Effects of high fat diet and maternal binge-like alcohol consumption and their influence on cocaine response in female mice offspring

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Significance Statement

Prenatal alcohol exposure is a leading cause of neurobehavioral and neurocognitive deficits, including eating disorders and increased risk for substance abuse as very common issues. Furthermore, eating conditions can also induce long-lasting modifications on the responses to drugs.

In this context, we considered of particular relevance to assess the interaction between alcohol exposure during gestation and lactation (PLAE) and a high fat diet (HFD) during childhood and adolescence of female offspring, and their consequences on nutrition-related parameters as well as on the response to the most consumed psychostimulant worldwide, cocaine.

Our results shed a light and suggest the existence of a crosstalk between the effects induced by a HFD and PLAE, so that there is a mutual attempt to compensate the effects induced by each condition (PLAE and HFD). However, further investigation would be needed to unveil what would occur with a longer exposure to a fatty diet.
Abstract

BACKGROUND: Prenatal alcohol exposure is a leading cause of neurobehavioral and neurocognitive deficits collectively known as fetal alcohol spectrum disorders (FASD), including eating disorders and increased risk for substance abuse as very common issues. In this context, the present study aimed to assess the interaction between alcohol exposure during gestation and lactation periods (PLAE) and a high fat diet (HFD) during childhood and adolescence.

METHODS: Pregnant C57BL/6 mice underwent a procedure for alcohol binge drinking during gestation and lactation periods. Subsequently, PLAE female offspring were fed with a HFD for 8 weeks and thereafter, nutrition-related parameters as well as their response to cocaine were assessed.

RESULTS: In our model, feeding young females with a HFD increased their triglyceride blood levels but did not induce an overweight compared to those fed with a standard diet. Moreover, PLAE affected how females responded to the fatty diet as they consumed less amount of food than water-exposed offspring, consistent with a lower gain of body weight. HFD increased the psychostimulant effects of cocaine. Surprisingly, PLAE reduced the locomotor responses to cocaine without modifying cocaine-induced reward. Moreover, PLAE prevented the striatal overexpression of cannabinoid 1 receptors induced by a HFD and induced an alteration of myelin damage biomarker in the prefrontal cortex, an effect that was mitigated by a HFD-based feeding.

CONCLUSION: Therefore, in female offspring, some effects triggered by one of these factors, PLAE or a HFD, were blunted by the other, suggesting a close interaction between the involved mechanisms.

Keywords: Drinking-in-the-dark; Alcohol; High fat diet; Cocaine; Female
1. Introduction

Prenatal alcohol exposure is a leading cause of significant neurobehavioral and neurocognitive deficits. This can contribute to increased risk for substance abuse among people with fetal alcohol spectrum disorders (FASD) (Grant et al., 2013). In this sense, Barbier et al. (2008) reported that alcohol-exposed male offsprings are more sensitive to the anxiolytic effect of ethanol, a feature that could partially explain the altered pattern of consumption of alcohol observed in these animals. Furthermore, it has been described that maternal binge-like alcohol consumption during gestation and lactation alters sensitivity to the reinforcing effects of cocaine and thus, enhances vulnerability to cocaine addiction in adult mice (Cantacorps et al., 2020).

Additionally, prenatal alcohol exposure increases the hypothalamic-pituitary-adrenal (HPA) axis tone, resulting in HPA dysregulation throughout life. This dysregulation is thought to affect energy homeostasis and eating behavior (Yau and Potenza, 2013). Nevertheless, the potential consequences of maternal alcohol exposure for eating behaviors and on nutritional issues of young offspring are relatively under-researched. In this regard, eating disorders seem to be common, whereby children with FASD show an altered food intake self-regulation. In this way, these children seem to prefer calorie-dense foods, which can increase obesity risk (Fuglestad et al., 2014; Amos-Kroohs et al., 2018). Indeed, rates of overweight and obesity are increased among children with FASD diagnosis, particularly in females (Werts et al., 2014). Altogether, evidence is suggestive of possible metabolic and/or endocrine disruptions in children with FASD. Furthermore, eating conditions, including body weight and type of food can also induce long-lasting modifications on the responses to drugs (therapeutic and recreational) (Baladi et al., 2010).

In this context, the present study aims to assess the interaction between alcohol exposure during gestation and lactation, and a high fat diet (HFD) during the female offspring’s childhood and adolescence, and their consequences on the psychostimulant-induced effects of cocaine (both behavioral and molecular) and on nutrition-related parameters.

It is important to highlight that the experimental model used in the present study intends to mimic as much as possible real-life exposition to these issues. Hence, C57BL/6 female mice were exposed to alcohol following the drinking-in-the-dark (DID) paradigm, during the entire gestational and lactational periods. Subsequently, prenatal and lactation alcohol exposed (PLAE) female offspring were fed ad libitum with a HFD for 8 weeks. Thereafter, they were
assessed for the psychostimulant and rewarding effects of cocaine using the horizontal locomotor test and the conditioned place preference (CPP) paradigm. In parallel, we investigated possible changes in some parameters related with neurotoxicity, cocaine response and its main target, the dopaminergic system, to explore how alcohol and HFD could influence the vulnerability to this psychostimulant in female mice. These changes were studied in specific brain areas: the ventral striatum (VS) and the prefrontal cortex (PFC), respectively. Additionally, nutrition-related parameters such as food intake, glucose and triglyceride blood levels, and the expression of genes induced by ghrelin were determined in the hypothalamus.

Cannabinoid CB1 receptor levels were also determined due to their implication in feeding regulation and drug abuse (Kirkham and Williams, 2001; Maldonado et al., 2006). The stimulation of CB1 receptors is a key component in the development of obesity, so that CB1 knockout mice are resistant to diet-induced obesity and antagonists of these receptors have anti-obesity potential (Di Marzo et al., 2001; Ravinet Trillou et al., 2004). On the other hand, endocannabinoids also play an important role in the modulation of synaptic plasticity in the dorsal striatum and NAcc (for reviews, see Freund et al., 2003; Gerdeman and Lovinger, 2003; and Piomelli, 2003). Moreover, dopamine agonists and psychostimulants increase the striatal release of endocannabinoids, suggesting that they could participate in the effects of psychostimulant drugs (Giuffrida et al., 1999; Patel et al., 2003; Centonze et al., 2004).

Our results point to a mutual interaction between some of the effects induced by PLAE and HFD, which deserves further investigation.

2. Materials and methods

2.1 Animals

Twelve-week-old male and female C57BL/6 inbred mice were purchased from Charles River (Barcelona, Spain) and shipped to our animal facility (UBIOMEX, PRBB) to be used as breeders. Upon arrival, they were housed in standard cages at constant temperature (21 ± 1°C) and humidity (55 ± 10%), under a reversed light-dark cycle (white lights on 20:00-08:00 h). After one week of acclimatization, breeding pairs were mated, and pregnant females were observed daily for parturition. For each litter, the date of birth was designated as PND 0. Pups remained with their mothers for 21 days and were then weaned (PND 21). After weaning,
male offspring were used for other experiments and female mice from each litter were randomly assigned to the different diets to avoid potential litter effect. All animal care and experimental procedures were approved by the local ethics committee (CEEA-PRBB) and conducted in accordance with the European Union Directive 2010/63/EU guidelines on the protection of animals used for scientific purposes. ARRIVE guidelines for reporting animal research have been followed.

2.2 Materials

Ethyl alcohol was purchased from Merck (Darmstadt, Germany) and diluted in tap water in order to obtain a 20% (v/v) alcohol solution for the Drinking-in-the-dark procedure. Cocaine hydrochloride was purchased in Alcaliber S.A. (Madrid, Spain) and prepared in 0.9% NaCl pH=7.4 (saline) immediately before administration.

[^3]H]Dopamine ([^3]H]DA) was from Perkin Elmer (Boston, USA). All the other reagents were of analytical grade and purchased from several commercial sources.

2.3 Experimental design

The experimental design and the four experimental groups involved in the study are depicted in Figure 1. Three different sets of animals were used in this study. The first one (N=106) had food intake, glucose and triglyceride blood levels measured. Then, the animals were subjected to the cocaine-induced conditioned place preference (CPP) that lasted for 10 days. Throughout the whole CPP process, the animals continued being fed with the corresponding diet. Afterwards, the animals which did not receive cocaine (N=20) were sacrificed to assess the expression of ghrelin-induced genes in the hypothalamus. The second set was used to assess the psychostimulant properties of cocaine (N=37), and the third to evaluate[^3]H]DA uptake (he mi-striatum) (N=33) and the expression of factors regarding dopaminergic neurotransmission (ventral hemi-striatum) and neuronal damage (frontal cortex) (N=20).

2.3.1 Drinking-in-the-dark (DID) procedure

Pregnant and nursing C57BL/6 mice were exposed to alcohol following the drinking-in-the-dark (DID) paradigm (Rhodes et al., 2005). The procedure was conducted as previously reported (Cantacorps et al., 2017; 2020), starting two days after mating. Pregnant female mice were randomly assigned to two groups: alcohol and water-exposed (control). Briefly, water bottles were replaced with 10-ml graduated cylinders fitted with sipper tubes
containing either 20% (v/v) alcohol in tap water or only tap water 3 h after the lights were turned off. Following a 2 h-access period, individual intake was recorded, and the original water drinking bottles were returned to the home cage. This procedure was repeated on days 2 and 3 and fresh fluids were provided each day. On day 4, alcohol or water cylinders were left for 4 h and fluid intakes were recorded. Fluid intakes (g/kg body weight) were calculated based on average 2-day body weight values, as dams were weighed at 2-day intervals. The procedure was maintained throughout the 3-week gestation and the 3-week lactation periods.

2.3.2. Feeding conditions

Two different types of diet were assigned to female offspring: a standard diet (SD, 831193 RM1; Special Diets Services, Essex, UK) and a high-fat diet (HFD, D05122301; Research Diets, Inc., USA). The SD had an energy density of 3.52 kcal/g (energy contribution from 75.1% carbohydrates, 17.49% protein and 7.42% fat). The HFD had an energy density of 4.73 kcal/g (35% carbohydrate energy, 20% protein energy and 45% fat energy). The fat source was composed of 91% hydrogenated coconut oil and 9% soybean oil.

On PND 29, female offspring were randomly divided in four experimental groups and assigned either a SD or a HFD (Figure 1B). Animals were fed with one of both diets from the weaning up to sacrifice. All behavioral experiments were carried out during the youth of the animals (PND 85-105) (Flurkey et al., 2007). The weight of the animals and the food intake were monitored weekly during the whole feeding period. The weight gain throughout the feeding period was calculated for each animal as the difference of weight between the last and the first week of diet exposure, expressed in grams.

2.3.3. Glucose and triglyceride determination

Fasting triglycerides and glucose levels were measured following 8 weeks of SD or HFD feeding, immediately before the conditioned-place preference test (CPP) (Set 1). In brief, mice were fasted for 6 h, and blood samples were extracted from the tail between 3 pm and 4 pm. The tests were performed in a quiet room using an appropriate measuring device (Accutrend Plus® system Cobas, Roche, Spain).
2.3.4. Behavioral Tests

Cocaine-induced conditioned place preference (CPP)

The CPP procedure consisted of three different phases: preconditioning, conditioning and testing day (Luján et al., 2019). During preconditioning, mice could freely explore both compartments for 20 min. The conditioning phase consisted of four pairings: mice received an i.p. injection of 15 mg/kg cocaine immediately prior to confinement to the drug-paired compartment for 30 min on days 2, 4, 6 and 8, while on alternate days (3, 5, 7 and 9) mice received physiological saline before being confined to the vehicle-paired compartment for 30 min. The time spent in each compartment during the preconditioning and testing sessions, as well as the distance travelled, were recorded by computerized monitoring software (CIBERTEC APL software). A CPP score was calculated for each subject as the difference between the time spent in the drug- paired compartment during the testing and the preconditioning sessions.

Cocaine-induced locomotor activity

After a 2-day habituation period (day -2), all mice (set 2) received a single dose of cocaine (8 mg/kg i.p.) and were immediately placed into the open field arena where their horizontal locomotor activity (HLA) was recorded by a computerized monitoring software (Smart 3.0 Panlab, Barcelona, Spain) for 30 min.

2.3.5. Molecular Determinations

Tissue samples preparation

Mice were sacrificed by cervical dislocation. The whole striatum, ventral striatum, prefrontal cortex or hypothalamus, when appropriate, were quickly dissected out and, except the whole striatum, they were stored at -80 °C until use.

For the [3H]DA uptake experiments, mice synaptosome from fresh hemi-striatums were prepared as described by Pubill et al. (2005).

Total protein extracts were isolated from ventral hemi-striatum and prefrontal cortex and processed as described by Pubill et al. (2013), with minor modifications. Briefly, tissue
samples were thawed and homogenized through sonication at 4°C in 20 volumes of lysis buffer (20 mM Tris-HCl, 1% NP40, 137 mM NaCl and 2 mM EDTA, pH=8) containing a protease and phosphatase inhibitor cocktail. Thereafter, the homogenates were shaken and rolled for 2 h at 4°C and subsequently centrifuged at 15,000 x g for 30 min at 4 °C. Protein content of the supernatants was determined using the Bio-Rad Protein Reagent (Bio Rad, Inc. Spain).

[^3]H]DA uptake in striatal synaptosomes

Reaction tubes were composed of 25 µl of the radioligand [^3]H]DA (final concentration 5 nM), 100 µl of the synaptosome suspension and 125 µl of Hank’s HEPES-buffered solution containing pargyline (20 mM) and ascorbic acid (1 mM). The incubation was performed for 5 min at 37 ºC. Uptake reactions were terminated by rapid vacuum filtration through Whatman GF/B glass fiber filters (Whatman Intl Ltd, Maidstone, UK) pre-soaked with 0.5% polyethyleneimine. Tubes and filters were washed three times with ice-cold 50 mM Tris-HCl. The radioactivity trapped on the filters was measured by liquid scintillation spectrometry. Non-specific uptake value was determined in parallel samples containing cocaine (final concentration 300 µM) at 4 ºC and was subtracted from total uptake to yield specific uptake.

Western blot analysis

A general Western blotting protocol was used as described by Duart-Castells et al. (2019), with minor modifications. Membranes were incubated overnight at 4 ºC with anti CB1 receptor (1:1000, Frontiers Institute, South Africa), MBP (1:1000, Abcam, Cambridge, UK), NeuN (1:10000 Abcam, Cambridge, UK), NFκB (1:1000, Cell Signaling Technology, Inc) or TH (1:5000, Transduction Laboratories, Lexington, USA) primary antibodies. After washing, membranes were incubated for 1 h at room temperature with their respective secondary peroxidase-conjugated anti-IgG antibody: donkey anti-rabbit, sheep anti-mouse or goat anti-rat (1:5000, GE Healthcare, USA). GAPDH (1:5000, Merck Millipore, USA) antibody was used as a control for loading.

Total RNA extraction and gene expression determination

RNA extraction and quantitative RT-PCR were performed from hypothalamus as described by Mir et al., (2018). The sequences of the primers used for each gene were 5'-TGCAGACCGAGCAGAAGAAG (forward) and 5'-GACTCGTGCGAGCCTTACACA (reverse) for AgRP; 5'-TATCTCTGCTCGTGTTTG (forward) and 5'-
GTTCTGGGGCGTTTTCTG (reverse) for Npy; and 5'- TATGCAGTCGCCCTTCCT (forward) and 5'- ACATCAATCAGGTGTGTCTGCT (reverse) for Cpt1c.

2.4 Data acquisition and statistical analysis

Data are expressed as mean ± SEM. Data from biochemical analysis (Western blot and [³H]DA uptake experiments) were normalized with 100% defined as the mean of the technical replicates in the control group (W-SD). In qPCR analysis, data were expressed as fold-change variations relative to W-SD.

Differences between groups were compared using one-way or two-way ANOVA with diet (SD or HFD) and maternal exposure (water or alcohol) as factors of variation. To analyze the food intake and cocaine induced CPP, a three-way ANOVA was performed, with diet (SD or HFD), mother’s exposure (water or alcohol) and time (day or week), as factors of variations. The α error probability was set at 0.05. Significant differences (P < 0.05) were analyzed using the Bonferroni post-hoc test for multiple comparison measures only if F achieved the necessary level of statistical significance (P < 0.05) and no significant variance in homogeneity was observed. Statistical calculations were performed using GraphPad Prism 8.0 software.

3. Results

3.1 Maternal alcohol consumption

Two-way ANOVA with repeated measures analysis of the volumes of water and alcohol consumed during DID testing showed a significant effect of day [F(23,1173) = 27.693; P < 0.001] and alcohol [F(1,51) = 66.795; P < 0.001], with interaction between factors [F(23,1173) = 10.482; P < 0.001] (Figure 2A). Bonferroni post-hoc comparisons revealed a significant increase in water consumption on day 4 compared with day 1 (P < 0.001), day 2 (P < 0.001) and day 3 (P < 0.001), on day 8 compared with day 5 (P < 0.05), day 6 (P < 0.05) and day 7 (P < 0.01), on day 16 compared with day 14 (P < 0.05) and on day 24 compared with day 21 (P < 0.05), day 22 (P < 0.01) and day 23 (P < 0.05). Furthermore, a significant increase in alcohol consumption on day 4 compared with day 1 (P < 0.01) and day 2 (P < 0.05), on day 8 compared with day 5 (P < 0.05) and on day 24 compared with day 22 (P < 0.01) was found. Additionally, one-way ANOVA with repeated measures analysis of alcohol
intake showed a significant effect of day $[F_{(23,575)} = 7.389; P < 0.001]$ (Figure 2B). Bonferroni post-hoc comparisons revealed a significant increase in alcohol intake on day 4 compared with day 1 ($P < 0.01$) and day 2 ($P < 0.05$), on day 8 compared with day 5 ($P < 0.05$) and on day 24 compared with day 22 ($P < 0.01$).

### 3.2 Body weight and food intake

Initially, the body weight after weaning was not affected by alcohol exposure $[F_{(3,85)} = 1.241; \text{n.s.}]$. From that moment on, all groups of animals gained weight over time across the 8-week exposure period. Two-way ANOVA of the weight gain revealed a significant effect of the alcohol factor $[F_{(1,100)} = 14.96; P < 0.001]$ but no effect of the diet $[F_{(1,100)} = 0.451; P > 0.05]$. Accordingly, no significant differences in weight gain between the groups fed with HFD vs. those fed with SD were observed. Bonferroni post-hoc multiple comparisons revealed that the weight gain of the A-HFD group was significantly lower than that of W-HFD. Conversely, no differences were observed in weight gain between animals fed with SD, whether exposed to water or alcohol (Figure 3A).

Weekly food intake, calculated either as an average of grams/day or Kcal/day, was also measured. Three-way ANOVA ($alcohol \times diet \times week$) with repeated measures of grams/day ingested yielded a significant effect of diet $[F_{(1,100)} = 305.5; P < 0.001]$ and time $[F_{(7,651)} = 68.85; P < 0.001]$, with interaction between $alcohol \times diet$ $[F_{(1,100)} = 6.034; P < 0.05]$ and $time \times diet$ $[F_{(7,651)} = 62.20; P < 0.01]$. Subsequent Bonferroni post-hoc tests revealed significant differences in food intake between SD and HFD groups in mice exposed to alcohol ($P < 0.001$). Furthermore, significant differences between SD and HFD levels of food intake were found on weeks 1, 2, 3, 6, 7 and 8 ($P < 0.001$; in all cases). Overall, mice receiving HFD ingested less grams of diet than those fed with SD.

Regarding the weekly food intake in Kcal/day, three-way ANOVA analysis ($alcohol \times diet \times week$) with repeated measures showed a significant effect of time $[F_{(7,651)} = 74.44; P < 0.001]$ and diet $[F_{(1,100)} = 27.86; P < 0.001]$, with significant interactions between $time \times diet$ $[F_{(7,651)} = 68.85; P < 0.001]$ and $alcohol \times diet$ $[F_{(1,100)} = 6.36; P < 0.05]$ (Figure 3B). Bonferroni post-hoc comparisons indicated that, from the second week of diet consumption until the sixth, the HFD-groups (water and alcohol) significantly ingested more Kcal/day than SD-fed mice ($P <$
0.001; in all cases). Nevertheless, from week 6 until the end, these groups reduced their Kcal intake until being equated to that of the SD-fed groups.

In accordance with the significant interaction found (alcohol x diet), maternal exposure to alcohol did not modify the food and Kcal intake when female offspring were fed with SD but it did when fed with HFD.

3.3 Glucose and triglyceride blood levels

Two-way ANOVA analysis of triglycerides blood levels showed a significant effect of the diet \( [F_{(1,20)} = 13.07; P < 0.01] \). Thus, HFD exposure significantly increased triglycerides blood levels in comparison to SD (Figure 3C). By contrast, neither the diet nor the maternal alcohol exposure altered the glucose blood levels by the end of exposure (Figure 3D).

3.4 Basal locomotor activity and cocaine-induced CPP

The first set of mice were finally subjected to the cocaine-induced conditioned place preference. Firstly, two-way ANOVA of the results did not show any significant effect of the diet nor alcohol exposure on the distance travelled by the animals in the apparatus, so no changes in basal locomotor activity were found between the different experimental groups (Figure 4, inset).

Regarding the CPP experiments, three-way ANOVA analysis (alcohol x diet x day) revealed a significant effect of cocaine treatment \( [F_{(1,78)}=93.314; P < 0.001] \), indicating that the repeated administration of cocaine (15 mg/kg, i.p.) produced a preference for the cocaine-paired compartment in all groups (Figure 4). However, no effect of alcohol \( [F_{(1,78)}=0.913; P > 0.05] \) or the diet \( [F_{(1,78)}=0.496; P > 0.05] \) was found.

3.5 Cocaine effect on locomotor activity

When the acute effect of cocaine was assessed in the second set of mice, an interesting result was obtained. Two-way ANOVA yielded a significant influence of the interaction alcohol x diet \( [F_{(1,34)}=7.12; P = 0.01] \). As shown in Figure 5A, an acute administration of cocaine (8 mg/kg i.p.) elicited a similar effect in all groups of animals, but an increased response in the W-HFD group compared to the W-SD (\( P < 0.05 \)).
3.6 Molecular determinations

In order to find a possible explanation to the differences observed in food intake, the expression of genes induced by ghrelin was determined in the hypothalamus (set 1 of animals). In parallel, as an attempt to explain the results observed from the cocaine-induced psychostimulation, $[^3]$H]DA uptake in striatal synaptosomes (set 2) and the expression of tyrosine hydroxylase (TH) and cannabinoid 1 receptor (CB1) in VS were evaluated (set 3). Finally, the expression levels of proteins involved in neuronal and myelin damage were measured in the PFC (set 3) in view of previous studies using the male offspring from mice subjected to the DID procedure (Cantacorps et al., 2017): NeuN, nuclear factor κB (NF-κB) and myelin basic protein (MBP).

3.6.1. Effects on ghrelin-induced genes

In the hypothalamus, we determined by qPCR the mRNA levels of genes that are induced by ghrelin and promote food intake: neuropeptide Y (NPY) (Figure 3E) and Agouti-related protein (AgRP) (W-SD: 1.000 ± 0.31, A-SD: 1.193 ± 0.425, W-HFD: 0.873 ± 0.191, A-HFD: 1.226 ± 0.409), and no significant changes were observed (N=5/group). We also measured the mRNA levels of CPT1C as a mediator of ghrelin signaling, showing reduced levels in the alcohol-exposed groups which were very close to reach statistical significance [$F_{(1,16)}= 4.263; P = 0.055$] (Figure 3F).

3.6.2 Effects on $[^3]$H]DA uptake

Striatal synaptosome suspensions were prepared to perform $[^3]$H]DA uptake experiments. Two-way ANOVA yielded a significant effect of diet [$F_{(1,29)}=16.28; P < 0.001$]. Accordingly, mice exposed to HFD presented lower DA uptake activity than those fed with SD (P < 0.01 and P < 0.05 vs. the water and alcohol-matching groups, respectively) (Figure 5B).

3.6.3 Effects on TH and CB1 receptor expression in ventral striatum

As shown in Figure 5C, the expression of TH was not affected neither by alcohol nor by the diet. By contrast, two-way ANOVA of the results of CB1 expression yielded a significant effect of the diet [$F_{(1,16)}=6.576; P < 0.05$] and alcohol [$F_{(1,16)}=21.14; P < 0.001$] with interaction between both factors [$F_{(1,16)}=10.27; P < 0.01$] (Figure 5D). Bonferroni post-hoc multiple comparisons showed a significant increase in CB1 receptor expression in W-HFD.
mice compared to W-SD (P < 0.01). However, such overexpression was not present in animals fed with HFD and exposed to alcohol. (Pictures of the whole immunoblot membranes can be seen as supplementary materials).

3.6.4. Neuronal and myelin damage within the prefrontal cortex

In our study, using female offspring, no differences were observed in NeuN or NFκB/p65 expression (Figure 6A and 6B). However, two-way ANOVA of the results of MBP expression revealed a significant effect of alcohol [F(1,16)=5.000; P < 0.05] (Figure 6C). Therefore, alcohol exposure significantly decreased MBP protein levels, mainly in animals fed with SD (P=0.05).

4. Discussion

The present study aimed to assess the interaction between alcohol exposure during gestation and lactation and a high fat diet during the female offspring’s childhood and adolescence on some nutrition-related parameters as well as on the response to cocaine.

In our study, there was no contribution of PLAE to offspring’s body weight either in agreement with previous studies from our group (Cantacorps et al., 2017). Moreover, we did not observe any significant weight gain in mice fed with HFD with respect to the SD group accordingly with previous studies (Hwang et al., 2010; Sims et al., 2013; Hicks et al., 2016). When food intake was measured, HFD-fed mice reduced the grams of ingested food from week 6 until the end of the exposure, thus equaling their Kcal/day intake to those of the SD-group. Importantly, the equal intake of Kcal is in line with the non-significant differences observed in body weight between W-HFD and W-SD mice at the end of the experiment. Lin et al. (2000), using a similar HFD-feeding protocol, showed that HFD-induced obesity in mice could be divided into early (1 week), middle (until 8 weeks) and late stages. Mice fed for 8 weeks with HFD showed a very slight increased body weight, but energy intake fell below control levels during this period, which began to increase gradually after 8 weeks. In addition, previous studies have described that male C57BL/6J mice on HFD are more susceptible to weight gain than females (Gelineau et al., 2017) and weight gain appears earlier in males (46-days old) than in females (129-days old) (Hwang et al., 2010). Overall, we can reasonably deduce that, in our model, female offspring were probably in a pre-obesity state and thus, they needed a longer exposure period on the HFD (more than 16-20 weeks) (Wang and Liao, 2013) to develop an obesity status.
TG and glucose blood levels were also assessed. The HFD significantly increased TG blood levels, but not those of glucose. It is already known that C57BL/6 mice, even when fed with high sucrose diets (65%), do not develop insulin resistance (Gajda et al., 2007). Nonetheless, female C57BL/6J mice are particularly resistant to the physiological changes caused by HFD, and they exhibit a better glucose tolerance than males (Gelineau et al., 2017). Although we used a strain of C57BL/6 mice with a slightly different genetic background (from the Charles River labs) and strain-dependent differences could not be completely ruled out, our results about glucose levels and metabolic effects are in line with those previously reported by other groups using the J strain. Importantly, no additional effects of PLAE were evidenced on TG and glucose blood levels.

Additionally, PLAE affected how C57BL/6 mice responded to the HFD in such a way that, during almost the whole feeding-period, A-HFD mice ate less grams of food, and thus, less Kcal, than the W-HFD, consistent with their lower weight gain. Indeed, similar results were observed by Amos-Kroohs et al. (2018). Importantly, such lower weight gain cannot be attributed to an increased basal locomotor activity of these animals, since no differences were observed in their basal locomotion. However, regarding the effects of PLAE and HFD on ghrelin-induced genes, although no changes were observed in NPY or AgRP mRNA expression, an apparent decrease of the CPTC1 expression in A-HFD mice should be noted. Hypothalamic CPT1C mediates the central effects of leptin and ghrelin on feeding behavior (Gao et al., 2011; Ramírez et al., 2013). More specifically, besides other mechanisms, ghrelin induces food intake through regulation of hypothalamic CPT1C, so the orexigenic action of ghrelin is totally blunted in CPT1C knockout mice (Ramírez et al., 2013). Accordingly, aside from the metabolic change induced by HFD, which caused an overall decrease in food intake, PLAE seemed to additionally induce an apparent decrease in CPTC1 expression, a feature that might be involved in the lower weight gain observed in the A-HFD animals.

FASD is associated with a higher risk of later developing drug abuse. Our results showed that PLAE and/or a HFD did not modify CPP acquisition, thus the rewarding effects of cocaine were not altered. Consistent with our findings, Blanco-Gandía et al. (2017) suggested that consumption of a HFD during adolescence in male mice induces neurobiochemical changes that increased sensitivity to cocaine but only when fat is withdrawn, acting as an alternative reward. Moreover, locomotion induced by an acute dose of cocaine was assayed as an indicative of its psychostimulant effect. Data in rats indicate that the consumption of fat and, perhaps, the resulting hormonal changes, markedly alter dopamine systems (Baladi et al.,
In agreement with Collins et al., (2015), we observed an increased sensitivity to the psychostimulant effect of acute cocaine in animals fed with HFD. This is of chief importance since it provides evidence that consuming a HFD during early development enhances the psychostimulant effect of cocaine, and thus, might favor and increase the probability of repeating and perpetuating the use of the substance, ultimately leading to its abuse. Surprisingly, the A-HFD mice did not show the increased response to the first dose of cocaine when compared with the SD group as the W-HFD group did.

It is known that reduced phasic dopamine release and slowed dopamine uptake occur in the nucleus accumbens after a HFD (Barnes et al., 2020). In our study, when assessing [3H]DA uptake in striatal synaptosomes we found out that the HFD reduced the overall [3H]DA uptake in both, A-HFD and W-HFD groups implicating diet in the regulation of DA function, which can lead to functional modifications in DA signaling (South and Huang, 2008; Cone et al., 2013). By decreasing DA uptake, HFD consumption could promote adaptions such as downregulation of DA receptors, a feature of both human and rodent models of obesity (Wang et al., 2009; Johnson and Kenny, 2010). Cone et al. (2013) described that prolonged HFD appears to reduce DAT trafficking or perhaps maturation, but not DAT gene expression. Therefore, diet-related decreases in membrane DAT could precede and contribute to the onset of DA receptors downregulation, obesity and compulsive eating behavior that develop over the course of HFD consumption (Johnson and Kenny, 2010).

Cocaine increases the motor activity by facilitating DA signaling via the mesostriatal pathway through both DAT- dependent and independent mechanisms (Gardner and Ashby, 2000). Given the effects on DA uptake, it seems reasonable that the W-HFD group elicited an enhanced psychostimulant response to an acute dose of cocaine, as more DA would be available in the synapses and the proportion of DAT blocked by the same dose of cocaine would be higher. However, the A-HFD mice did not show enhanced hyperlocomotion by cocaine. Several mechanisms can be hypothesized to explain such difference: probably alcohol reduced the cocaine-induced hyperlocomotion in the HFD group by other mechanisms than DA synthesis or DA uptake, which could not be explained in this study, but correlates with other parameters induced by HFD but dampened by PLAE. Another possible cause may be the increased CB1 receptors, which appeared only in the W-HFD group and play an important role in dopamine transmission, as discussed below. Finally, an increased sensitivity of postsynaptic dopamine receptors as a result of CB1 increase or any other undetermined mechanism cannot completely be ruled out.
We also assessed two proteins related to dopaminergic neurotransmission in the ventral striatum: TH and CB1 receptor. In our model, the expression of the rate-limiting enzyme in DA synthesis, TH, was affected neither by alcohol nor by the diet. By contrast, and as observed for the acute response to cocaine, HFD induced an increase in CB1 receptors but only in the animals not exposed to alcohol. Stimulation of CB1 receptors is a key component in the development of diet-induced obesity, in such a way that CB1 receptor knockout mice display an hypofagic behavior and reduced body weight (Ravinet Trillou et al., 2004). The fact that the A-HFD group in our study did not present an increase in CB1 receptors probably contributed to a reduced predisposition to obesity due to early alcohol exposure.

There are functional interactions between endocannabinoid and dopaminergic systems that may contribute to striatal signaling. More concretely, the endogenous cannabinoid system is a key regulatory element of the plastic changes that are associated with cocaine-induced behavioral responses in the rat. Corbillé et al. (2007) investigated the role of CB1 receptor in the effects of a single injection of psychostimulants and found that locomotor responses to cocaine and D-amphetamine were decreased in CB1 receptor-deficient mice. Their results provided strong evidence for the role of the endocannabinoid system in regulating neuronal circuits critical for cocaine effects, as ERK phosphorylation, presumably by acting on CB1 receptors located on terminals of striatal medium spiny neurons. Additionally, Gessa et al. (1998) used CB1−/− mice with a C57BL/6J genetic background to further investigate the role of CB1 receptors in cocaine action. CB1−/− mice displayed a significant reduction in cocaine-enhanced locomotion related to a reduction in DA release. Accordingly, pharmacological blockade of CB1 receptors inhibited cocaine-induced hyperlocomotion and DA release in CB1+/+ mice. All these findings suggest an important role for CB1 receptors in mediating the psychostimulant effects of cocaine. Therefore, an increase of CB1, as seen in W-HFD group, combined with the reduced DA uptake could contribute to the enhanced hyperlocomotion induced by this psychostimulant.

Recent studies have demonstrated that alcohol intake activates the innate immune system in the central nervous system, leading to neuroinflammation and contributing to brain damage and behavioral dysfunctions (Montesinos et al., 2016). In this context, we have already reported that PLAE increases pro-inflammatory markers, alters the expression of myelin proteins, and induces neuron cell damage in the PFC of male C57BL/6 mice (Cantacorps et al., 2017). Reduction of MBP levels in certain brain areas after alcohol exposure during development has also been described by other research groups (Bichenkov and Ellingson,
2009). In the present study, effects of PLAE were evident in MBP protein expression, in such a way that alcohol exposure induced a decrease in MBP. Regarding NeuN expression, although a significant decrease was found in A-SD mice compared to W-SD ($t_7 = 2.557, P < 0.05$), such difference was blunted until being non-significant when considering also the variable diet in the analysis. Therefore, we could confirm that the effects of PLAE on NeuN, and MBP expression were the same in both sexes albeit female offspring seemed more resistant to such effects than males. Furthermore, all the alterations induced by PLAE were blunted in HFD groups, thereby a short exposure to a fatty diet seems to counteract the deleterious effects of prenatal and lactational alcohol exposure. It could be hypothesized that the fatty diet would supply extra lipids that could overcome the deleterious effects of alcohol on myelinization.

In summary, in our model, feeding young female mice with a HFD for 8 weeks increased their TG blood levels but did not induce an overweight compared to those fed with SD. Moreover, PLAE affected how females responded to the fatty diet as they consumed less amount of food, consistent with their lower weight gain. Regarding their response to cocaine, the HFD reduced DA uptake and increased the acute hyperlocomotion induced by the drug without modifying its rewarding effects. Surprisingly, PLAE attenuated such increment in the cocaine-induced locomotion and the overexpression of CB1 receptors induced by the HFD. At the same time, the effect of PLAE on MBP was reduced by the HFD.

Altogether, the results suggest the existence of a crosstalk among the mechanisms involved in such changes, so that in the first period of exposition to a HFD there might be a mutual attempt to compensate the effects induced by both conditions (PLAE and HFD). However, further investigation is needed to unveil what would occur with a longer exposure to a fatty diet.
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Conflict of interest
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Figure captions

Figure 1. (A) Experimental design. Pregnant C57BL/6 females were exposed to alcohol or water (control) during the gestation and lactation periods, following the drinking-in-the-dark (DID) procedure. On PND 29, female offspring were assigned a standard (SD) or a high-fat diet (HFD). After feeding for 8 weeks, their glucose (Glc) and triglyceride (TG) blood levels were determined and mice were tested either for cocaine-induced psychostimulation, cocaine-induced conditioned place preference (CPP) or for molecular determinations. (B) Experimental groups: there are four experimental groups according to the exposure of their mothers to water (W) or alcohol (A) during the gestation and lactation periods, and the diet they were fed (SD or HFD). PND: postnatal day.

Figure 2. Maternal alcohol drinking. (A) Volume of water or alcohol consumed and (B) Alcohol intake (g ethanol (EtOH)/kg), throughout the gestation (week 1-3) and lactation (week 4-6) periods during the DID procedure test. Results are expressed as mean ± SEM. *P < 0.05, **P < 0.01 day 4 of each week compared with the 3 previous days in the alcohol-exposed group. #P < 0.05, ##P < 0.01 and ###P < 0.001 day 4 of each week compared with the 3 previous days in the water-exposed group. (N=26-27/group).

Figure 3. (A) Body weight gain after 8 weeks of feeding, expressed in grams (N=25-27/group). (B) Mean daily Kcal ingested by mice exposed to water (W) or alcohol (A) during the gestation and lactation periods and exposed to a standard diet (SD) or a high fat diet (HFD) (N=25-27/group). (D) Fasting triglycerides (TG) and Glucose blood levels (N=6/group) and (E) NPY and (F) CPT1C gene expression in the hypothalamus (N=5/group) measured following 8 weeks of SD or HFD feeding. Results are expressed as mean ± SEM. ###P < 0.001 vs W-HFD; *P < 0.05, **P < 0.01, ***P < 0.001 vs. the corresponding group (water or alcohol) fed with SD; &&&P < 0.01 vs. SD.

Figure 4. Cocaine-induced CPP in mice exposed to water (W) or alcohol (A) during the gestation and lactation periods, after 8 weeks of feeding with a standard diet (SD) or a high fat diet (HFD). Bars represent the CPP score, that is the difference in seconds between the time spent in the drug-paired compartment during the testing and the pre-conditioning
sessions. (Inset) distance travelled by each experimental group during the preconditioning phase. Results are expressed as mean ± SEM (N=10-12/group). &&& P < 0.001 vs. saline-treated animals. *P < 0.05 and **P < 0.01 vs. the corresponding SD-group (water or alcohol).

Figure 5. (A) Cocaine-induced locomotor activity immediately after the drug injection (8 mg/kg i.p.) for 30 min. Results are expressed as mean ± SEM of the distance travelled (cm) (N=9-10/group). (B) [³H]DA uptake in striatal synaptosomes. Data are expressed as a percentage of control uptake relative to W-SD (mean ± SEM), (N=7-9/group). (C) Effects on TH and (D) CB1 protein expression in ventral striatum induced by PLAE and the subsequent exposure to a standard diet (SD) or a high fat diet (HFD) for 8 weeks. Representative pictures of the immunoblots are shown below the bar charts. Images of the whole immunoblot membranes can be found as Supplementary Figure S1. Results are expressed as mean ± SEM (N=5/group). * P < 0.05, **P < 0.01 vs. W-SD.

Figure 6. Effects on (A) NeuN, (B) NFκB and (C) MBP protein expression in prefrontal cortex induced by PLAE and the subsequent exposure to a standard diet (SD) or a high fat diet (HFD) for 8 weeks. Results are expressed as mean ± SEM. #P < 0.05 vs. W-SD. (N=5/group).
Set 1: The animals which did not receive cocaine in CPP were used for gene expression (PND 95).

- Set 1: Food intake. After blood samples for Glc and TG analysis, the animals were subjected to CPP (PND 85-95).
- Set 2:
  - Cocaine-induced psychostimulation (PND 85-86)
- Set 3:
  - [3H]DA uptake (PND 87)
  - Molecular determinations (PND 87)
Figure 2

A

Volume consumed

B

Alcohol intake
Figure 4

- W-SD
- A-SD
- W-HFD
- A-HFD

Cocaine

Saline

CPP score (seconds)

Distance (cm)

Group

\[***\]
Figure 5

A

B

C

D

TH expression (%)

CB1 expression (%)

SD     HFD

W-SD   A-SD   W-HFD   A-HFD
Figure 6

(A) NeuN expression (%)

(B) NFkB expression (%)

(C) MBP expression (%)

- W-SD
- A-SD
- W-HFD
- A-HFD