was induced with high-fat diet (HFD) and the offspring were studied at an early phase [postnatal day 21 (P21)], a late phase (P70) and finally P120. In the early phase we found a sex-independent increase in WAT in offspring of obese dams using magnetic resonance imaging (MRI), which was more pronounced in females than males. While the adipocyte size increased in both sexes, the distribution of WAT differed in males and females. As mechanistic hints, we identified an inflammatory response in females and a senescence-associated reduction in the preadipocyte factor DLK in males. In the late phase, the obese body composition persisted in both sexes, with a partial reversal in females. Moreover, female offspring recovered completely from both the adipocyte hypertrophy and the inflammatory response. These findings were linked to a dysregulation of lipolytic, adipogenic and stemness-related markers as well as AMPKα and Akt signaling. Finally, the sex-dependent metabolic programming persisted with sex-specific differences in adipocyte size until P120. In conclusion, we do not only provide new insights into the molecular mechanisms of sex-dependent metabolic programming of WAT dysfunction, but also highlight the sex-dependent development of low- and high-grade pathogenic obesity.

Introduction

Childhood overweight and obesity are global epidemic threats with a steadily rising incidence and the risk for chronic health problems later on [1–3]. The increasing prevalence of obesity results in a higher incidence of obese pregnant and lactating women. Accumulating evidence demonstrates an intergenerational cycle of obesity, with children of obese women being more susceptible to early-onset obesity. This concept has been coined as metabolic programming [4–6]. Clinical and experimental studies confirm that maternal obesity during a critical window of development determines metabolic programming of organ structure and physiology in the offspring, and induces metabolic disorders beyond infancy [7–10]. For
example, offspring of obese dams exhibit marked accumulation of white adipose tissue (WAT), elevated levels of leptin and inflammatory cytokines, insulin resistance and aggravated metabolic response when exposed to obesogenic diet later in life [10–13]. Understanding the molecular mechanisms of adipose dysfunction after maternal obesity may provide novel avenues to prevent long-term sequelae of metabolic programming.

Obesity is characterized by an accumulation and a dysfunction of WAT that results in an elevated secretion of adipocytokines and hormones. This chronic subacute inflammatory state predisposes to higher risk for metabolic pathologies, including diabetes mellitus and cardiovascular diseases [14,15]. The WAT orchestrates various metabolic pathways and contributes to the metabolic syndrome in a sex-dependent manner [16–19]. The progression of obesity-related disease is often inevitable, and intimately linked to adipocyte dysfunction [20]. In general, WAT dysfunction includes inflammation, senescence, insulin resistance, preadipocyte dysfunction and adipocyte hypertrophy [21]. Interestingly, these changes in adipogenic function do not only trigger obesity-related diseases, but are also hallmarks of ageing [22–24]. However, the molecular mechanisms as well as sex differences in adipogenesis during metabolic programming and WAT function remain elusive.

The timing of adipose tissue development determines the window of vulnerability to metabolic programming. The number of adipocytes is primarily determined early in life and is mostly stable through adulthood [25]. Moreover, adipogenesis and adipocyte function are tightly regulated by the concerted interaction of various growth factors and by the microenvironment, including inflammatory cells, matrix, cytokines, hormones and mechanical stress [26,27]. Disruption of these processes by maternal obesity during development can result in WAT dysfunction with adverse auto- and paracrine as well as systemic effects, ultimately promoting metabolic disorders and diseases [25]. Thus, identification of early-onset molecular mechanisms disrupting adipogenesis and determining adipocyte function in a sex-dependent manner in health and disease may define new targets to prevent or reverse WAT dysfunction and attenuate obesity-related pathologies. Here, we present a comprehensive study, in which we investigated adipogenesis, the senescence-associated secretory phenotype (SASP), inflammatory response and the deposition of WAT up to postnatal day 120 (P120). Our data provide evidence of sex-dependent mechanisms in metabolic programming of WAT and most interestingly, a protection of females from long-term adipocyte hypertrophy.

Materials and methods

Animal procedures

All animal procedures for this manuscript were performed in accordance with the German regulations and legal requirements and were authorized by the local government authorities (LANUV; 2012.A424). All mice (C57BL/6N) were placed in a room with 22 ± 2 °C, were exposed to a light/dark cycle of 12 h each and had ad libitum access to water and their respective chow. The animal model of metabolic programming was performed as previously described [10]. Virgin female mice (C57BL/6N) from our own colony received a high-fat diet (HFD; modified catalog no. C1057; Altromin, Lage, Germany) or a standard diet (Co; ssniff catalog no. R/M-H, V1534-0) for 8 weeks after weaning (P21). HFD and Co dams were time-mated with standard diet-fed male mice and continued on their respective diets throughout gestation and lactation. At birth, the litter size of all dams was normalized to six for each litter. Water and chow were available ad libitum, and food was withdrawn only for experimental reasons. After weaning at P21, the offspring of HFD and Co dams were fed standard diet until P120. Both male and female offspring were studied: Co\textsuperscript{male}, Co\textsuperscript{female}, HFD\textsuperscript{male} and HFD\textsuperscript{female}. The exact number of animals are listed in the figure legends; the animals of each group and time point were obtained from three to four different litters. After killing the mice at P21 and P70, WAT was excised as previously described [28]. In addition, some mice underwent a magnetic resonance imaging (MRI) analysis at P21, P70 or P120. The animal experiments took place in the Laboratory of the the Department of Pediatric and Adolescent Medicine and the Department of Diagnostic and Interventional Radiology, University Hospital Cologne, Cologne, Germany.

MRI-analysis of body composition

A whole body MRI scan was performed to assess total body volume, total fat volume and fat fraction. MRI scans were acquired on a Philips Ingenia 3.0 T system (Philips Healthcare) combined with a commercially available small animal coil (Philips Research Hamburg, Germany) with heating function to preserve body temperature during the examination. All mice underwent inhalational anesthesia with isoflurane (2.0–2.5%) in air during image acquisition. The MR protocol consisted of a 3D mFFE T1-weighted sequence with following parameters: echo time (TE) 3.41 ms, repetition time (TR) 8.13 ms, flip angle 45°, field of view (FOV) = 80 × 32 mm, matrix 256 × 256, slice thickness 0.35 mm, gap 0 mm, acquired voxel size 0.35 × 0.35 mm and reconstructed voxel size 0.17 × 0.17 mm, number of signal averages (NSA) 2 and standard scan time for 183 slices was 2 min. First, the total body volume was quantified
by circling the offspring’s body, starting at the cerebellum and ending when both legs were separated from the ilium. Second, the total fat volume was measured by defining signal-intensity thresholds in order to detect the adipose tissue; adipose tissue has a high signal intensity in T1 when compared with other organs. The intestinal area was excluded to avoid false-positive measurements. Finally, the total fat volume was related to the total body volume to determine the fat fraction.

**Physiological data of the offspring**
The body weight (in grams) was obtained at each time point (P21, P70 and P120). Subsequently, mice were killed and perigonadal (pg), retroperitoneal and subcutaneous WAT were excised and weighed (grams).

**Tissue preparation**
Perigonadal WAT (pgWAT) was excised for molecular studies at P21 and P70. A fraction of the pgWAT was immediately frozen and stored at −80°C for protein analysis and assessment of gene expression. Another fraction was fixed in 4% paraformaldehyde (PFA) in phosphate-buffered saline, followed by paraffin embedding for quantitative histomorphological studies. Finally, a third fraction was used to assess β-galactosidase-positive cells as an indicator of senescence.

**Hematoxylin and Eosin staining**
Five-micrometer cross-sections of the PFA-fixed and paraffin-embedded pgWAT were stained with Hematoxylin and Eosin as previously described [29]. The tissue sections were imaged in a magnification of 20× using the slide scanner (Leica SCN400, Germany). The mean linear intercept (MLI) of adipocytes was measured by using the program ‘cell∧D’ and a grid pattern of 50 μm × 50 μm (version 5.1, Olympus Europe SE & Co. KG, Hamburg, Germany). Blood vessels and connective tissue were avoided. The MLI was assessed in six to eight random fields of view per pgWAT section; three sections per animal.

**Senescence-associated β-galactosidase staining**
Senescence-associated β-galactosidase (SA-β-gal) solution was prepared using 1 mg/ml β-X-Gal (5-Bromo-4-Chloro-3-indolyl β-D-galactopyranoside; #B4252, Sigma–Aldrich, Germany), 5 mmol/l potassium ferrocyanide [potassium hexacyanoferrate(II) trihydrate; #P3289, Sigma–Aldrich, Germany], 5 mmol/l potassium ferricyanide [potassium hexacyanoferrate(III) trihydrate; #244023, Sigma–Aldrich, Germany], 150 mmol/l NaCl, 2 mmol/l MgCl₂, 0.01% SDS and 0.02% Nonidet-40 (NP40 Alternative, #492016, Calbiochem, Germany). After that, pgWAT was incubated in β-galactosidase for 2 h at 38°C. Subsequently, the tissue was fixed with Tissue-Tec (Sakura Finetek™; #4583 Sakura; Thermo Fisher Scientific, Germany) and frozen at −80°C. The frozen WAT was sectioned (20 μm) using Research Cryostat Leica CM3050S and counterstained with Nuclear Fast Red (Hoelzel Biotech; #BOS-AR0008, Germany). The number of SA-β-gal positive cells was assessed in two fields of view per pgWAT section; one section per animal.

**RNA extraction and reverse transcription polymerase chain reaction**
Total RNA was isolated from pgWAT using TRizol (Thermo Fisher Scientific, Germany), and real time reverse transcription polymerase chain reaction (RT-PCR) was performed at P21 and P70 using the 7500 real-time PCR system (Applied Biosystems, Foster City, CA) as described previously [30]. Gene expression was normalized to Hypoxanthine-guanine phosphoribosyltransferase (HPRT) and β-Actin at P21 and P70, respectively. Primers and Taq-Man probes were designed using Primer Express and are listed in Table 1.

**Protein isolation and immunoblotting**
Protein isolation of pgWAT, measurement of protein concentration and immunoblots were performed as described previously [29]. The blots were incubated with the following antibodies: monoclonal rabbit anti-phospho Akt (pAkt; Cell Signaling Technology, #4058; 1:1000); monoclonal rabbit anti-Akt (Cell Signaling Technology, #9272; 1:2000); monoclonal rabbit anti-phospho 5’ adenosine monophosphate-activated protein kinase α (pAMPKα; Thr172; Cell Signaling Technology, #2535; 1:1000); monoclonal rabbit anti-AMPKα (Cell Signaling Technology, #2603; 1:1000); polyclonal rabbit anti-Δ like non-canonical notch ligand [DLK; Preadipocyte factor 1 (Pref1); abcam, #ab21682; 1:2000]; monoclonal rabbit anti-phospho p38 mitogen-activated protein kinase (pp38; Thr180/Tyr182; Cell Signaling Technology, #4511; 1:1000) and polyclonal rabbit anti-p38 (Cell Signaling Technology, #9212; 1:1000). Monoclonal mouse anti-β-Actin (Cell Signaling Technology, #3700; 1:4000) served as a loading control. Anti-mouse IgG,
Table 1 List of primers used for real-time RT-PCR; TaqMan and SYBR-Green primers

| Gene                                    | Method | Primer | Sequence                                |
|-----------------------------------------|--------|--------|-----------------------------------------|
| Leptin (Lep)                            | Taq    | for    | 5’TCAOCAGGATCATAGCATTTCAC’            |
|                                         |        | rev    | 3’AGCCGGAAATGACGGCAG’                  |
|                                         |        | probe  | 5’ACGGATGTCAGCGCCGCG’                  |
| Interleukin 1-β (Il1β)                  | Taq    | for    | 5’TGAAGAGATGACGACTTC’                  |
|                                         |        | rev    | 3’GCAAGCGGCAAGTCGAC’                   |
|                                         |        | taq    | 5’ACCCAAGGACGTCAC’                     |
| Metalloproteinase12 (Mmp12)             | Taq    | for    | 5’CAGCGCTTTCTCTGAGTA’                  |
|                                         |        | rev    | 3’GTACATGCAGGGACCTCAC’                 |
|                                         |        | taq    | 5’GTCGGCAAGTGCACACACTGG’               |
| Cyclin-dependent kinase inhibitor 1b (Cdtn1b) | Taq    | for    | 5’GAGACAGTGTGCAAGGATT’                 |
|                                         |        | rev    | 3’GGGGTTCGTGCATCCAC’                   |
|                                         |        | taq    | 5’ATGCAGCCTTCTGCTTACC’                 |
| Cyclin D1 (Ccndf1)                      | Taq    | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Sirtuin 1 (Sirt1)                       | Taq    | for    | 5’GGCTTTGAGATGTAACGTGG’                |
|                                         |        | rev    | 3’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | taq    | 5’ATGCAGCCTTCTGCTTACC’                 |
| Catenin β 1 (Ctnnb1)                    | Taq    | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Krüppel-like factor 4 (Klf4)            | Taq    | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Sterol regulatory element binding transcription factor 1 (Srebp1a) | Taq    | for    | 5’CATGACTTGGCCGCTGC’                   |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Adiponectin (Adipoq)                    | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Cyclin-dependent kinase inhibitor 1a (Cdkn1a) | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Cyclin-dependent kinase inhibitor 1b (Cdkn1b) | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Cyclin-dependent kinase inhibitor 2a (Cdkn2a) | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Perilipin 1 (Plin1)                     | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Perilipin 2 (Plin2)                     | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Monoacylglycerol lipase (Mgll)          | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |
|                                         |        | taq    | 5’AGGGTCCGCAAGTGGAAA’                  |
| Fatty acid-binding protein 4 (Fabbp4)   | SYBR   | for    | 5’GAGACGTGTCGAGTCAAC’                  |
|                                         |        | rev    | 3’GGGGCAAGGCAAGTGCAC’                  |

HRP-linked (Cell Signaling Technology, #7076, 1:5000) and anti-rabbit IgG, HRP-linked (Cell Signaling Technology, #7074, 1:2000 or 1:1000) were used as secondary antibodies.
Figure 1. Measurement of body weight and relative WAT in the offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 and P70

(A) Postnatal Day 21

(B) Postnatal Day 70

Figure 1. Measurement of body weight and relative WAT in the offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 and P70.

(A) Body weight and total WAT (g) relative to body weight in male and female offspring at P21. (B) Body weight and total WAT (g) relative to body weight in male and female offspring at P70. Total WAT includes pg, retroperitoneal and subcutaneous WAT. Co, black bar; HFD, white bar; n = 4–10 animals per group; mean ± SEM; *Student’s t test and Mann–Whitney test: *P < 0.05; **P < 0.01.

Analysis of data

The results of real-time RT-PCR were calculated based on the $-\Delta\Delta C_t$ method and expressed as fold induction of mRNA expression compared with the housekeeping gene (1.0-fold induction). Values are shown as means ± standard error of the mean (SEM). Mann–Whitney test or Student’s t test were used to test significance at the given time points. A P-value less than 0.05 was considered significant. Densitometric analysis of protein bands was performed using Bio-Rad ImageLab software (Bio-Rad, Munich, Germany). Band intensities from samples were normalized for loading using the β-Actin band from the same sample.

Results

Maternal obesity induces obese body composition with sex-dependent distribution in the offspring

Maternal obesity was induced in virgin female mice by feeding HFD for 8 weeks prior mating; the future control dams (Co) received standard diet. Both groups continued on their respective diets during pregnancy and lactation. The offspring were investigated and divided in four groups: Co male, Co female, HFD male and HFD female. First, we queried if there are sex-dependent differences regarding the impact of maternal obesity on body weight. At P21, both body weight and relative WAT were slightly higher in HFD male when compared with Co male, respectively; in contrast, females did not exhibit differences in body weight, but a mild increase in fat mass after maternal obesity (Figure 1A). At P70, we found the opposing effect: the body weight was rather reduced, whereas the relative WAT was significantly higher in HFD male when compared with Co male. On the contrary, HFD female exhibited a similar body weight like Co female and the relative WAT was not significantly higher (Figure 1B). We next performed MRI studies to measure body volume (ml) and fat fraction (%) in the offspring. Representative MRI images illustrate fat mass and distribution in HFD male and HFD female in comparison with their respective controls Co male and Co female at P21 (Figure 2A) and P70 (Figure 2C). At P21, the body volume was slightly augmented in males after HFD and significantly higher in females. Likewise, the fat fraction was markedly greater in HFD male than Co male; this effect was even more pronounced in HFD female (Figure 2B). At P70, we found no effect of maternal obesity on body volume, but the fat fraction remained significantly higher in males and slightly in females after maternal obesity. However, the fat fraction in females after maternal obesity compared with Co female was less augmented at P70 than at P21, indicating a partial reversal coupled with a physiological increase in WAT in females (Figure 2D). Finally, we studied the long-term effects on body composition and adipocyte size in the offspring. Representative MRI images at P120 are shown in Figure 2E. There were no differences in body volume between Co and HFD in males or females. The fat fraction, however, was higher by almost two-fold in HFD male and HFD female when compared with the sex-matched control (Figure 2F).

We next analyzed WAT distribution by measuring the mass of the different compartments after dissection. At P21, we determined a marked increase in retroperitoneal and pg WAT in HFD male when compared with Co male, whereas in females only pg WAT was higher after maternal obesity (Figure 2G). At P70, we observed a shift in WAT distribution in males with a marked increase in subcutaneous WAT in both HFD males and HFD females when compared with Co male or Co female, respectively (Figure 2H). In conclusion, maternal obesity has a modest impact on body weight in both sexes, but induces a persistent obese body composition with a sex-dependent fat distribution.
Maternal obesity programs adipocyte size in a sex-dependent manner

Having shown that maternal obesity induces an obese body composition and a sex-dependent fat distribution let us study the adipocyte size. Since our previous studies investigated pgWAT and the distribution of pgWAT was markedly different in male and female offspring after maternal obesity in the present study, we performed quantitative histomorphometric analysis using pgWAT. Representative H&E-stained images are displayed in Figure 3A (P21), B (P70) and C (P120). Assessment of MLI as an indicator of adipocyte size showed a significant increase in both HFDmale and HFDfemale when compared with the respective Co group at P21 (Figure 3D). At P70, however, the MLI remained slightly higher in HFDmale ($P=0.07$), but not in HFDfemale (Figure 3E). Finally, we analyzed the size of adipocytes at...
P120 and determined a sex-dependent increase in adipocyte size in the offspring after maternal obesity. HFDmale exhibited a significant greater MLI when compared with Comale. Adipocyte size of females, however, was not altered by HFD (Figure 3F). These data demonstrate a persistent effect of maternal obesity on adipocyte size in males, whereas females recover. In summary, the data show a sex-dependent long-term programming effect of maternal obesity with
a persistent obese body composition in both female and male offspring, whereas adipocyte hypertrophy only persisted in males.

**Sex-dependent metabolic programming of the inflammatory response in pgWAT**

Inflammation is a hallmark of adipose tissue dysfunction and metabolic diseases. Hence, we next analyzed the gene expression of inflammatory mediators in pgWAT in the *early* (P21) and *late* phase (P70). *Early phase* (P21): we found a moderate increase in leptin (*Lep*) mRNA in HFD<sub>male</sub>, but not HFD<sub>female</sub> compared with Co<sub>male</sub> and Co<sub>female</sub>, respectively (Figure 4A). In contrast, adiponectin (*AdipoQ*) mRNA expression was higher in females after maternal obesity, whereas males were not affected (Figure 4B). Interestingly, key inflammatory macrophage markers, such as interleukin 1β (*II1β*) and metalloproteinase 12 (*Mmp12*), were up-regulated in females after maternal obesity when compared with HFD<sub>male</sub> (Figure 4C,D). We next assessed the activation of the inflammatory p38 signaling and found a two-fold greater phosphorylation of p38 (pp38) relative to total p38 in females after maternal obesity than in males (Figure 4E). When p38 was compared with the loading control (β-Actin) we only determined a significantly greater activation in SD<sub>female</sub> than SD<sub>male</sub>, possible due to interindividual variability.

*Late phase* (P70): we found a marked sex-dependent reversal of the early inflammatory and metabolic response of pgWAT. The *Lep* expression was not only lower in HFD<sub>male</sub> when compared with Co<sub>male</sub>, but also in general lower in females than males. *AdipoQ* mRNA was neither significantly regulated by sex nor by maternal obesity (Figure 4F,G). Regarding inflammatory markers, HFD did not affect the expression of *II1β* and *Mmp12* in either sex; however, both genes were significantly lower by up to 80% in females when compared with males (Figure 4H,I). While phosphorylated p38 relative to total p38 was not regulated by maternal obesity, it was lower in Co<sub>female</sub> than in Co<sub>male</sub> (Figure 4J), suggesting a time- and sex-dependent inflammatory expression pattern in adipose tissue.

**Maternal obesity has long-term sex-dependent effects on the cell cycle machinery of adipocytes**

The preceding results demonstrate changes in adipocyte size and the inflammatory expression pattern. We next tested, if these changes are related to altered expression of cell cycle-regulating genes in pgWAT. At the *early phase* (P21; Figure 5A–E) we determined no sex- or HFD-dependent effect on gene expression of cyclin-dependent kinase inhibitor 1a (*Cdkn1a*, P21), cyclin-dependent kinase inhibitor 1b (*Cdkn1b*, P27), cyclin D1 (*Ccnd1*) and NAD-dependent deacetylase sirtuin-1 (*Sirt1*); however, gene expression of cyclin-dependent kinase inhibitor 2a (*Cdkn2a*, P16) was regulated by sex with a significant increase in Co<sub>female</sub> when compared with Co<sub>male</sub>

On the contrary, we determined marked regulation of the aforementioned cell cycle regulating genes at the *late phase* (P70, Figure 5F–I): Cell cycle inhibitory markers (*Cdkn1b* and *Cdkn2a*, *Sirt1*) were significantly reduced in a sex-dependent manner in HFD<sub>female</sub> compared with Co<sub>female</sub>. Gene expression of *Cdkn1a* was neither regulated by sex or maternal obesity. However, the proliferative marker *Ccnd1* was significantly up-regulated in HFD<sub>male</sub> when compared with Co<sub>male</sub> and not affected in females. In summary, these findings demonstrate a time- and sex-dependent expression of genes encoding for cell cycle-regulating markers, favoring proliferation in females, but less in males after maternal obesity.

**Differentiation of adipocytes is time- and sex-dependent after maternal obesity**

Preadipocytes are central in adipogenesis and adipose tissue function. They not only serve as progenitors of adipocytes, but they also contribute to adipose tissue dysfunction by adopting inflammatory macrophage-like or senescence-associated phenotype. Therefore, we assessed differentiation markers, such as Perilipin1 and 2 (*Plin1* and *Plin2*) in pgWAT at P21 and P70. At the *early phase* (P21, Figure 6A), we found that both HFD<sub>female</sub> and Co<sub>female</sub> express more *Plin1* and *Plin2* than HFD<sub>male</sub> and Co<sub>male</sub>, respectively. Interestingly, maternal obesity reduced *Plin2* mRNA significantly in females, but not males when compared with the sex-matched controls. Similarly, gene expression of Krüppel-like factor 4 (*Klf4*; regulator of cell survival and differentiation) was significantly lower in HFD<sub>female</sub> than in HFD<sub>male</sub>. Gene expression of Catenin β 1 (*Ctnnb1*; regulator of stemness) was not regulated by sex or maternal obesity. At the *late phase* (P70, Figure 6C), we found a lower *Plin1* and *Plin2* mRNA after maternal obesity independent of sex. Interestingly, the gene expression of *Klf4* and *Ctnnb1* was significantly lower in HFD<sub>female</sub> when compared with either Co<sub>female</sub> or HFD<sub>male</sub>. To further investigate if the differentiation of preadipocytes is regulated in a sex-specific manner after maternal obesity, we assessed protein abundance of DLK, which inhibits preadipocyte differentiation as well adipogenesis. At P21, but not P70 we found a markedly lower DLK in males, but not females.
Figure 4. Sex-dependent inflammatory response in pgWAT after maternal obesity

Assessment of adipokine expression and p38 signaling in pgWAT after maternal obesity at P21 (A–E) and P70 (F–J). (A–D, F–I) Measurement of gene expression in pgWAT in male and female offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 (A–D) and P70 (F–I) using qRT-PCR: Comale, HFDmale, Cofemale and HFDfemale. Adipokines: leptin (Lep; A,F); adiponectin (AdipoQ; B,G); interleukin 1β (Il1β; C,H) and metalloproteinase 12 (Mmp12; D,I). (E,J) Immunoblot showing phosphorylated p38 (pp38) and total p38 in pgWAT of Comale, HFDmale, Cofemale and HFDfemale at P21 (E) and P70 (J); β-Actin served as loading control. The quantitative densitometric summary is shown next to respective immunoblot; pp38 was related to total p38 or β-Actin; total p38 was related to β-Actin. Co, black bar; HFD, white bar; n=4–8 animals per group; mean ± SEM; *Mann–Whitney test: *P < 0.05; **P < 0.01.
Figure 5. Sex-dependent regulation of cell cycle-regulating genes in pgWAT
Assessment of the expression of genes encoding cell cycle regulators in pgWAT from male and female offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 (A–E) and P70 (F–J) using qRT-PCR: Co-male, HFD-male, Co-female and HFD-female. Genes: Cdkn1a (p21; A,F); Cdkn1b (p27; B,G); Cdkn2a (p16; C,H); Ccnd1 (D,I); NAD-dependent deacetylase Sirt1 (E,J). Co, black bar; HFD, white bar; n=4–6 animals per group; mean ± SEM; *Mann–Whitney test: *P<0.05; **P<0.01.

Maternal obesity regulates AMPKα and Akt signaling in a sex-dependent manner
Energy metabolism and differentiation of adipocytes are important in the maintenance of adipose tissue function. Since both AMPKα and Akt signaling are central in these processes and regulated by cytokines and hormones, we studied their activation in pgWAT using immunoblots. Early phase (P21; Figure 7A,C): first, we assessed phosphorylated AMPKα (pAMPKα) relative to total AMPKα and found lower activation in Co-female when compared with Co-males. Maternal obesity, however, increased pAMPKα in males significantly and in females slightly (P=0.06). Second, we measured phosphorylated Akt (pAkt) and found sex-dependent differences with a greater activation in HFD-females than in HFD-males. At the late phase (P70; Figure 7B,D) we determined opposing effects: pAMPKα was significantly lower in Co-females than in Co-males. Maternal obesity only reduced AMPKα signaling in males, but did not have any effect in females. On the other hand, pAkt was greater in HFD-males when compared with HFD-females. In summary, maternal obesity regulates AMPKα and Akt signaling in a time- and sex-dependent manner. At the early phase, AMPKα and Akt are markedly activated in males and females, respectively; whereas this shift at the late phase, possibly reflecting a sex-dependent imbalance in adipokines and insulin sensitivity.

Maternal obesity regulates adipocyte metabolism in a sex-dependent manner
To determine if adipocyte metabolism is differently regulated in the long-term in males and females after maternal obesity we assessed gene expression of monoacylglycerol lipase (Mglil), fatty acid-binding protein 4 (Fabp4) and sterol regulatory element-binding protein (Srebp1) in pgWAT at P70. We found that Mglil (Figure 8A) and Fabp4 (Figure 8B) were significantly lower in HFD-females when compared with Co-females. This effect was not detected in male offspring...
Figure 6. Maternal obesity regulates the expression of differentiation markers of adipocytes in pgWAT in a time- and sex-dependent manner.

(A,C) Measurement of the expression of genes encoding regulators of adipocyte differentiation and stemness in pgWAT from male and female offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 (A) and P70 (C) using qRT-PCR: Plin1; Plin2; Klf4; Ctnnb1. Genes: Plin1; Plin2; Klf4; Ctnnb1. (B,D) Immunoblot for DLK (Pref1) as an adipocyte differentiation marker in WAT at P21 (B) and P70 (D); β-Actin served as loading control. The quantitative densitometric summary is shown below to respective immunoblot; DLK was related to β-Actin. Co, black bar; HFD, white bar; n=4-6 animals per group; mean ± SEM; *Mann–Whitney test: *P<0.05; **P<0.01.
Figure 7. Maternal obesity regulates AMPKa and Akt signaling in pgWAT in a time- and sex-dependent manner
(A, B) Immunoblot for pAMPKα and total AMPKα as an indicator of adipokine signaling and regulator of adipocyte homeostasis in pgWAT from male and female offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P21 (A) and P70 (B); β-Actin served as loading control. The quantitative densitometric summary is shown below to respective immunoblot; pAMPKα was related to total AMPKα. (C, D) Immunoblot for pAkt and total Akt as an indicator of insulin signaling in pgWAT at P21 (C) and P70 (D); β-Actin served as loading control. The quantitative densitometric summary is shown below to respective immunoblot; pAkt was related to total Akt. Co, black bar; HFD, white bar; n=4–6 animals per group; mean ± SEM; *Mann–Whitney test: *P<0.05; **P<0.01; ***P<0.001.

Figure 8. Assessment of expression of genes encoding regulators of adipocyte metabolism in pgWAT from male and female offspring of obese high-fat diet (HFD)-fed and control (standard diet-fed, Co) dams at P70 using qRT-PCR: Comale, HFDmale, Cofemale and HFDfemale
Genes: Mgl1 (A); Fabp4 (B); Srebp1 (C). Co, black bar; HFD, white bar; n=5–8 animals per group; mean ± SEM; *Mann–Whitney test: *P<0.05.
Maternal obesity induces early-onset senescence in adipocytes of male offspring

Since we found a markedly lower differentiation markers in male offspring after maternal obesity, we next assessed senescence using SA-β-gal staining. We found a higher number of SA-β-gal positive adipocytes per field of view and relative to all cells in pgWAT of male mice at the early phase (P21) (Figure 9A). At P120, however, we did not determine any differences in SA-β-gal staining between Co and HFD (Figure 9B).

Discussion

The present study demonstrates two phases of sex-dependent metabolic programming of adipose tissue in offspring of obese dams. First, we identified an early phase (P21) after maternal obesity with (i) an obese body composition, which was more pronounced in females than males; (ii) a sex-dependent distribution of the WAT; (iii) a hypertrophy of adipocytes in both sexes; (iv) a greater inflammatory response and leptin expression in females and males, respectively; (v) a sex-dependent activation of AMPKα and Akt pathway; and (vi) an increase in senescent adipocytes in male offspring.

Second, a late phase after maternal obesity was defined by (i) a persistent obese phenotype in both sexes, with a reversal of the hypertrophy of adipocytes and the inflammatory response in females and senescence in males; (ii) sex-dependent pro-proliferative and stemness inhibiting gene expression in females, but not males; (iii) a dysregulation of AMPKα and Akt signaling in WAT from male offspring; (iv) a sex-dependent dysregulation of lipid metabolism in pgWAT. Finally, the obese body composition persisted in both females and males, whereas the histological assessment of pgWAT revealed sex-specific differences in adipocyte size.

Sex determines metabolic programming of an obese body composition and fat distribution

Clinical and experimental studies identified maternal obesity as a rising risk factor for the offspring's health and the origin of chronic diseases beyond infancy [5,6,10,31]. A hormonal imbalance coupled with the chronic subacute inflammatory state in obese individuals triggers metabolic disorder, such as insulin resistance and diabetes mellitus [32]. Prior studies confirmed that impaired metabolism and obesity in the offspring after maternal HFD are intimately linked to an inflammatory response. Moreover, we demonstrated higher levels of adipocytokines and hormones of the pgWAT in offspring of obese dams, which then contributed to renal and pulmonary metabolic programming [10,33]. There is accumulating evidence that susceptibility to metabolic disorders is sex-dependent. A significant heterogeneity exists between men and women developing the metabolic syndrome, in large part related to hormonal regulation of...
Figure 10. Working model, in which we propose a sex-dependent metabolic programming of adipocyte tissue with an early phase (P21) and a late phase (P70) in offspring of obese dams

This two-step metabolic programming results in a persistent obese phenotype with sex-specific persistent hypertrophy of adipocytes in males, but not females. Based on these findings we speculate that metabolic programming after maternal obesity favors low-grade pathogenic obesity in females and high-grade pathogenic obesity in males.

Sex-dependent metabolic programming of adipocyte size and adipogenic capacity

Whether WAT is a friend or foe is not necessarily defined by its amount, but rather by the localization, the size and the function of adipocytes. For example, hypertrophic adipocytes correlate with systemic insulin resistance, and differ from biochemical functions when compared with smaller adipocytes. These differences involve the expression of anti-inflammatory cytokines, such as adiponectin [40–42]. Prior studies showed that maternal obesity increases adipocyte size and expression of adipokines, including leptin and Il6, in male offspring [43]. Our present study linked a sex-specific inflammatory response in pgWAT of female offspring after maternal obesity to hypertrophy of adipocytes at the early stage; however, this greater adipocyte size was reversed in the later phase. In contrast, in males the hypertrophy of adipocytes persisted throughout life. Dysfunction of the adipose tissue has been identified as a hallmark of age-related diseases, including cardiovascular and metabolic disorders as well as cancer. In particular, an inflammatory microenvironment is characteristic of adipose tissue dysfunction in this context. Preadipocytes are not only central in adipogenesis, but also involved in inflammation through a senescent phenotype, characterized...
Sex regulates metabolic programming of adipocyte metabolism

Adipokines and insulin are crucial in the maintenance of adipocyte homeostasis through AMPKα and Akt signaling pathways, respectively [55,56]. Insulin/Akt promotes the differentiation of preadipocytes to increase the adipose tissue storage capacity [57], and is thereby important in adipogenesis. In contrast, insulin resistance as seen in obese conditions is characteristic for adipose tissue dysfunction [56]. Similarly, AMPK is a key regulator of glucose and energy metabolism as well as stem cell function [55,58]. There is evidence that a lack of AMPKα induces obesity in response to nutrient-overload, identifying thereby AMPKα as a central regulator of adipocytes [55]. In the present study, AMPKα is activated in WAT of male and female offspring during the early phase, and contributes to maintenance of differentiation capacity of adipocytes. On the other hand, during the late phase AMPKα signaling was blocked in males, but not females, and thereby may affect adipocyte function and hypertrophy. Moreover, the hypertrophy of adipocytes in male offspring of obese dams compared with females may also be related to a marked activation of Akt signaling in males. Dysregulation of lipolysis and lipogenesis resulting in altered storage of lipids represents another possible explanation for sex-specific hypertrophy of adipocytes. Indeed, key enzymes involved in these processes were regulated in a sex-dependent manner. The expression of monoacylglycerol lipase (MglL), Fabp4 and Srebp1 was down-regulated in female mice after maternal obesity when compared with either male offspring of obese dams or control females. Interestingly, MglL null mice fed a HFD gained less body weight than wild-type mice and were protected from insulin resistance and hepatic steatosis [59]. Similarly, SREBP1 promotes hypertrophy of adipocytes and FABP4 is associated with metabolic health [60,61]. Reduced expression of these enzymes may in part explain improved adipocyte function and reversal of early metabolic programming in female offspring.

There are few limitations of the work that need to be discussed and addressed in future studies. Since previous studies from our group investigated pgWAT, we have focused on this compartment of adipose tissue in the present study as well. However, the subcutaneous WAT seems to be markedly affected by perinatal HFD and should be studied in-depth in future studies. Moreover, the question arises whether susceptibility to diet-induced obesity differs between male and female after birth. To this end, a second exposure of adult offspring of obese and lean dams should be performed in the future.

In conclusion, the present study provides a comprehensive analysis of sex-dependent metabolic programming of adipose tissue dysfunction and persistent obese body composition. We do not only demonstrate the long-term impact of maternal obesity on fat fraction and distribution, but we also show two phases of metabolic programming and how sex determines function of the pgWAT. Most intriguingly, female offspring show the capacity to reverse adipocyte hypertrophy and functional changes after the initial metabolic insult by maternal obesity. As depicted in Figure 10, the present study highlights the importance of sex-dependent differences in the pathogenesis of metabolic diseases, and provides new insights into the molecular mechanisms of sex-dependent metabolic programming of adipose tissue dysfunction, suggesting a low-grade and high grade pathogenic obesity in female and male after maternal obesity, respectively.
Clinical perspectives

- Adipogenesis and adipocyte function are tightly regulated by the microenvironment. Disruption of these processes by maternal obesity can result in metabolic programming of WAT, ultimately promoting obesity-related diseases. Our study aimed to identify molecular mechanisms determining adipocyte function in a sex-dependent manner.

- Maternal obesity induces sex-independent obese body composition and adipose tissue dysfunction in the offspring. While inflammatory signaling is activated in females early in life, the male offspring show early features of SASP and persistent adipocyte hypertrophy. Obesity persists in both sexes up to 3 months. Interestingly, WAT dysfunction and adipocyte hypertrophy reversed in female offspring, but not in male.

- Our data provide new insights into the molecular mechanisms of sex-dependent metabolic programming of WAT dysfunction and possible low- and high-grade pathogenic obesity. This sex-dependent metabolic programming could account for higher clinical susceptibility of males to obesity-related diseases.

Competing Interests

The authors declare that there are no competing interests associated with the manuscript.

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Author Contribution

K.D., T.P. and M.A.A.A. conceived and designed the research. T.L., E.-K.H., K.D., R.W., C.V., J.S., M.K., C.H. and M.A.A.A. performed the experiments. T.L., E.K.H., K.D., R.W., J.S., T.P. and M.A.A.A. analyzed the data. T.L., E.K.H., K.D., T.P., C.H., J.D. and M.A.A.A. interpreted the results of experiments. T.L., E.K.H. and M.A.A.A. prepared the figures. T.L., E.K.H. and M.A.A.A. drafted the manuscript. T.L., E.K.H., K.D., R.W., C.V., J.S., M.K., T.P., C.H., J.D. and M.A.A.A. approved the final version of manuscript. T.L., E.K.H. and M.A.A.A. edited and revised the manuscript.

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Abbreviations

DLK, \(\Delta\) like non-canonical notch ligand; HFD, high-fat diet; MLI, mean linear intercept; MRI, magnetic resonance imaging; P, postnatal day; PFA, paraformaldehyde; pg, perigonadal; pgWAT, perigonadal white adipose tissue; pp38, phosphorylation of p38; Pref1, preadipocyte factor 1; RT-PCR, reverse transcription polymerase chain reaction; SASP, senescent-associated secretory phenotype; SA-\(\beta\)-gal, senescence-associated \(\beta\)-galactosidase; WAT, white adipose tissue.

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