Microglial and peripheral immune priming is partially sexually dimorphic in adolescent mouse offspring exposed to maternal high-fat diet

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Abstract
Background
Maternal nutrition is critical for proper fetal development. While increased nutrient intake is essential during pregnancy, an excessive consumption of certain nutrients, like fat, can lead to long-lasting detrimental consequences on the offspring. Animal work investigating the consequences of maternal high-fat diet (mHFD) revealed in the offspring a maternal immune activation (MIA) phenotype associated with increased inflammatory signals. This inflammation was proposed as one of the mechanisms causing neuronal circuit dysfunction, notably in the hippocampus, by altering the brain-resident macrophages –microglia. However, the understanding of mechanisms linking inflammation and microglial activities to pathological brain development remains limited. We hypothesized that mHFD-induced inflammation could prime microglia by altering their specific gene expression signature, population density and/or functions.

Methods
We used an integrative approach combining molecular techniques, confocal and scanning electron microscopy, to investigate the effects of mHFD vs control diet on inflammatory priming, as well as microglial density, distribution, morphology, and ultrastructure in mice. These analyses were performed on the mothers, and/or their adolescent offspring at postnatal day (P) 30.

Results
Our study revealed that mHFD results in MIA defined by increased circulating levels of interleukin (IL)-6 in the mothers. This phenotype was associated with an exacerbated inflammatory response to peripheral lipopolysaccharide in mHFD-exposed offspring of both sexes. Microglial morphology was also altered and there were increased microglial interactions with astrocytes in the hippocampus CA1 of mHFD-exposed male offspring, as well as decreased microglia-associated extracellular space pockets in the same region of mHFD-exposed offspring of the two sexes. A decreased mRNA expression of the inflammatory-regulating cytokine Tgfb1 and microglial receptors Tmem119, Trem2 and Cx3cr1 was additionally measured in the hippocampus of mHFD-exposed offspring, especially in males.

Conclusions
Here, we described how dietary habits during pregnancy and nurturing, particularly the consumption
of an enriched fat diet, can influence peripheral immune priming in the offspring. We also found that microglia are affected in terms of gene expression signature, morphology and interactions with the hippocampal parenchyma, in a partially sexually dimorphic manner, which may contribute to the adverse neurodevelopmental outcomes on the offspring.

**Background**

Maternal obesity and dietary overconsumption are risk factors for several health conditions in the offspring, from metabolic syndrome to neurodevelopmental disorders [1–3]. In fact, excess weight has been on the rise in both middle and high income countries, affecting over one third of the global population and about 38.9 million pregnant women worldwide [4]. This increase in the number of overweight or obese pregnant women has been linked to several elements of modern-day environment such as urbanization rate and increased caloric supply [4], which are often associated with increased fast food consumption [5]. This global issue stresses the importance of studying the impact of energy-dense, high-sugar and high-fat food diets during pregnancy [5].

High-fat diet (HFD) successfully models in animals the excessive intake of energy-dense, high-sugar and high-fat food [6]. Of these overconsumed nutrients, fat is of utmost importance for brain growth and development [7]. However, several independent studies using different animal models of maternal (m)HFD have shown a broad range of lasting behavioral alterations in the offspring related to neurodevelopmental disorders from increased anxiety-like behaviors [8–13] to cognitive [8, 14–18], social and motor deficits [9]. Although the mechanisms linking mHFD to the neurodevelopmental alterations remain unclear, several pathological processes such as decreased placental function, hormonal dysregulation, epigenetic alterations [2] and increased central as well as systemic inflammation [1] have been proposed.

In the past decade, epidemiological studies identified “maternal immune activation” (MIA), which refers to maternal systemic inflammation, as a risk factor for several neurodevelopmental disorders [19–22]. This has mostly been observed and researched in the context of infection, where a tight association between bacterial or viral (e.g. influenza) infections during pregnancy and a higher incidence of neurodevelopmental disorders, such as schizophrenia, was uncovered in the progeny
[19, 20, 23]. It is now known that MIA is not limited to infection [19, 20, 23] and may occur after exposure to a large variety of maternal environmental risk factors common to modern-day life including stress [24, 25], smoking, alcohol consumption [26], air pollution [27, 28] and dietary imbalance or overconsumption such as HFD [1]. Postmortem studies revealed neuroimmune alterations in the brain of individuals with developmental disorders, as highlighted by the alteration of microglia – the brain-resident macrophages, in terms of morphology and gene expression signature [29–31], together with changes in the brain levels of cytokines – the molecules involved in signalling and modulation of the inflammatory state, among other important functions [19, 20, 23].

In animal models of mHFD, the inflammatory status in the offspring brain, notably in hippocampus, has been characterized by measuring changes of several inflammatory mediators during the lifespan. The mRNA expression of Cd11b and Tlr4 – two genes involved in the innate immune response – were found to be increased in the hippocampus of male and female rat offspring as early as postnatal day (P)1, followed by interleukin (Il)1β at juvenile stages [8]. During adolescence, both male and female rat offspring exhibited greater levels of the cytokine Il6, as well as of the inflammatory regulators nuclear factor kb (Nfk) and mitogen-activated kinase protein (Mkp) 1 in the hippocampus [32].

Adolescent male rat offspring also expressed increased levels of Nfkb inhibitor (IkB) in the hippocampus. Later in adult life, a few studies observed no change in cytokines [10] while others found greater expression of Il1β in the hippocampus of male rats [8], and lower expression of IkB and Il1ra in the hippocampus of female rats [13] exposed to mHFD. However, it remains unclear if this increased gene expression of inflammatory mediators is directly linked to a global inflammatory status or to altered microglial activities in the offspring brain. So far, the reported microglial changes induced by mHFD in the offspring brain comprise a higher density of ionized calcium-binding adapter 1 (IBA1)-positive (⁺) cells in the hippocampus of adult male and female rats [8] suggesting changes in microglial function.

Studies performed in animal models and human postmortem samples have revealed that microglia modulate key neurodevelopmental processes – cell migration and maturation, followed by the formation of neuronal circuits and myelination – in a sexually dimorphic manner [33–35]. Microglia also
contribute to neuronal circuit refinement during adolescence [36] and adulthood [29, 35, 37-39]. Paralleling the immune alterations described above, mHFD was shown to be associated with synaptic changes resulting in decreased synapses density and/or spines density and stability in rodent offspring across brain regions and stages of the lifespan [16, 25, 40-42] including the hippocampus of adolescent mice [16]. Microglial cell density, distribution, gene expression signature, physiological functions and/or response to immune challenges may be altered upon exposure to various environmental factors, thus impacting on their sculpting of the brain network [43, 44]. This phenomenon is referred to as immune priming [43, 44]. Studying microglial priming during sensitive periods like adolescence, in which the brain and especially the hippocampus experience important synaptic changes, may help to understand the pathological cascade underlying mHFD.

Using a mouse model, we explored the changes of microglia in the offspring exposed to mHFD vs control diet (CD). Specifically, we investigated microglia-related gene expression in whole hippocampus, as well as microglial density, distribution, morphology and ultrastructure in the dorsal hippocampus CA1. We focused on this region considering its crucial importance for memory, learning [45, 46] and executive [47] functions that are affected with MIA [19-22] including with mHFD [8, 15, 16, 42]. The measurements were made at P30, which corresponds to early adolescence [48]. To characterize our model, we also assessed metabolic changes in the mothers and offspring, together with peripheral immune priming in the adolescent offspring, using a single systemic immune challenge with lipopolysaccharide (LPS) at P30. Male and female offspring were compared in all experiments to investigate possible sex differences.

Methods

Animal and tissue processing

All animal protocols were approved by McGill University’s Facility Animal Care Committee under the guidelines of the Canadian Council on Animal Care. Mice were submitted to 12 hours dark/light cycles (8:00–20:00) with an ad libitum access to water and food. They were housed two to five per cage, except during pregnancy and nurturing, when the dams were housed single or alone with their litter. Paired-housed C57BL/6N female mice aged 5–6 weeks were obtained from Charles River (St-Constant,
QC, Canada) and habituated for one week prior to the protocol onset. They were then provided with either a HFD (diet rich in saturated and unsaturated fats; 60% kcal by lipids; Teklad TD.06414, ENVIGO, Indianapolis, IN, United States; Supplementary Fig. 1e) or CD (Teklad 2014, ENVIGO; Supplementary Fig. 1e) *ad libitum* starting 4 weeks prior to mating, throughout gestation and nurturing, until weaning of their litter (Fig. 1). During the diet protocol, food consumption and weight were recorded in the dams to identify potential metabolic changes. Blood samples were collected by submandibular puncture after 8 weeks on the diet to evaluate the inflammatory profile and assess circulating glucose levels. Dams were anesthetized with a rodent cocktail (0.3 mL/100 g) containing ketamine [100 mg/mL], xylazine [20 mg/mL] and aceprozamine [10 mg/mL], fresh-decapitated at the end of the protocol, after 9–10 weeks on the diet, and their fat deposits were dissected (Fig. 1). Male and female offspring were compared to study mHFD effects in the two sexes. Animals presenting major anomalies (e.g. unopened or abnormal eye, dwarf or severe tooth malformation) were excluded. Only 1-2 offspring of each sex per litter were used for each experiment to prevent litter effects. At P21, the offspring were weaned and switched to CD. At P30, offspring were anesthetized with rodent cocktail and their blood, brain and fat tissues were collected. One cohort of offspring (n = 5–6 animal/sex/diet) was fresh-decapitated, their brain rapidly extracted, and the hippocampus dissected, flash-frozen on dry ice and stored at -80 °C until mRNA analysis by real-time quantitative polymerase chain reaction (rt-qPCR). Another cohort of offspring was anesthetised as described above, perfused with 15 mL of phosphate-buffered saline (PBS) followed by ~ 180 mL of 4% paraformaldehyde (PFA) in [50mM] phosphate-buffer (PB) for histological analysis (n = 5 animals/sex/diet). PFA-fixed brains were post-fixed in 4% PFA for 24 hours at 4°C, then immersed in 30% glucose solution (in [50mM] PBS, pH = 7.4) at -4 °C for 48 hours and flash-frozen. Frozen brains were cut into 30 µm-coronal sections using a cryostat (CM3050S, Leica Biosystems, Wetzlar, Germany) and stored free-floating in cryoprotectant (30% ethylene glycol, 30% glycerol in [50mM] PBS, pH = 7.4) at -20 °C until use for histochemistry. The last cohort of animals was anesthetised as described above, flushed with 15 mL of PBS and perfused with 75 mL of 3.5% acrolein in [100mM] PB (pH = 7.4) and 150 mL of 4% PFA in [100mM] PB (pH = 7.4) for ultrastructural analysis (n = 4
animals/sex/diet). PFA/acrolein-fixed brains were post-fixed for 2 hours in 4% PFA at 4°C, washed in PBS, cut into 50 µm-coronal sections with a vibratome (VT1200S, Leica Biosystems) and stored in cryoprotectant solution at -20 °C until use (Fig. 1).

Adolescent peripheral immune challenge
To evaluate peripheral immune priming, a subset of animals (n = 5–7 animals/sex/diet/treatment) received a single intraperitoneal injection at P30, of either 100 µg/kg lipopolysaccharide (LPS) from Escherichia coli O111:B4 (cat# 62325, Millipore Sigma, Burlington, MA, United States) –a component of the gram-negative bacterial wall used to model an inflammatory response– or saline (SAL; 0.9% NaCl solution, cat# 01966010, Hospira, United States). Eight hours after the administration, animals were anesthetized with rodent cocktail (0.3 mL/100 g) and decapitated. Trunk blood was collected in heparinized tubes and centrifuged at 3600 rpm for 10 minutes. Plasma was collected and flash-frozen and stored at -80 °C until analysis by multiplex-ELISA (see section Cytokines measurement by multiplex-ELISA) (Fig. 1).

Physiological changes
To determine the phenotype induced by our diet model, a thorough characterization of the gestation and litter, as well as metabolic changes including weight, food intake, fat distribution and glucose levels was performed in the dams. The methods and results are presented as Supplementary material.

Cytokine measurement by multiplex-ELISA
To determine changes in the inflammatory profile of the dams, peripheral inflammation was assessed at the end of nurturing (after 8–9 weeks on the diet) (n = 5–6 dams/diet). Similarly, the peripheral inflammatory profile of the pubertal offspring (P30) was characterized in homeostatic (SAL-treated) vs immune (LPS-treated) contexts (n = 5–6 animals/sex/diet/treatment). Levels of the cytokines IL-1β, IL-6, IL-10, IL-17 and tumor necrosis factor (TNF)-α were measured in maternal and offspring plasma using Luminex Multiplex Assay (MILLIPLEX MAP kit Mouse Cytokine/Chemokine Magnetic Bead Panel, cat# MCYTOMAG-70K, Millipore Sigma). Prior to the experiment, the multiplex assay reader Luminex MAGPIX (ThermoFisher Scientific, Waltham, MA, United States) was calibrated (#cat MPXCALK25, Thermo Fisher Scientific) and its performance verified (#cat MPXPVERK25, Thermo Fisher Scientific)
according to the manufacturers’ guidelines. Plasma samples were diluted 1:2 by combining 30 µL of plasma with the provided drive fluid buffer. Immunoassays were performed following the manufacturer’s instructions. Diluted plasma was incubated overnight (~16 hours) with primary antibodies at 4 °C, 1 hour with detection antibodies at room temperature and 30 minutes with Streptavidin-beads at room temperature. Samples were resuspended in 150 µL drive fluid and run through the Luminex MAGPIX to assess cytokines levels.

**Real-time quantitative polymerase chain-reaction**

At P30, the two hippocampi of each offspring (n = 5–6 animals/sex/diet) were homogenized in Trizol (cat#15596-026, Ambion, Austin, TX, United States) and RNA was extracted using the Trizol/chloroform method followed by an isopropanol precipitation. The RNA pellet was washed once in 75% ethanol, let dry, then eluted in Nuclease-free water (cat#AM9937, Ambion). Samples were dosed using the NanoDrop ND-1000 kit (ThermoFisher Scientific).

Genomic DNA was removed from 1 µg isolated RNA sample by enzymatic degradation (cat# G488, Applied Biological Materials Inc, Richmond, BC, Canada). Purified RNA was used to obtain complementary DNA (cDNA) by reverse transcriptase reaction with iScript 5X MasterMix (cat#1708890, BioRad Laboratories, Hercules, CA, United States) using a TI thermocycler (Biometra, Göttingen, Germany). Using diluted cDNA, rt-qPCR was performed with the SybrGreen technology and cycle threshold (Ct) was determined using a LightCycler 480 II (Roche, Basel, Switzerland). RT-qPCR measured the microglial function-related genes *Tmem119* (primer 5’ - TTC TTC CGG CAG TAC GTG AT; primer 3’ - CGA GGA TGG GTA GTA GGC TG), *Aif1* (primer 5’ – TCT GCC GTC CAA ACT TGA AG; primer 3’ – GCC ACT GGA CAC CTC TCT AA), *Trem2* (primer 5’ – ACC CTC TAG ATG ACC AAG ATG C, primer 3’ – TTG GGC ACC CTC GAA ACT C), *Cx3cr1* (primer 5’ – CAA GCT CAC GAC TGC CTT CT; primer 3’ – TGT CCG GTT GTT CAT GGA GTT), *Tgfb1* (primer 5’ – ACA TGT GGA ACT CTA CCA GAA A; primer 3’ – CTG CCG TAC AAC TCC AGT GA) as well as housekeeping gene *Rpl32* (primer 5’-TTG TGT CTC CCA TAA CCG ATG, primer 3’-TTA AGC GAA ACT GGC GGA AAC). Other than the cytokine *Tgfb1*, we attempted to assess mRNA levels of *Il1b* and *Il6*, however we failed to obtain robust amplification by rt-qPCR. Relative expression was calculated by determining the difference of Ct between the genes of interest
and the housekeeping gene (2^{−ΔΔCt}), while arbitrarily considering CD-exposed male offspring as the reference group [49]. Results were presented in ratio fold and statistical analysis was assessed on ΔΔCt which are normally distributed.

**Microglial density, distribution and morphology analysis**

PFA-perfused brain sections were double-immunostained against IBA1 (labels all myeloid cells including microglia) and transmembrane protein 119 (TMEM119; microglia-specific [50]) to evaluate microglial density, distribution and morphology, as well as myeloid cell infiltration as previously described [51]. Three to four brain sections containing the dorsal hippocampus CA1 (Bregma: -1.31 to -1.91, stereotaxic atlas of Paxinos and Franklin 4th edition [52]) were selected for each offspring (n = 5 animals/sex/diet). Sections were washed, incubated 40 minutes in sodium citrate buffer to expose epitopes, washed again and treated with 0.1% NaBH₄ (cat#480886, MilliporeSigma) to quench autofluorescence. Afterwards, brain sections were incubated in blocking solution (0.5% gelatin, 5% donkey serum, 0.1% Triton X-100) for 1 hour at room temperature, washed and incubated overnight at 4 °C with a cocktail of primary antibodies in blocking solution: mouse anti-IBA1 (1:190; cat# MABN92, Millipore Sigma) and rabbit anti-TMEM119 (1:300; cat# ab209064, Abcam). The sections were washed and incubated with the secondary antibodies donkey anti-mouse Alexa555-conjugated (1:300 in PBS; cat#A31570, Invitrogen, ThermoFisher Scientific) and donkey anti-rabbit Alexa647-conjugated (1:300 in PBS; cat#A31573, Invitrogen, ThermoFisher Scientific) for 1.5 hour at room temperature. Sections were then mounted on slides and coverslipped in Fluoromount mounting medium (cat# 0100-01, SouthernBiotech, Birmingham, AB, United States).

For density and distribution analysis, all stained sections were imaged in a single plane at 20X using an Axio Imager M2 epifluorescence microscope equipped with an AxioCam MRm camera (Zeiss, Oberkochen, Germany). For morphology analysis, z-stacks (stratum radiatum (st rad): 18–20 z-stacks/animal; stratum lacunosum moleculare (st lac mol): 10–15 z-stacks/animal) were captured at 40X using a Quorum Wave FX spinning disc confocal microscope (Quorum Technologies, Guelph, ON, Canada) equipped with an ORCA-R2 camera (512 × 512 pixels; Hamamatsu Photonics, Hamamatsu, Japan). The z-stacks were merged into a single plane using Volocity software (Version 5.4,
PerkinElmer, Waltham, MA, United States). 

All analyses were performed blind to the experimental condition using ImageJ software (v1.51j8; National Institute of Health, Bethesda, MD, United States). Total count of IBA1⁺/TMEM119⁺ (microglia) and IBA1⁺/TMEM119⁻ (peripheral macrophages) cells was compiled for the dorsal hippocampus CA1, st rad and st lac mol, across 6–8 hippocampi per animal, using the analyze particles plugin [51, 53]. Afterwards, cellular distribution was assessed using the nearest neighbour distance (NDD) plugin. For microglial morphology analysis, the number of branches and junctions, as well as average and longest branch length were obtained (st rad: n = 20–25 microglia/animal, st lac mol: n = 15–20 microglia/animal, N = 5 animals/sex/diet) using a semi-automatic method adapted from [54]. For each microglial cell, soma and manual arbor were also traced using the freehand tool and polygon selection tool to obtain area values. Morphological index for each microglia was calculated by dividing the soma area by the manual arborization area to help identify microglial changes from their steady state [51, 53]. Manual arbor selection was further processed in a semi-automated manner to obtain unsharp mask of the cell, adjusted by the observer when needed, and area of the cell and shape descriptors (i.e. circularity, solidity and aspect ratio) values were measured. Circularity was calculated by: \(4\pi \times \frac{\text{Area}}{\text{Perimeter}^2}\), for which a value of 1.0 represent a perfect circle and towards 0.0 an elongated shape. Solidity was calculated by dividing the cell area by the convex cell area meaning that a value close to 0.0 indicate a porous shape and close to 1.0 a convex shape. Finally, the aspect ratio was calculated by dividing the major axis of the cell by the minor axis of the cell, meaning a value of 1.0 similar ratio of minor and major axis and the higher the value the more elongated the cell is. The mask of the cell was skeletonized and analyzed using skeleton 2D/3D plugin. Skeleton analysis allowed us to determine number, average length, and maximal length of branches as well as number of junctions.

**Microglial ultrastructure analysis**

Two PFA/acrolein-perfused brain sections containing the dorsal hippocampus CA1 (Bregma − 1.67 mm; stereotaxic atlas of Paxinos and Franklin 4th edition [52]) were selected in each of four
animals per group. Sections were washed in PBS, then quenched 10 minutes in 0.3% \( \text{H}_2\text{O}_2 \) in PBS and permeabilized 30 minutes in 0.1% \( \text{NaBH}_4 \) in PBS. Sections were first incubated 1 hour at room temperature in blocking solution (10% fetal bovine serum, 3% bovine serum albumin, 0.01% Triton X-100 in [50mM] tris-buffered saline (TBS) before incubation with rabbit anti-IBA1 primary antibody (1:1000; \text{cat#019-19741, FUJIFILM Wako Chemical, Osaka, Japan}) in blocking solution overnight at 4 °C. The following day, antibody was washed out and the sections were incubated with biotinylated goat anti-rabbit secondary antibody (\text{cat#111-066-046, Jackson ImmunoResearch, West Grove, PA, United States}) in TBS for 1.5 hour, followed by avidin-biotin complex solution (1:1:100 in TBS; \text{cat#PK-6100, Vector Laboratories, Burlingame, CA, United States}) for 1 hour. The staining was revealed in 0.05% diaminobenzidine (DAB; \text{cat# D5905-50TAB, Millipore Sigma}) with 0.015% \( \text{H}_2\text{O}_2 \) in TBS for 4.5 minutes at room temperature.

The immunostained sections were next post-fixed flat in osmium-thiocarbohydrazide-osmium for scanning electron microscopy (SEM). In particular, sections were incubated in 3% ferrocyanide (\text{cat#PFC232.250, BioShop, Burlington, ON, Canada}) diluted in water combined (1:1) with 4% aqueous osmium tetroxide (\text{cat#19170, Electron Microscopy Sciences, Hatfield, PA, United States}) for 1 hour, in 1% thiocarbohydrazide diluted in water (\text{cat# 2231-57-4, Electron Microscopy Sciences}) for 20 minutes, in 2% osmium tetroxide diluted in water, then dehydrated in ascending concentration of ethanol (2 × 35%, 50%, 70%, 80%, 90%, 3 × 100%) followed by propylene oxide (3x) for 5 minutes each. After post-fixation, tissues were embedded in Durcupan ACM resin (\text{cat# 44611–44614, Millipore Sigma}) for 24 hours and carefully placed between two ACLAR® embedding films (\text{cat# 50425-25, Electron Microscopy Sciences}) and the resin was let to polymerize at 55 °C for 72 hours.

Regions of selection –dorsal hippocampus CA1– were excised from the embedded sections on ACLAR® sheets and re-embedded on top of a resin block for ultrathin sectioning (Ultracut UC7 ultramicrotome, Leica Biosystems). Ultrathin sections (~75 nm thickness) were collected and placed on a silicon nitride chip and glued on specimen mounts for SEM. Seven to 12 microglial cell bodies in each animal/layer of interest (\textit{st rad} and \textit{st lac mol}) were imaged at 5 nm of resolution using a
Crossbeam 540 field emission SEM with a Gemini column (Zeiss).

Ultrastructural analysis was performed blind to the experimental conditions using ImageJ software (st rad: n = 32-41 microglia/sex/diet, st lac mol: n = 28-36 microglia/sex/diet, N = 4 animals/sex/diet).

Microglial endoplasmic reticulum, Golgi apparatus, lysosomes, lipofuscin, mitochondria and endosomes were first analyzed quantitatively [55]. Dilation of the endoplasmic reticulum and/or Golgi apparatus was noted when the distance between the cisternal membranes was 50 nm or greater [56, 57]. Lysosomes were identified by their dense heterogeneous contents enclosed by a single membrane [58, 59]. Secondary lysosomes were differentiated from primary lysosomes by their contacts with fusion endosomes. Tertiary lysosomes were identified by their contacts with lipofuscin and often also with fusion endosomes [58, 60]. Lipofuscin granules, for their part, were identified by their oval- or round structure and finely granular composition with a unique fingerprint-like pattern [60]. Mitochondria were considered as elongated when their length was greater than 1 µm [55].

Microglial contacts with the cell bodies from other brain cells (i.e. astrocytes, neurons, oligodendrocytes) as well as blood vessels were quantified. For the neurons, contacted myelinated axons and synaptic elements –pre-synaptic axon terminal or post-synaptic dendritic spine– were further identified. Astrocytic cells were identified by their pale nuclei with a thin rim of heterochromatin and pale irregular cytoplasm, often containing intermediary filaments [61]. Neurons were distinguished by their pale nuclei and pale cytoplasm, often with an apical dendrite and innervation from axon terminals [61]. Pre-synaptic axon terminals were differentiated by their synaptic vesicles, while post-synaptic spines were in contact with a pre-synaptic axon terminal, often with a visible post-synaptic density at their junction [61]. Microglia were recognized by their dark irregular nuclei with a heterogenous chromatin pattern and a dark irregular cytoplasm, often containing short endoplasmic reticulum cisternae and lipidic inclusions (i.e. lipofuscin, lipid bodies or droplets, lysosomes) [61]. Similar to microglia, oligodendrocytes were identified by their dark nuclei with a heterogenous chromatin pattern and dark squarish or rectangular-shape cytoplasm, often containing short and wide endoplasmic reticulum cisternae organised in the vicinity of the nucleus and ribosomes, as well as a wider space between nuclear membranes than microglia [61]. In the
vicinity of microglia, the occurrence of degradation activities (degenerating myelin, extracellular digestion) was also noted. Extracellular digestion, also named “exophagy”, was identified by extracellular space pockets containing degraded elements or debris [62, 63]. In contrast, degenerating myelin was recognized by ballooning, swelling or distancing of myelin sheaths [61].

In addition, the density of dark cells and apoptotic cells was assessed (N = 4 animals/sex/diet). Then, dark cells were analyzed in a semi-quantitative manner (st rad: n = 0–6 dark cells/sex/diet, st lac mol: n = 13–19 dark cells/sex/diet, N = 4 animals/sex/diet). Dark cells were distinguished by their electron-dense nuclei showing a loss of the chromatin pattern and electron-dense cytoplasm presenting several signs of cellular stress (i.e. dilated endoplasmic reticulum and Golgi apparatus cisternae, elongated mitochondria) [64]. In the present study, we distinguished between two types of dark cells: microglia and perivascular cells. In addition to their dark features, dark microglia were recognised by their microglial characteristics [55, 64] and were located inside the brain parenchyma. Dark perivascular cells were identified by their localization enclosed in the perivascular space and possessed the dark features mentioned above. Apoptotic cells were also dark, and recognised with their pyknotic nucleus and accumulation of autophagic endosomes [65].

Sample size of dark cells in st lac mol (n = 59 individual dark cells total) was considered sufficient to attempt statistical data analysis. This assumption was based on the sample size calculated using G*Power software (v3.1.9.6) [66] to detect a large effect size of 0.4 that was estimated to 52 individuals cells.

**Statistical analyses**

Data are reported as means ± standard error of the mean (SEM). Sample size (n) refers to individual animals for metabolic parameters, immune priming, gene expression as well as microglial density and morphology analyses, while it refers to individual microglia or dark cells for ultrastructural analyses. Statistical analyses were conducted using Prism 8 (v.8.3, GraphPad Software, San Diego, CA, United States). Normality was verified using Shapiro-Wilk and assessed by QQ plot. For normally distributed dataset, Grubbs’ test was used to identify outliers that were removed from the datasets prior to performing parametric statistic tests. To compare CD vs HFD in the dams, a Student t-test was used
for non-repeated measures including glucose levels, weight and fat deposit measurements, while a 2-way analysis of variance (ANOVA) for repeated measures test was used for comparing weight and dietary follow up data across time. In the offspring, a 2-way ANOVA was used to compare CD vs mHFD, as well as male and female animals for metabolic parameters (i.e. weight and fat deposits), gene expression, as well as microglial density, distribution, morphology and ultrastructure. Significant ANOVA tests with a Sex*Diet interaction were followed by Bonferroni post-hoc test to identify significant differences between individual groups. For non-normally distributed dataset, a Mann-Whitney test was used to compare CD vs HFD in the dams for cytokines profile, gestation duration and litter size. For non-normally distributed offspring dataset, a mixed-effect model was used to compare CD vs mHFD, males vs females in terms of myeloid cells infiltration, as well as to compare SAL vs LPS groups after immune challenge to assess cytokines profile. Significant mixed-effect model was followed by a Bonferroni post-hoc test. Statistically significant differences were considered for p value < 0.05.

Results
High-fat diet induces long-term increase of peripheral IL-6 and fat deposits in dams and male offspring
Previously, rodent models of mHFD were shown to induce peripheral inflammation, notably by increasing the levels of cytokines in maternal blood circulation [14, 28, 67, 68]. To evaluate the maternal immune profile in our mouse model of mHFD, we measured plasma levels of pro- (IL-1β, IL-6, IL-17, TNF-α) and anti-inflammatory (IL-6, IL-10) cytokines in the dams at weaning of their litter by multiplex-ELISA. Plasma levels of IL-6 were significantly increased (p = 0.0079) in HFD-fed dams (17.74 ± 8.79 pg/mL) compared to CD-fed dams (3.636 ± 0.933 pg/mL) (Fig. 2d), while both diet groups had similar levels of IL-1 β, IL-10, IL-17 and TNF-α (Fig. 2a-b, e). Overall, these results suggest a MIA phenotype induced by mHFD in our model, confirmed by the increase of IL-6.

Other than peripheral inflammation, HFD has been associated with a variety of metabolic changes including increased body fat, obesity, diabetic-like phenotype (glucose and/or insulin intolerance) [69] and decreased fertility [70]. To characterize further our model, we assessed weight and food consumption, glucose levels, fat deposits, as well as gestation duration and litter size in the dams.
These measurements revealed that mHFD does not induce obesity, although it is associated with an overconsumption of fats to the detriment of carbohydrates (Supplementary Fig. 1). HFD nevertheless resulted in an increase of retroperitoneal, subcutaneous and perigonadal fat deposition at the measured endpoint, without driving an overall increase of total body mass or an obese phenotype (Supplementary Fig. 2). Importantly, it did not lead to glycemia or fertility alterations, ruling out major metabolic alterations as often seen in diet-induced animal models of obesity (Supplementary Fig. 2).

Previous work on mHFD revealed that offspring are more prone to developing metabolic syndrome, which includes increase in fat deposits and body weight [2]. In our study, mHFD offspring had similar body weight compared to CD offspring at P30, but the mHFD males showed increased perigonadal fat deposits compared to CD males (Supplementary Fig. 3), which highlights a sexually dimorphic effect of mHFD on offspring fat deposition.

**mHFD offspring have higher IL-6 plasma levels after LPS-induced immune challenge**

To assess peripheral immune priming in the adolescent offspring, we measured circulating cytokines by multiplex ELISA at P30, eight hours after systemic injection of low dose LPS vs SAL. This time point corresponds to a period of inflammatory resolution after the immune challenge. In LPS-injected animals, plasma levels of IL-6 were significantly elevated ($F_{(1,37)} = 10.44, p = 0.0317$) in mHFD vs CD offspring regardless of their sex ($436.8 \pm 107.0$ pg/mL vs $197.9 \pm 74.1$ pg/mL) (Fig. 3d). Moreover, regardless of their maternal diet, LPS-treated female offspring had significantly ($F_{(1,37)} = 5.324, p = 0.0004$) increased plasma levels of IL-6 compared to LPS-treated male offspring ($504.1 \pm 113.8$ pg/mL vs $152.3 \pm 44.0$ pg/mL) (Fig. 3d). Levels of TNF-α and IL-10 were also significantly increased (TNF-α: $F_{(1,37)} = 16.43, p = 0.0002$; IL-10: $F_{(1,21)} = 30.63, p < 0.0001$) in LPS-treated offspring compared to SAL-treated offspring regardless of their sex or maternal diet (TNF-α: $15.23 \pm 2.66$ pg/mL vs $4.435 \pm 0.745$ pg/mL; IL-10: $82.54 \pm 8.03$ pg/mL vs $10.76 \pm 7.40$ pg/mL) (Fig. 3c, e-f). Finally, levels of IL-1β and IL-17 were similar between SAL-treated and LPS-treated offspring regardless of their sex and maternal diet (Fig. 3a-b, f). Together, these results indicate that systemic LPS administration induced IL-6, IL-10 and TNF-α release detected eight hours afterwards in CD and mHFD offspring, while IL-6
release was significantly exacerbated in mHFD offspring, indicating either a sustained inflammation or a stronger response to the immune challenge specific to this cytokine upon exposure to mHFD. Male offspring have altered gene expression after mHFD whereas both male and female offspring exhibit altered microglial morphology

Other than a peripheral inflammatory response, immune priming has been associated with changes in gene expression and/or morphology of immune cells -including microglia in the brain [1, 28, 72]. To characterize microglia-related gene changes, we used rt-qPCR to study mRNA expression in whole hippocampus of mHFD vs CD offspring at P30 (Fig. 4a). We focused on *Tgfb1* -a cytokine that modulates inflammation and microglia survival [73]- as well as on receptors mainly expressed by microglia in the brain that are involved with the regulation of inflammation (i.e. *Aif1* [74]), microglial survival (i.e. *Tmem119* [50], *Trem2* [75]) or synaptic remodeling (i.e. *Cx3cr1* [76-78], *Trem2* [76, 79-81]). mHFD-exposed male offspring had significantly reduced expression of *Tgfb1* \(F\,(1,19) = 10.05, \text{CD male}: p = 0.0045, \text{CD female}: p = 0.0018, \text{mHFD female}: p = 0.0008\) compared to other offspring groups (detailed in Table 1; Fig. 4b). Expression of *Aif1* was however similar between groups (Fig. 4c). mHFD-exposed male offspring also had a significantly lower ratio fold of *Tmem119* \(F\,(1,19) = 19.47, \text{CD male}: p < 0.0001, \text{CD female}: p < 0.0001, \text{mHFD female}: p < 0.0001\), *Trem2* \(F\,(1,19) = 20.88, \text{CD male}: p < 0.0001, \text{CD female}: p < 0.0001, \text{mHFD female}: p < 0.0001\) and *Cx3cr1* \(F\,(1,19) = 16.44, \text{CD male}: p = 0.0002, \text{CD female}: p < 0.0001, \text{mHFD female}: p < 0.0001\) compared to the other offspring groups (detailed in Table 1; Fig. 4d-f). Together, these results indicate that mHFD leads to altered expression, specifically in males, of inflammatory-regulating (*Tgfb1*) as well as microglial function-related (*Tmem119, Trem2 and Cx3cr1*) genes.
Table 1
mHFD effect on hippocampal mRNA levels of P30 offspring. Transcripts level of Tgf1b, Aif1, Tmem119, Trem2 and Cx3cr1 were normalized by Rpl32, where CD male offspring represent the reference group. $2^{-\Delta\Delta Ct}$: ratio fold compared to expression of the reference group, $\Delta Ct$: difference of cycle threshold between $\Delta Ct$ of the target gene and $\Delta Ct$ of Rpl32, CD: control diet, mHFD: maternal high-fat diet.

| Gene     | Male CD | mHFD | Female CD | mHFD | F       | p       |
|----------|---------|------|-----------|------|---------|---------|
| Tgf1b    | $2^{-\Delta Ct}=1.000$, $\Delta Ct=0.000$ ± 0.218 | $2^{-\Delta Ct}=0.2764$, $\Delta Ct=1.860$ ± 0.262 | $2^{-\Delta Ct}=1.139$, $\Delta Ct=0.190$ ± 0.407 | $2^{-\Delta Ct}=1.282$, $\Delta Ct=0.360$ ± 0.335 | Sex*Diет:10.05 | Sex: 14.16, Diет: 6.965 |
| Aif1     | $2^{-\Delta Ct}=1.000$, $\Delta Ct=0.000$ ± 0.328 | $2^{-\Delta Ct}=0.7169$, $\Delta Ct=0.480$ ± 0.340 | $2^{-\Delta Ct}=1.069$, $\Delta Ct=0.100$ ± 0.290 | $2^{-\Delta Ct}=1.196$, $\Delta Ct=0.260$ ± 0.299 | Sex*Diет:1.035 | Sex: 1.783, Diет: 0.2588 |
| Tmem119  | $2^{-\Delta Ct}=1.000$, $\Delta Ct=0.000$ ± 0.313 | $2^{-\Delta Ct}=0.1346$, $\Delta Ct=2.89$ ± 0.269 | $2^{-\Delta Ct}=1.260$, $\Delta Ct=-0.330$ ± 0.350 | $2^{-\Delta Ct}=1.101$, $\Delta Ct=-0.140$ ± 0.265 | Sex*Diет:19.47 | Sex: 30.15, Diет: 25.34 |
| Trem2    | $2^{-\Delta Ct}=1.000$, $\Delta Ct=0.000$ ± 0.282 | $2^{-\Delta Ct}=0.1690$, $\Delta Ct=2.570$ ± 0.229 | $2^{-\Delta Ct}=0.9781$, $\Delta Ct=0.030$ ± 0.326 | $2^{-\Delta Ct}=0.9598$, $\Delta Ct=0.060$ ± 0.242 | Sex*Diет:20.88 | Sex: 19.90, Diет: 21.88 |
| Cx3cr1   | $2^{-\Delta Ct}=1.000$, $\Delta Ct=0.000$ ± 0.203 | $2^{-\Delta Ct}=0.1724$, $\Delta Ct=2.520$ ± 0.479 | $2^{-\Delta Ct}=1.147$, $\Delta Ct=-0.200$ ± 0.315 | $2^{-\Delta Ct}=1.207$, $\Delta Ct=-0.270$ ± 0.276 | Sex*Diет:16.44 | Sex: 21.92, Diет: 14.71 |

After performing rt-qPCR, we characterized the density, distribution, morphology and ultrastructure of microglia in mHFD vs CD-exposed offspring at P30. We focused on the dorsal hippocampus CA1, particularly the st rad and st lac mol –two main layers where neuronal plasticity occurs during cognitive processes [82] and that are associated to behavioral deficits previously reported in mHFD animal models [8, 14–18]. In both layers (Fig. 5a-e, n-r), the density and distribution of microglia (IBA1+/TMEM119+) and infiltrated myeloid cells (IBA1+/TMEM119−) were similar between groups (Table 2; Fig. 5j, w). Of note, infiltrated myeloid cells were marginal, accounting for 0.207% of IBA1+ cells in the st rad and 0.240% in st lac mol. Regardless of their sex and maternal diet, adolescent offspring displayed similar values for microglial soma, arbor and cell area, as well as morphological index (soma area/manual arborization area) in both CA1 st rad and st lac mol (Table 3–4; Fig. 5f-i, s-v). In st rad, further analysis of ‘skeletonized’ microglia revealed no significant difference between offspring groups in terms of number of branches, junctions, as well as average and maximal branch length (Table 3). However, shape descriptor analysis of microglia identified a significant decrease in their circularity value ($F_{(1,16)} = 4.683, p = 0.0459$) in mHFD offspring compared to controls ($0.0265 ± 0.0004$ vs $0.0295 ± 0.0006$), but solidity and aspect ratio remained unchanged (Table 3; Fig. 5k-m). In st lac mol, microglia of mHFD-exposed offspring had significantly shorter branch length ($F_{(1,16)} = 4.553, p = 0.0487$) compared to CD-exposed offspring ($3.276 ± 0.015 \mu m$ vs $3.442 ± 0.022 \mu m$).
In addition to their shorter branch length, microglia of mHFD-exposed offspring had a significantly increased solidity \( (F_{(1,16)} = 5.616, p = 0.0307) \) compared to CD offspring \( (0.2845 \pm 0.0030 \text{ vs } 0.2603 \pm 0.0023) \), regardless of the sex (Table 4; Fig. 5y), which could indicate a difference in microglial arborization distribution and/or organization with mHFD. In this layer, microglial branch number, maximal branch length, junction number, circularity and aspect ratio were also similar between groups. Together, these morphological changes align with a microglial priming hypothesis, in which mHFD alters microglial morphology.

### Table 2

mHFD effects on microglial density, distribution, and peripheral myeloid cell infiltration in the dorsal hippocampus CA1 of P30 offspring. %Infiltration: Average percentage of IBA1\(^+\)/TMEM119\(^-\) cells on total myeloid cells count, a.u.: arbitrary unit, CD: control diet, St lac mol: stratum lacunosum moleculare, mHFD: maternal high-fat diet, St rad: stratum radiatum.

| Parameters       | Mean ± standard error of the mean | F     | p     |
|------------------|----------------------------------|-------|-------|
|                  | Male CD | mHFD | Female CD | mHFD | Sex*Diet | Sex | Diet |
| **St rad**       |         |      |           |      |           |     |      |
| Density (cells/mm\(^2\)) | 228.5 ± 12.1 | 233.8 ± 2.1 | 225.4 ± 10.1 | 214.9 ± 5.4 | Sex*Diet: 0.8838 | Sex: 1.725 | Diet: 0.09162 |
|                  |         |      |           |      |           |     |      |
| Spacing Index (a.u.) | 0.452 ± 0.014 | 0.469 ± 0.006 | 0.456 ± 0.011 | 0.454 ± 0.013 | Sex*Diet: 0.6857 | Sex: 0.2303 | Diet: 0.4325 |
|                  |         |      |           |      |           |     |      |
| Cluster          | 0.261 ± 0.138 | 0.051 ± 0.031 | 0.182 ± 0.084 | 0.222 ± 0.091 | Sex*Diet: 1.759 | Sex: 0.2359 | Diet: 0.8177 |
|                  |         |      |           |      |           |     |      |
| %Infiltration    | 0.201 ± 0.131 | 0.150 ± 0.092 | 0.270 ± 0.143 | 0.000 ± 0.000 | Sex*Diet: 1.031 | Sex: 0.1433 | Diet: 2.241 |
|                  |         |      |           |      |           |     |      |
| **St lac mol**   |         |      |           |      |           |     |      |
| Density (cells/mm\(^2\)) | 248.8 ± 14.5 | 272.7 ± 14.2 | 239.4 ± 11.8 | 259.6 ± 15.0 | Sex*Diet: 0.01730 | Sex: 0.6481 | Diet: 2.507 |
|                  |         |      |           |      |           |     |      |
| Spacing Index (a.u.) | 0.473 ± 0.019 | 0.466 ± 0.004 | 0.442 ± 0.016 | 0.470 ± 0.022 | Sex*Diet: 1.105 | Sex: 0.6448 | Diet: 0.4185 |
|                  |         |      |           |      |           |     |      |
| Cluster          | 0.122 ± 0.062 | 0.179 ± 0.093 | 0.029 ± 0.029 | 0.147 ± 0.065 | Sex*Diet: 2.179 | Sex: 0.9011 | Diet: 1.739 |
|                  |         |      |           |      |           |     |      |
| %Infiltration    | 0.073 ± 0.073 | 0.437 ± 0.151 | 0.233 ± 0.154 | 0.216 ± 0.091 | Sex*Diet: 2.421 | Sex: 0.06403 | Diet: 2.006 |


Table 3
mHFD effects on microglial morphological parameters in the dorsal hippocampus CA1 stratum radiatum of P30 offspring. #: number, %: percentage on total myeloid cells, a.u.: arbitrary unit, CD: control diet, mHFD: maternal high-fat diet.

| Parameters                  | Mean ± standard error of the mean |   |   |   |   |
|-----------------------------|-----------------------------------|---|---|---|---|
|                            | Male CD                          | mHFD | Female CD | mHFD | Male mHFD | Female mHFD | Sex*Diet | Sex | Diet | P |
| Soma area (µm²)             | 46.59 ± 1.92                     | 46.88 ± 2.54 | 44.42 ± 1.38 | 45.82 ± 1.84 | 45.88 ± 2.54 | 44.42 ± 1.38 | 45.82 ± 1.84 | 0.08109 | 0.6758 | 0.1842 | 0.7795 | 0.4231 | 0.6735 |
| Arbor area (µm²)            | 1352.68 ± 154.93                 | 1355.63 ± 112.22 | 1374.23 ± 160.96 | 1377.16 ± 129.59 | 1335.63 ± 112.22 | 1374.23 ± 160.96 | 1377.16 ± 129.59 | 4.378 x 10⁻⁹ | 0.02340 | 0.0004357 | 0.7795 | 0.8803 | 0.9836 |
| Morphological index (a.u.)  | 0.036 ± 0.003                    | 0.035 ± 0.001 | 0.034 ± 0.003 | 0.035 ± 0.004 | 0.035 ± 0.001 | 0.034 ± 0.003 | 0.035 ± 0.004 | 0.09893 | 0.1851 | 0.05674 | 0.7795 | 0.7110 | 0.8148 |
| Cell area (µm²)             | 425.49 ± 86.24                   | 430.71 ± 55.12 | 473.40 ± 79.36 | 435.13 ± 49.86 | 425.49 ± 86.24 | 430.71 ± 55.12 | 473.40 ± 79.36 | 0.09819 | 0.1422 | 0.05674 | 0.7795 | 0.7110 | 0.8148 |
| Circularity (a.u.)          | 0.029 ± 0.001                    | 0.027 ± 0.001 | 0.030 ± 0.001 | 0.027 ± 0.001 | 0.029 ± 0.001 | 0.027 ± 0.001 | 0.030 ± 0.001 | 0.03614 | 0.7875 | 4.683 | 0.8516 | 0.3880 | 0.0459 |
| Solidity (a.u.)             | 0.281 ± 0.020                    | 0.290 ± 0.018 | 0.302 ± 0.012 | 0.285 ± 0.006 | 0.281 ± 0.020 | 0.290 ± 0.018 | 0.302 ± 0.012 | 0.7345 | 0.2907 | 0.07122 | 0.7930 | 0.5972 | 0.7900 |
| Aspect ratio (a.u.)         | 1.666 ± 0.055                    | 1.648 ± 0.053 | 1.633 ± 0.065 | 1.663 ± 0.069 | 1.666 ± 0.055 | 1.648 ± 0.053 | 1.633 ± 0.065 | 0.1587 | 0.02512 | 0.008885 | 0.7930 | 0.8761 | 0.9261 |
| # Branches                  | 101.92 ± 21.70                   | 104.47 ± 13.79 | 113.11 ± 17.14 | 102.23 ± 12.52 | 101.92 ± 21.70 | 104.47 ± 13.79 | 113.11 ± 17.14 | 0.1622 | 0.07200 | 0.06261 | 0.7919 | 0.7919 | 0.8056 |
| Average branch length (µm) | 3.318 ± 0.127                    | 3.218 ± 0.075 | 3.172 ± 0.071 | 3.237 ± 0.041 | 3.318 ± 0.127 | 3.218 ± 0.075 | 3.172 ± 0.071 | 0.9630 | 0.5690 | 0.04635 | 0.3410 | 0.4616 | 0.8323 |
| Longest branch (µm)         | 13.29 ± 0.24                     | 13.26 ± 0.49 | 12.77 ± 0.21 | 13.99 ± 0.21 | 13.29 ± 0.24 | 13.26 ± 0.49 | 12.77 ± 0.21 | 4.048 | 0.1145 | 3.750 | 0.6925 | 0.7924 | 0.7839 |
| # Junctions                 | 52.38 ± 12.05                    | 53.49 ± 7.28 | 58.43 ± 9.08 | 52.28 ± 6.78 | 52.38 ± 12.05 | 53.49 ± 7.28 | 58.43 ± 9.08 | 0.1616 | 0.07164 | 0.07779 | 0.6930 | 0.7924 | 0.7839 |
Table 4

mHFD effects on microglial morphological parameters in the dorsal hippocampus CA1 stratum lacunosum moleculare of P30 offspring. #: number, %: percentage on total myeloid cells, a.u.: arbitrary unit, CD: control diet, mHFD: maternal high-fat diet.

| Parameters                  | Mean ± standard error of the mean | F   | p     |
|-----------------------------|-----------------------------------|-----|-------|
|                             | Male CD                          | mHFD | Female CD | mHFD | Sex*Diet: | Sex: | Diet: |
| Soma area (µm²)             | 40.18 ± 1.74                     | 42.45 ± 2.06 | 40.14 ± 0.92 | 40.64 ± 1.57 | 0.2961 | 0.3230 | 0.7283 |
| Arbor area (µm²)            | 988.76 ± 125.44                  | 908.98 ± 111.46 | 974.04 ± 118.48 | 973.34 ± 157.09 | 0.09350 | 0.03685 | 0.09685 |
| Morphological index (a.u.)  | 0.043 ± 0.004                    | 0.049 ± 0.005 | 0.044 ± 0.006 | 0.047 ± 0.009 | 0.04385 | 0.009177 | 0.02427 |
| Cell area (µm²)             | 272.57 ± 46.35                   | 272.50 ± 36.57 | 269.99 ± 40.25 | 283.11 ± 43.67 | 0.02479 | 0.009177 | 0.02427 |
| Circularity (a.u.)          | 0.028 ± 0.003                    | 0.033 ± 0.003 | 0.031 ± 0.003 | 0.032 ± 0.006 | 0.2333 | 0.1162 | 0.6419 |
| Solidity (a.u.)             | 0.258 ± 0.012                    | 0.287 ± 0.008 | 0.263 ± 0.007 | 0.282 ± 0.013 | 0.2641 | 0.004823 | 5.616 |
| Aspect ratio (a.u.)         | 1.803 ± 0.089                    | 1.787 ± 0.058 | 1.793 ± 0.078 | 1.752 ± 0.094 | 0.02490 | 0.07486 | 0.1260 |
| # Branches                  | 72.83 ± 12.86                    | 73.24 ± 9.70 | 73.90 ± 11.91 | 77.18 ± 12.65 | 0.01466 | 0.04465 | 0.02438 |
| Average branch length (µm)  | 3.464 ± 0.077                    | 3.262 ± 0.048 | 3.420 ± 0.120 | 3.291 ± 0.037 | 0.2168 | 0.008423 | 4.553 |
| Longest branch (µm)         | 13.61 ± 0.46                     | 12.51 ± 0.55 | 12.96 ± 0.57 | 12.98 ± 0.61 | 1.024 | 0.02989 | 0.9634 |
| # Junctions                 | 36.88 ± 6.99                     | 37.01 ± 5.19 | 37.61 ± 6.37 | 39.29 ± 6.71 | 0.01468 | 0.05614 | 0.02034 |

Microglia from mHFD male offspring show increased interactions with astrocytes whereas both male and female offspring have decreased extracellular space pockets

To provide insights into microglial functions, we further performed SEM analysis to reveal, at nanoscale resolution, possible changes in their organelles and intercellular relationships in the dorsal hippocampus CA1 of mHFD- vs CD-exposed offspring, upon sacrifice at P30. We determined the number of microglial organelles involved in phagolysosomal activity (primary, secondary and tertiary lysosomes, lipofuscin, endosomes with or without content) and alterations to organelles that serve as markers of cellular stress (dilated cisternae of endoplasmic reticulum and Golgi apparatus, elongated
and total mitochondria). We also evaluated microglial interactions with their microenvironment, particularly direct contacts with astrocytic cell bodies, neuronal cell bodies, axon terminals, dendritic spines, oligodendrocytic cell bodies, myelinated axons, and blood vessels, as well as associations with extracellular space pockets containing degraded elements or debris indicative of extracellular digestion or “exophagy” [62, 63].

In the st rad, no significant difference in microglial organelles was observed between offspring groups, but there was a trend for a main diet effect regarding the total number of mitochondria per microglial cell body ($F_{(1,146)} = 3.870, p = 0.0511$) (Table 5). In terms of microglial interactions with their microenvironment, we detected a Sex*Diet interaction for the number of microglial contacts with astrocytes ($F_{(1,146)} = 0.0446$). Post-hoc analysis revealed that microglial cell bodies from mHFD-exposed male offspring made more cell-cell contacts with astrocytic cell bodies compared to CD male offspring ($p = 0.0182, 0.225 \pm 0.067 \text{ contacts vs } 0.054 \pm 0.038 \text{ contacts}$) (Table 5; Fig. 6a-e).

Microglial interactions with synaptic elements, myelinated axons, neurons as well as oligodendrocytes remained unchanged (Table 5). We also identified a Sex*Diet interaction for the prevalence of microglia-associated extracellular digestion ($F_{(1,146)} = 0.0433$), however, post-hoc analysis revealed no significant different between offspring groups (Table 5).

| Parameters       | Mean ± standard error of the mean | $F$     | $p$     |
|------------------|-----------------------------------|---------|---------|
|                  | CD                                | mHFD    | CD      | mHFD    | Sex*Diet: | Sex:  | Diet: |
| Organelles       | Male                             | Female  |         |         |           |       |       |
| # Lysosomes      | 1ary                             | 1.405 ± 0.461 | 1.150 ± 0.317 | 0.878 ± 0.213 | 1.438 ± 0.258 | Sex*Diet: | 1.552 | Sex: 0.1344 |
|                  | 2ary                             | 0.108 ± 0.052 | 0.200 ± 0.096 | 0.146 ± 0.075 | 0.188 ± 0.083 | Sex*Diet: | 0.1028 | Sex: 0.02644 |
|                  | 3ary                             | 0.027 ± 0.027 | 0.025 ± 0.025 | 0.000 ± 0.000 | 0.031 ± 0.031 | Sex*Diet: | 0.5148 | Sex: 0.5148 |
| # Lipofuscin     |                                  | 0.135 ± 0.079 | 0.100 ± 0.048 | 0.073 ± 0.041 | 0.063 ± 0.043 | Sex*Diet: | 0.04895 | Sex: 0.8091 |
| # Endosome       | Empty                            | 0.108 ± 0.052 | 0.200 ± 0.089 | 0.098 ± 0.058 | 0.063 ± 0.043 | Sex*Diet: | 0.9319 | Sex: 1.267 |

Table 5: mHFD effects on microglial ultrastructure in the dorsal hippocampus CA1 stratum radiatum of P30 offspring. #: number, 1ary: primary, 2ary: secondary, 3ary: tertiary. Astro: astrocyte, CD: control diet, Dil ER/golgi: dilated endoplasmic reticulum and Golgi apparatus cisterna, Elong: elongated mitochondria, mHFD: maternal high-fat diet, Mito: mitochondria, N/A: not applicable, Oligo: oligodendrocyte.
### In st lac mol, microglial organelle content and ultrastructure were unaffected by offspring groups (Table 6), but their interactions with the microenvironment differed. Similar to microglia in the st rad, microglial cell bodies in st lac mol of mHFD-exposed male offspring had increased interactions with astrocytic cell bodies ($F_{(1,128)} = 4.604, p = 0.0446$) compared to CD male offspring ($0.028 \pm 0.028$ contacts vs $0.214 \pm 0.094$ contacts) (Table 6; Fig. 6a-e, h-k, q). Of note, a significant main sex effect was also detected for microglial interactions with neurons ($F_{(1,128)} = 6.062, p = 0.0151$), where
microglial cell bodies from male offspring compared to female offspring made more cell-cell contacts with neuronal cell bodies, regardless of maternal diet (0.081 ± 0.026 contacts vs 0.000 ± 0.000 contacts) (Table 6; Fig. 6h-k, r). Microglial interactions with synaptic elements, oligodendrocytes and myelinated axons were also unchanged across sex and diet groups (Table 6). Lastly, mHFD-exposed offspring had a significant decrease ($F_{(1,128)} = 7.666, p = 0.0065$) in microglia-associated extracellular space pockets compared to CD offspring (0.1374 ± 0.0768 contacts vs 0.5349 ± 0.0206 contacts) (Table 6; Fig. 6l-o, s) in the two sexes.
Table 6

mHFD effects on microglial ultrastructure in the dorsal hippocampus CA1 stratum lacunsum moleculare of P30 offspring. #: number, 1ary: primary, 2ary: secondary, 3ary: tertiary, Astro: astrocyte, CD: control diet, Dil ER/golgi: dilated endoplasmic reticulum and Golgi apparatus cisterna, Elong: elongated mitochondria, mHFD: maternal high-fat diet, Mito: mitochondria, N/A: not applicable, Oligodendrocyte.

| Parameters | Mean ± standard error of the mean | F     | p     |
|------------|----------------------------------|-------|-------|
|            | Male CD | mHFD | Female CD | mHFD | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: | Sex*Diet: | Sex: | Diet: |
| Organelles | Lysosomes | 1ary | 1.556 ± 0.419 | 1.357 ± 0.258 | 1.229 ± 0.239 | 1.182 ± 0.202 | Sex*Diet: 0.06292 | Sex: 0.6902 | Diet: 0.1644 | Sex*Diet: 0.8023 | Sex: 0.4076 | Diet: 0.6858 | 2ary | 0.361 ± 0.090 | 0.250 ± 0.098 | 0.229 ± 0.092 | 0.212 ± 0.104 | Sex*Diet: 0.2389 | Sex: 0.7742 | Diet: 0.4337 | Sex*Diet: 0.6259 | Sex: 0.3806 | Diet: 0.5113 | 3ary | 0.028 ± 0.028 | 0.000 ± 0.000 | 0.000 ± 0.000 | 0.030 ± 0.030 | Sex*Diet: 1.817 | Sex: 0.003345 | Diet: 0.003345 | Sex*Diet: 0.1800 | Sex: 0.9534 | Diet: 0.9534 |
|              | Lipofuscin | # | 0.111 ± 0.053 | 0.107 ± 0.060 | 0.029 ± 0.029 | 0.061 ± 0.042 | Sex*Diet: 0.1493 | Sex: 1.918 | Diet: 0.09070 | Sex*Diet: 0.6999 | Sex: 0.1684 | Diet: 0.7638 | Empty | 0.194 ± 0.078 | 0.107 ± 0.079 | 0.229 ± 0.101 | 0.091 ± 0.051 | Sex*Diet: 0.09748 | Sex: 0.01231 | Diet: 1.945 | Sex*Diet: 0.7554 | Sex: 0.9118 | Diet: 0.1655 | Content | 0.361 ± 0.144 | 0.214 ± 0.079 | 0.171 ± 0.077 | 0.152 ± 0.063 | Sex*Diet: 0.3989 | Sex: 1.579 | Diet: 0.6886 | Sex*Diet: 0.5288 | Sex: 0.2113 | Diet: 0.4082 |
| Dil ER/golgi | 5.611 ± 0.788 | 7.643 ± 1.413 | 5.800 ± 0.805 | 7.512 ± 1.168 | Sex*Diet: 0.1075 | Sex: 0.02125 | Diet: 2.659 | Sex*Diet: 0.7436 | Sex: 0.8843 | Diet: 0.1054 | Mito | 0.472 ± 0.180 | 0.500 ± 0.159 | 0.257 ± 0.118 | 0.485 ± 0.235 | Sex*Diet: 0.3099 | Sex: 0.4110 | Diet: 0.05061 | Sex*Diet: 0.5787 | Sex: 0.5226 | Diet: 0.4781 | Elong | 3.111 ± 0.534 | 2.071 ± 0.430 | 2.400 ± 0.341 | 2.970 ± 0.543 | Sex*Diet: 2.856 | Sex: 0.03862 | Diet: 0.2435 | Sex*Diet: 0.8445 | Sex: 0.6225 | Diet: 0.6225 |
| Interactions with Synaptic microenvironment | Pre | 6.250 ± 0.745 | 5.821 ± 0.587 | 5.171 ± 0.527 | 5.125 ± 0.442 | Sex*Diet: 0.1168 | Sex: 2.138 | Diet: 0.1406 | Sex*Diet: 0.7331 | Sex: 0.1462 | Diet: 0.7083 | Post | 3.306 ± 0.378 | 3.036 ± 0.369 | 3.629 ± 0.482 | 3.030 ± 0.300 | Sex*Diet: 0.1726 | Sex: 0.1614 | Diet: 1.206 | Sex*Diet: 0.6785 | Sex: 0.6885 | Diet: 0.2742 | Myelinated axon | 0.417 ± 0.108 | 0.536 ± 0.167 | 0.486 ± 0.161 | 0.545 ± 0.151 | Sex*Diet: 0.04059 | Sex: 0.07164 | Diet: 0.3689 | Sex*Diet: 0.8406 | Sex: 0.7894 | Diet: 0.5447 | Degenerating myelin | 0.083 ± 0.047 | 0.179 ± 0.090 | 0.200 ± 0.069 | 0.121 ± 0.058 | Sex*Diet: 1.763 | Sex: 0.2047 | Diet: 0.01575 | Sex*Diet: 0.1867 | Sex: 0.6517 | Diet: 0.9003 |
| Contacts with brain cells | Astro | 0.028 ± 0.028 | 0.214 ± 0.094 | 0.114 ± 0.055 | 0.061 ± 0.042 | Sex*Diet: 4.604 | Sex: 0.3601 | Diet: 1.408 | Sex*Diet: 0.0338 | Sex: 0.5495 | Diet: 0.2376 | Neuron | 0.056 ± 0.039 | 0.107 ± 0.060 | 0.000 ± 0.000 | 0.000 ± 0.000 | Sex*Diet: 0.6095 | Sex: 6.062 | Diet: 0.6095 | Sex*Diet: 0.4364 | Sex: 0.0151 | Diet: 0.4364 | Oligo | 0.000 ± 0.000 | 0.036 ± 0.036 | 0.000 ± 0.000 | 0.000 ± 0.000 | Sex*Diet: 1.384 | Sex: 1.384 | Diet: 1.384 | Sex*Diet: 0.2417 | Sex: 0.2417 | Diet: 0.2417 | Blood vessel | 0.056 ± 0.039 | 0.143 ± 0.067 | 0.114 ± 0.055 | 0.152 ± 0.063 | Sex*Diet: 0.2002 | Sex: 0.3625 | Diet: 1.238 | Sex*Diet: 0.6553 | Sex: 0.5482 | Diet: 0.2679 | Extracellular space | 0.556 ± 0.176 | 0.214 ± 0.094 | 0.514 ± 0.180 | 0.061 ± 0.042 | Sex*Diet: 0.1533 | Sex: 0.4611 | Diet: 7.666 | Sex*Diet: 0.6961 | Sex: 0.4984 | Diet: 0.0065 | Extracellular digestion | 0.528 ± 0.216 | 0.321 ± 0.219 | 0.200 ± 0.090 | 0.364 ± 0.105 | Sex*Diet: 1.231 | Sex: 0.7334 | Diet: 0.01641 | Sex*Diet: 0.2693 | Sex: 0.3934 | Diet: 0.8983 |

Dark microglia and perivascular cells display increased number of dilated endoplasmic reticulum and Golgi apparatus cisterna in mHFD offspring.
Previously, our laboratory identified a microglial subset, the “dark microglia”, which are characterized by a distinct ultrastructural signature compared with typical microglia. These cells are found within the brain parenchyma, notably in the ventral/dorsal hippocampus CA1 st rad and st lac mol. Dark microglia exhibit several markers of cellular stress (dilatation of endoplasmic reticulum and Golgi, elongated mitochondria) as well as a dark, electron-dense cytoplasm and nucleoplasm [64]. These stressed microglia are rare in healthy mature mice, but become abundant in pathological conditions [64] including in a MIA mouse model induced with polyinosinic:polycytidulic acid (poly I:C) [49]. In the current study, we characterized the density and ultrastructure of dark microglia in the dorsal hippocampus CA1, st rad and st lac mol, comparing mHFD with CD offspring at P30. While imaging, we also noticed intriguing dark perivascular cells, localized inside the perivascular space yet displaying dark features similar to the dark microglia (i.e., dark, electron-dense cytoplasm and nucleoplasm, as well as markers of cellular stress). We further encountered apoptotic cells, identified by their dark cytoplasm, which was accompanied in this case by a distinctive pyknotic and fragmented nucleus. We decided to also quantify their density. The quantitative analysis of dark microglia and apoptotic cells revealed no significant difference in their density among the st rad and st lac mol of the adolescent offspring, regardless of their sex and maternal diet (Supplementary Table 1; Fig. 7a-c). Notwithstanding, more than half of the apoptotic cells we observed (four out of seven) were identified as microglia by their IBA1⁺ staining. In the st rad, dark perivascular cells also displayed a similar density between offspring groups. In the st lac mol, however, a sex difference was observed, with the female offspring showing a significantly increased density of dark perivascular cells \(F_{(1,12)} = 5.692, p = 0.0344\) compared to male offspring \(12.58 \pm 1.99 \text{ cell/mm}^2\) vs \(0.94 \pm 0.94 \text{ cell/mm}^2\) (Supplementary Table 1; Fig. 7d-e); regardless of maternal diet.

To study changes in organelles among the stressed dark cells in the st lac mol, we pooled together dark microglia and dark perivascular cells to obtain a sufficient sample size, required to be around 50 individual cells total for a large effect size (~ 0.4) (also see Supplementary Table 2 for the semi-descriptive analysis of the dark microglia and dark perivascular cells considered separately). This
quantitative analysis of dark cells revealed a main diet effect on their number of dilated endoplasmic reticulum or Golgi apparatus cisternae, which significantly increased \( (F_{(1,55)} = 4.264, p = 0.0437) \) in mHFD compared to CD offspring \((14.38 \pm 0.62 \text{ dilated cisterna} \text{ vs} \ 9.805 \pm 3.090 \text{ dilated cisterna)} \) (Supplementary Table 3; Fig. 7f-j). In addition, secondary lysosomes were significantly more abundant in female offspring compared to male offspring regardless of their maternal diet (Female offspring: \( 0.453 \pm 0.120 \text{ lysosomes} \text{ vs} \ 0.077 \pm 0.109 \text{ lysosomes} \) (Supplementary Table 3). This finding may describe a sex difference, regardless of maternal diet, in terms of dark cells phagolysosomal pathways. Across groups, the two types of stressed dark cells lastly displayed in the \textit{st lac mol} similar numbers of lysosomes, lipofuscin, endosomes and mitochondria, and their relationships with the microenvironment did not differ between groups.

Discussion

Our study investigated the effects of a fat enriched maternal diet on peripheral immune and microglial properties in the adolescent offspring of each sex. Characterization of our mouse model revealed a MIA phenotype defined by elevated circulating IL-6 in the mothers, together with increased fat deposition in the mothers and male offspring without other major metabolic changes. This MIA model led to peripheral immune priming demonstrated by exacerbated release of IL-6 upon a LPS-induced immune challenge in mHFD-exposed male and female offspring. The mHFD-induced MIA resulted in significant microglial morphological changes in the dorsal hippocampus CA1. We also found sexually dimorphic hippocampal transcriptomic changes, with mHFD-exposed male offspring showing reduced mRNA expression of the inflammatory-regulating cytokine \textit{Tgf1b} and microglial receptors \textit{Tmem119}, \textit{Trem2} and \textit{Cx3cr1}. In parallel to these changes, mHFD-exposed male offspring had increased microglial interactions with astrocytes in the dorsal hippocampus CA1, while both mHFD males and females had decreased microglia-associated extracellular space pockets in the same region. Taken together, this data highlights the emergence of a partially sex-dependent priming that lasted until adolescence in the mHFD offspring.

Previous work studying the metabolic consequences of mHFD in rodents have reported variable effects. With the same diet protocol that we used but in rats, Sasaki \textit{et al.} observed that weight
differences varied with the age of the offspring across their lifespan upon exposure to mHFD [13, 32]. mHFD offspring were significantly heavier during pubertal stages (P8-21) [13], while at adolescence (P35) [32] and adulthood (P90), their weight was similar to control animals [13]. Similarly, we did not find any significant difference of body weight in the P30 adolescent mouse offspring. By contrast, during early postnatal ages (P1 to P10), rat offspring exposed to lower fat mHFD (43% kCal/fat vs our 60% kCal/fat) became heavier with an increased fat mass [83], similar to our mHFD-exposed male offspring. In contrast to our observations, this study also reported heavier dams with lower blood glucose levels prior to mating [83], describing a diet-induced obesity model. Bilbo and Tsang compared two types of mHFD in rats – one with saturated fat and the other with trans-fat – with the same protocol duration used here, and found that both diets led to heavier dams [8]. In offspring, exposure to mHFD rich in saturated fats, and not trans-fats, led to significant weight gain in both sexes at puberty (P20) and in males at adulthood (P60) [8]. Overall, these studies suggest that the diet composition and exposure time may differently impact on the dams and offspring metabolism. In comparison, our mHFD mouse model did not result in severe endocrinologic alterations in the dams thus suggesting that the effects observed in our offspring were primarily due to the immunologic consequences of a fat enriched diet during pregnancy and nurturing, without confounding endocrine risk factors.

Few studies have examined the MIA induced by mHFD as well as the mechanisms underlying these MIA effects of mHFD. Using a macaque model of mHFD, Thompson et al. have noted an elevation of the pro-inflammatory cytokine IL-12 and a decrease of macrophage-derived chemokine (also known as C-C motif chemokine 22) in the blood of the mothers [9]. This study was the first to directly highlight the interplay among metabolic and inflammatory maternal changes resulting from the diet and their consequences on the offspring behaviors. In rodents, Bilbo and Tsang did not observe an increase in IL-6 in rat dams fed with a high saturated fat diet [8], contrary to our study. However, subsequent studies using a mouse model as well as a more sensitive approach involving multiplex-ELISA observed increased levels of circulating cytokines, including IL-6, in dams, during pregnancy [68] and the end of nurturing [14]. Discrepancy between Bilbo and Tsang with ours’ and others’
results may be explained by the difference in the sensitivity between the techniques. In addition, differences between the diets or species used could also underlie this discrepancy. Interestingly, pregnant female rat fed with a HFD had increased mRNA levels of \textit{Il-6} in the placenta [68], suggesting that IL-6 could cross the placental barrier to modulate fetal development [84] in our model.

In the mHFD-exposed offspring, alterations of the inflammatory status has notably been highlighted by increased cytokines levels (i.e. IL-5) in the blood of male mice at P7 and of female mice at P21 [14]. Moreover, another independent study has reported increased peripheral IL-6 four hours after a LPS immune challenge in adult rat male and female offspring [8]. In agreement with our results, these data suggest that during both adolescence and adulthood, mHFD-exposed offspring have a stronger response to a systemic immune challenge. IL-6 could contribute to inducing immune programming changes in the offspring by modifying transcriptional regulation of inflammatory mediators that are part of IL-6 downstream signaling target (i.e. \textit{AP1}, \textit{NFkB}, \textit{Sp1}) [85]. It is also possible that a stronger response to immune challenge occurs through increased release of cytokines by fat deposits. In the homeostatic central nervous system, Bilbo and Tsang reported an increase of protein levels of IL-1\textbeta without change of mRNA levels of \textit{Il-1}\textbeta and \textit{Il-6} in P20 pubertal and P60 adult offspring, from both sexes [8]. Another group, Sasaki \textit{et al.}, revealed greater mRNA levels of \textit{Il-6} in the adolescent hippocampus [32] and no difference compared to control in the adult hippocampus [13] using a similar diet protocol in male and female rats. In the present study, we did not detect \textit{Il-1}\textbeta or \textit{Il-6} mRNA, suggesting levels below threshold for detection by rt-qPCR or a lower efficiency of the primers used. Although Sasaki \textit{et al.} focused on an adolescent timepoint, they looked later, at P45. Therefore, it is plausible that the expression of inflammatory molecules –at protein and mRNA levels– varies between the two adolescent time points. It would be interesting to determine if adolescence represents a shifting or transition period for inflammatory signals, from a pubertal-like to an adult-like inflammatory phenotype.

In parallel to the changes of pro-inflammatory genes, neuroendocrine regulators genes (i.e. \textit{Gr}, \textit{Mr}, \textit{Nfkb}) were also reported to be differently expressed in the rat hippocampus across adolescence and adulthood [13, 32, 86]. These neuroendocrine receptors can modulate microglial functions such as
their release of inflammatory mediators (i.e. IL-1β, IL-6, TNF-α) during a LPS immune challenge [87–89], their morphology [90], cellular dynamics and proliferation [89]. Moreover, neuroendocrine receptors are differently expressed by microglia between the sexes [91]. In the current study we observed a sexual dimorphism of the inflammatory-regulating cytokine Tgfb1, with lower expression levels detected in whole hippocampus of mHFD-exposed male offspring compared to other offspring groups.

mHFD studies have mainly examined its effects on global inflammation and gene expression changes, without focusing on microglia. However, microglia may represent one of the key actors mediating the pathological consequences of mHFD during neurodevelopment. Bilbo and Tsang previously reported higher density of IBA1+ cells in the hippocampus CA1, CA3 and dentate gyrus of mHFD-exposed male and female rat offspring at adulthood [8]. In the present study, we did not observe any change in microglial density in the st rad and st lac mol of the dorsal hippocampus CA1. This could be explained by the method of analysis where microglia are identified by integrated density of IBA1+ staining among hippocampal regions (CA1, CA3, dentate gyrus) vs count of IBA1+/TMEM119+ cells in individual CA1 layers, together with the age or the species of animals used. Nevertheless, we observed microglial morphological changes in the st rad (circularity) and in st lac mol (solidity), as well as a decreased average branch length in the st lac mol. The reduced microglial circularity in the st rad of mHFD-exposed offspring suggests that these cells took a more elongated shape. Microglia with an elongated, rod-like morphology have been previously proposed to play a role in the response to acute brain insults due to their increased prevalence [92]. Elongated bipolar microglia were also previously described in a viral MIA model, in which ex vivo analyses also revealed that microglial chemotaxis and phagocytosis are increased in response to treatment (chemotaxis: CCL-2, IL-8, phagocytosis: LPS, poly I:C) [93]. The increased microglial solidity may represent a change in the distribution of the arborization, particularly a decreased distance between branches which would reflect a more convex-like shape. Others also hypothesized that an increase of the solidity value occurs during the morphological shift from a ramified to an amoeboid shape upon neuroinflammatory
insults, and represents a de-ramified or bushy morphology [94, 95], a phenotype seen upon exposure to stress [96]. In a bacterial MIA mouse model, microglia were previously shown to shift their morphology to an ameboid shape in adolescent (P40) offspring amygdala [97]. Although the microglial morphological changes we observed varied between layers, similarly they could indicate a general shift from a surveillant-ramified state to an ameboid shape. This morphological shift may also accompany microglial functional alterations in the hippocampus CA1 of mHFD offspring.

Microglia from the mHFD vs CD offspring interacted differently with their microenvironment. In st lac mol, the presence of extracellular space pockets surrounding microglial cell bodies decreased, which could be explained by various mechanisms. For instance, changes in the composition of the extracellular matrix could lead to a global decrease in extracellular space [98]. Microglia themselves were shown to contribute to modifying the extracellular matrix by releasing proteases (e.g. cathepsins, heparinases, metalloproteinases) [99, 100] that can promote their cell body migration or process motility, notably in inflammatory contexts [99, 100] or during brain plasticity [36]. Indeed, during visual deprivation in juvenile mice, the pockets of extracellular space surrounding microglia were shown to undergo reorganization [36] that was hypothesized to promote microglia motility and/or even dendritic spine structural plasticity and pruning [36, 101]. Hence, the decrease of extracellular space pockets-associated with microglial cell bodies that we observed could indicate a decrease in their motility and synaptic pruning function necessary for proper neuronal network maturation, which could be examined using ex vivo or in vivo imaging [98, 102].

Furthermore, these morphological and functional changes were associated with a decreased expression of microglial receptors (i.e. Cx3cr1, Tmem119, Trem2) in adolescent male offspring. TMEM119 is a protein considered to be expressed constitutively and exclusively by microglia in the brain [50]. Although its role remains under investigation, TMEM119 is required for microglial survival [75]. The absence of changes in the size of the microglial and infiltrating myeloid cells populations suggest no major issue with microglia survival in the st rad and st lac mol. Nevertheless, it is plausible that the turnover rate of microglia may change in the mHFD-exposed male offspring without affecting the population at a given time point. In this case, future investigation should consider examining
microglial density, but also proliferation and apoptosis. Further investigation into TMEM119 function would help to assess if this receptor is also involved in cell-cell interactions. Similar to TMEM119, TREM2 aids microglia to survive, in addition to mediating other functions including blood flow regulation [103], phagocytosis and synaptic pruning [76, 79-81]. CX3CR1 is also involved in phagocytosis and synaptic remodeling by microglia [77, 78, 104]. Hence, a decrease of Cx3cr1 and Trem2 mRNA expression may reflect reduced microglial interactions with neurons or synaptic elements as well as phagocytic activities, which was not observed at the ultrastructural level. It should be noted however that our ultrastructural analysis focused on microglial cell bodies, leaving the possibility that microglial processes make less contacts with synapses, which remains to be explored. Further investigation into the protein expression of Cx3cr1 and Trem2 is also required.

While the exact signaling pathway or receptor involved in microglia-astrocyte crosstalk has not been ascertained, a recent study suggested that microglial TREM2 may be involved. Indeed, using a TREM2 knockout mouse model, TREM2 deficiency led to a lack of microglia-astrocyte interactions in the cerebral cortex and hippocampus, which prevented the astrocytic-mediated phagocytosis of axon terminals [105]. The decrease of TREM2 we observed may represent a partial loss of microglial control over this newly defined function of astrocytes at synapses. Taken with the male-specific increase of microglial interactions with astrocytes that we observed in our ultrastructural analysis, these results stress the importance of determining the molecular crosstalk between microglia and astrocytes that is at play in mHFD-exposed male offspring. Characterization by in vivo imaging of astrocyte-axon terminals interactions while performing microglial modulation could allow to investigate the microglia-astrocyte-synapse relationships further.

Since transcriptomic analyses were performed on whole hippocampi, it remains possible that microglial survival, density and/or synaptic remodeling were altered in other regions and/or layers of the hippocampus beyond the CA1 st rad and st lac mol. Subsequent studies could characterize differences in other regions and layers, and also, how transcriptomic changes occur between sexes in the mHFD model. It would be important for instance to assess whether the mechanisms recruited in female involve differences in post-translational modifications and/or protein expression. Interestingly,
Glendining and Jasoni investigated epigenetic changes of the oxytocin receptor in the hippocampus of mHFD-exposed male and female mouse offspring by chromatin immunoprecipitation-qPCR (ChIP-qPCR), which revealed a sexually dimorphic pattern of acetylation and methylation leading to an increased expression of the oxytocin receptor in male offspring only [106]. Further investigation looking at whole genome epigenomic changes could hence provide, in parallel with proteomic analysis, better understanding of how microglial gene expression differs between sexes upon mHFD. Our laboratory has previously identified dark microglia as a subset of microglia associated with pathological conditions that are characterized by a dark cytoplasm and nucleoplasm without a clear chromatin pattern [49, 64]. Not only microglia can undertake a ‘dark’ appearance, but several other types of brain cells such as neurons, oligodendrocytes [53, 61] and astrocytes [107, 108] as well. Here, we not only observed dark microglia, but also stressed dark perivascular cells showing several signs of cellular stress (i.e. elongated mitochondria and dilated cisternae of the endoplasmic reticulum and Golgi apparatus) and dying apoptotic cells in the dorsal hippocampus CA1 of the P30 offspring. No difference in the density of stressed dark cells and apoptotic cells was observed with the maternal diet. However, an interesting observation we made was the increased density of stressed dark perivascular cells in the st rad of female offspring compared to male offspring from both maternal diets. Unfortunately, the low sample size of stressed dark cells and apoptotic cells did not allow us to perform quantitative analysis in the dorsal hippocampus CA1 st rad. Similar to previous observations by our team in adult pathological conditions, dark microglia were mainly found to be located in the st lac mol among the CA1 [64]. We thus assessed the global changes in stressed dark cells organelles without discriminating dark microglia from dark perivascular cells to increase our sample size. This analysis of dark microglia and dark perivascular cells revealed a significantly increased number of dilated endoplasmic reticulum and Golgi apparatus cisternae. Dilation of the endoplasmic reticulum and the Golgi apparatus are well characterized ultrastructural signs of oxidative stress [109]. The increase in their incidence thus reflects a higher level of cellular stress in the dorsal hippocampus CA1 of mHFD-exposed adolescent offspring. It would be interesting to investigate the cellular stress signature of the two cell types at the molecular level, and the effects of...
dark microglia as well as dark perivascular cells on the blood-brain barrier. In fact, increased stress within the neurovascular unit could alter blood-brain barrier integrity and permeability, as well as the regulation of blood flow. Consequently, a dysfunctional blood-brain barrier may leak into the brain peripheral inflammatory signals released upon mHFD, such as IL-6, thus contributing to the microglial changes observed.

Conclusion
Although mHFD induced similar phenotypes in both sexes for immune priming and microglial morphology, we identified sex-specific effects in the mHFD male offspring in terms of transcriptomic and functional ultrastructural changes during adolescence. Considering their key role in shaping brain neuronal networks, microglia in dorsal hippocampus may be partially responsible for several pathological neurodevelopmental outcomes described in mHFD models. Adolescence is a period of intense brain plasticity and maturation, where several microglia-mediated processes, such as synaptic pruning and myelination, are ongoing and may be profoundly impacted by the mHFD-driven microglial priming. Notably, synaptic pruning involving microglia-astrocyte crosstalk may be especially impacted negatively in mHFD-exposed male offspring. Investigation into these specific neurodevelopmental processes across the brain and the kinetics of their alterations during the offspring neurodevelopment, from embryonic stages to young adulthood, will be central to understanding the sexually dimorphic pathological cascade involving microglia.

List Of Abbreviation
AP-1: activator protein-1
Aif1: allograft inflammatory factor 1
a.u.: arbitrary units
ANOVA: analysis of variance
CA: cornu ammonis
CD: control chow
cDNA: complementary deoxyribonucleic acid
Cx3cr1: fractalkine receptor
DAB: diaminobenzidine
E: embryonic day
ELISA: enzyme-linked immunosorbent assay
HFD: high-fat diet
IBA1: ionized calcium-binding adapter 1
IL: interleukin
Il1ra: IL1 receptor antagonist
IkB: Nfkb inhibitor
LPS: lipopolysaccharide
mHFD: maternal HFD
MIA: maternal immune activation
MKP-1: mitogen-activated kinase protein 1
mRNA: messenger RNA
Nfkb: nuclear factor κ-light-chain-enhancer activator of B cells)
P: postnatal day
PB: phosphate buffer
PBS: phosphate buffered saline
PFA: paraformaldehyde
RNA: ribonucleic acid
RPL32: ribosomal protein L32
Rt-qPCR: real-time quantitative polymerase chain reaction
SAL: saline
SEM: scanning electron microscopy
Sp-1: specificity protein 1
St lac mol: stratum lacunosum moleculare
St rad: stratum radiatum
TBS: tris buffered saline
Tgfβ R: Transforming growth factor beta 1 receptor
TMEM119: transmembrane protein 119
TNF: tumor necrosis factor
Trem2: triggering receptor expressed on myeloid cells 2
W: week

Declarations
Ethics approval and consent to participate
All animal protocols were approved by McGill’s Facility Animal Care Committee under the ethical guidelines of the Canadian Council on Animal Care.

Consent for publication
Not applicable

Availability of data and materials
The data that support the findings of this study are available from the corresponding author, MET, upon reasonable request.

Competing interests
Not applicable

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Authors’ contributions
MB designed the initial experiments and coordinated most of the experiments. CL and NV helped with the optimization of RNA isolation and, NV gave precious guidance for rt-qPCR design and taught MB electron microscopy tissue processing. JCS helped with experimental design for ultrastructural imaging and analysis using SEM. MET provided constructive insights into experimental design and data interpretation throughout the study.
MB performed and analyzed all the physiological follow-ups of the pregnant mothers and offspring as well as multiplex-ELISA on the mothers. MB mainly performed tissue processing, and all data analyses. CL and LFC assisted with part of the animal care, the sacrifice of the offspring and performed multiplex-ELISA experiments on the offspring. CL and LFC discussed possible avenue to better answer the experimental questions of this study. JCS and FGI provided their expertise and imaged for ultrastructural analysis using SEM.

MB mainly wrote the manuscript under guidance from MET. All authors read, edited and approved the final manuscript.

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Figures
Figure 1

Experimental outline of HFD protocol on the dams and the offspring. Diet protocol period is identified by a pale blue-colored bar on the dams and the offspring timelines. At P21, all offspring were on normal chow diet before sacrifice at P30. Part of the animals received an intraperitoneal LPS injection 8 hours before characterizing their peripheral inflammatory profile. ♀: female, ♂: male, E: embryonic days, LPS: lipopolysaccharide, P: postnatal days, SAL: saline, W: weeks.

Figure 2

HFD effect on peripheral inflammatory profile in the dams at weaning of their litters. Plasma levels was measured by multiplex-ELISA for (a) IL-1β, (b) IL-17, (c) TNF-α, (d) IL-6 and (e) IL-10. Detection limit of each graphs is represented by a dotted line. Data are shown as mean ± standard error of the mean. p<0.01, p<0.01** by Mann-Whitney test. CD: control diet, HFD: high-fat diet, IL: interleukin, TNF-α: tumor necrosis factor α.
Figure 3

mHFD effect on plasma cytokine profile of P30 offspring eight hours after LPS treatment. (a) IL-1β, (b) IL-17, (c) TNF-α, (d) IL-6 and (e) IL-10 were measured by multiplex-ELISA in offspring plasma after administration of SAL or LPS by intraperitoneal injection. Dotted line on the graphs indicates the detection limits. (f) Radar graph illustrates cytokines levels in offspring from both diet groups regardless of their sex eight hours after LPS immune challenge. Data are shown as mean ± standard error of the mean. p<0.001***, p<0.0001**** (treatment) by mixed-effects analysis, p<0.05 # (treatment*diet), p<0.01 # # # # (treatment*sex) by mixed-effects analysis. CD: control, LPS: lipopolysaccharide, IL: interleukin, SAL: saline, mHFD: maternal high-fat diet, TNF-α: tumor necrosis factor α.
mHFD effect on microglia-related mRNA expression in the hippocampus of P30 offspring. (a) Scheme illustrating the region of interest, the hippocampus, in sagittal and coronal views of a mouse brain. mRNA expression ($2^{-\Delta\Delta Ct}$) was evaluated for Tgfb1 (b), Aif1 (c), Tmem119 (d), Trem2 (e) and Cx3cr1 (f) normalized by housekeeping gene Rpl32. Data are shown as $2^{(-\Delta\Delta Ct \text{ Mean} \pm \text{ standard error of the mean})}$. p<0.001**, p<0.001***, p<0.0001**** (sex*diet) by 2-way ANOVA followed by Bonferroni post-hoc test. Aif1: allograft inflammatory factor 1, a.u.: arbitrary units, CD: control, Ct: cycle threshold, Cx3cr1: fractalkine receptor, mHFD: maternal high-fat diet, Tgfb1: transforming growth factor β1, Tmem119: transmembrane protein 119, Trem2: triggering receptor expressed by myeloid cells 2.
mHFD effect on microglial density, distribution and morphology in the dorsal hippocampus
CA1 of P30 offspring. Representative scheme of the (a) st rad and (n) st lac mol to illustrate the layers analyzed. Immunofluorescence (IBA1+ low. Immunofluorescence IBA1 (red) and TMEM119 (green) allowed analysis of (b-e, o-r) microglial number and morphology (f-i, s-v).

Scale bar for the density pictures is equivalent to 50µm (b-e, o-r), whereas scale bar for morphology pictures is 25µm (f-i, s-v). Only main differences are presented on the figure: (j, w) microglial density, (k,x) circularity, (l, y) solidity and (m, z) branch length average. Data are shown as mean ± standard error of the mean. P<0.05* (diet) by 2-way ANOVA. ♀: Female, ♂: male, CD: control, Conv area: convex area, mHFD: maternal high-fat diet, Peri: perimeter.
mHFD effect on microglial ultrastructure in the dorsal hippocampus CA1 of P30 offspring.
Ultrastructural analysis of microglia was performed in the (a-g) st rad and (h-s) st lac mol. (a-d, h-k) Representative pictures of microglia as well as (l-o) higher magnification views of extracellular space pockets are provided. The interactions of microglia with their microenvironment differed between groups, here we present main results for microglial contacts with (e, q) astrocytes, (f, r) neurons and (g, s) extracellular space pockets. On the representative pictures, astrocytes, blood vessels, microglia and neurons are respectively identified by a ‘A’, by ‘BV’, by a ‘M’ and by a ‘N’. Presynaptic and postsynaptic elements are pseudo-colored respectively in green and orange. Mitochondria are pseudo-colored in purple and extracellular space pockets in yellow. Data are shown as mean ± standard error of the mean. P<0.05* (sex) by 2-way ANOVA, p<0.05 # (sex*diet) by 2-way ANOVA followed by Bonferroni post-hoc test. ♀: Female, ♂: male, CD: control, mHFD: maternal high-fat diet.
mHFD effect on dark perivascular cells, dark microglial cells and apoptotic cells in dorsal hippocampus CA1 of P30 offspring. (a) Dark perivascular cells, (b) dark microglia and (c) apoptotic cells examples are presented. Dark perivascular cell density was counted in the (d) st rad and the (e) st lac mol. Ultrastructural analysis of dark cells in the st lac mol revealed differences in their (f-j) dilation of the endoplasmic reticulum and Golgi apparatus cisternae. Dark processes are pseudo-colored in red. Presynaptic and postsynaptic elements
are pseudo-colored respectively in green and orange. Mitochondria are pseudo-colored in purple. Blood vessel are identified by ‘BV’. Dark perivascular cell, dark microglia and apoptotic cell are respectively identified using ‘DP’, ‘DM’ and ‘AC’. Dilatated endoplasmic reticulum and Golgi apparatus are identified by a white asterisk. Scale bar for representative picture of cell type is equivalent to 5µm while scale bar for picture showing endoplasmic reticulum and Golgi apparatus is equivalent to 500 nm. Data are shown as mean ± standard error of the mean. p<0.05* (sex) by 2-way ANOVA, p<0.05 # (sex*diet) by 2-way ANOVA followed by Bonferroni post-hoc test. ♂: Female, ♂: male, CD: control, Dilated ER/Golgi: dilatated endoplasmic reticulum and Golgi apparatus cisternae, mHFD: maternal high-fat diet.

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