Research Article

Early Maternal Deprivation Enhances Voluntary Alcohol Intake Induced by Exposure to Stressful Events Later in Life

Sara Peñasco,1 Virginia Mela,1 Jose Antonio López-Moreno,2 María-Paz Viveros,1 and Eva M. Marco1

1Departamento de Fisiología (Fisiología Animal II), Facultad de Ciencias Biológicas, Universidad Complutense de Madrid, Instituto de Investigación Sanitaria del Hospital Clínico San Carlos (IDISSC), 28040 Madrid, Spain
2Departamento de Psicobiología, Facultad de Psicología, Universidad Complutense, 28223 Madrid, Spain

Correspondence should be addressed to María-Paz Viveros; pazviver@bio.ucm.es and Eva M. Marco; emmarco@bio.ucm.es

Received 15 November 2014; Revised 14 January 2015; Accepted 29 January 2015

Academic Editor: Nina N. Karpova

Copyright © 2015 Sara Peñasco et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

In the present study, we aimed to assess the impact of early life stress, in the form of early maternal deprivation (MD, 24h on postnatal day, pnd, 9), on voluntary alcohol intake in adolescent male and female Wistar rats. During adolescence, from pnd 28 to pnd 50, voluntary ethanol intake (20%, v/v) was investigated using the two-bottle free choice paradigm. To better understand the relationship between stress and alcohol consumption, voluntary alcohol intake was also evaluated following additional stressful events later in life, that is, a week of alcohol cessation and a week of alcohol cessation combined with exposure to restraint stress. Female animals consumed more alcohol than males only after a second episode of alcohol cessation combined with restraint stress. MD did not affect baseline voluntary alcohol intake but increased voluntary alcohol intake after stress exposure, indicating that MD may render animals more vulnerable to the effects of stress on alcohol intake. During adolescence, when animals had free access to alcohol, MD animals showed lower body weight gain but a higher growth rate than control animals. Moreover, the higher growth rate was accompanied by a decrease in food intake, suggesting an altered metabolic regulation in MD animals that may interact with alcohol intake.

1. Introduction

Epidemiological data indicate that adverse life events during the first few years of life can increase the risk for psychopathology, including drug addiction [1, 2]. In fact, early life stress has been proposed to predict alcohol drinking in adolescence and alcohol abuse and dependence at adulthood (see [3] for review). Alcohol dependence affects millions of people worldwide, and alcohol consumption at adolescence has reached alarming rates [4]. In rodents, the effects of early life stress can be studied in a controlled manner by removal of the dam during the first 2 weeks of life (for a review on the animals models of dam-litter interaction disruption, please consult [5–7]). The effects of early life stress on alcohol consumption currently available are inconsistent and highly dependent on environmental factors such as the species and strains used and the protocol used for the maternal separation protocol (see [8] as an example). In rats, repeated maternal separation during early postnatal life (180–360 min. a day during the first three weeks of life) has been reported to induce an increase in alcohol intake [9–12], although no changes in alcohol consumption have also been reported [13–15]. It is worth mentioning that the investigation of possible sex differences in this regard has been neglected. In the last years, we have focused on the investigation of the early maternal deprivation (MD) animal model, that is, 24h of dam-litter separation on postnatal day (pnd) 9. Adolescent animals exposed to MD have been reported to exhibit a trend to increased impulsivity, depressive-like responses, and impairments in cognitive function [16–18]. At adolescence, an increase in striatal dopamine levels has also been reported as a consequence of MD, possibly reflecting an altered brain reward system [19]. Actually, MD seems to modify the response of
animals to diverse drugs of abuse, including cannabino-
id [7, 18, 20], 3,4-methylenedioxymethamphetamine (MDMA) [21], and cocaine [22].

There is evidence about a relationship between stress and alcohol consumption (see [23, 24] for review) although the nature of this relationship is far from being understood. At this point, it is worth mentioning that, in a previous study, alcohol consumption during adolescence rendered animals more susceptible to acute stressor-specific effects in terms of alcohol consumption [25]. Therefore, in the present study alcohol deprivation was employed as a potential stressful stimulus that may influence motivation to drink [26, 27], and alcohol deprivation was also combined with restraint stress since exposure to restraint stress has been reported to increase alcohol consumption in rats [14].

In the present study we aimed to analyze the effects of MD on adolescent alcohol consumption, as well as addressing to which extent a previous history of early life stress (MD) could modulate the consequences of subsequent stressful life events (alcohol deprivation and alcohol deprivation combined with the concomitant exposure to restraint stress) on voluntary alcohol intake. Given the amount of evidence supporting sex differences in the consequences of MD, both male and female animals were employed in the present study.

2. Material and Methods

Experiments were performed in agreement with the European Directive 2010/63/EU on the protection of animals used for scientific purposes and in compliance with the Spanish Royal Decree 1201/2005, October 21, 2005 (BOE nº 252), about protection of experimental animals. All efforts were made to minimize animal suffering and distress.

2.1. Animals. Experiments were carried out on the offspring of rats purchased from Harlan Laboratories (Rossdorf, Germany), which were mated (one male × two females for ten consecutive days) in our animal facilities approximately 2 weeks after their arrival. After mating, female rats were housed individually in standard Plexiglas cages (50 × 25 × 17.5 cm.) and the animals were monitored daily for parturi-
tion. On the day of birth, postnatal day (pnd) 0, the litters were culled and sex balanced to 8 pups per dam (4 males and 4 females). No cross-fostering was allowed. Litters were randomly assigned to each experimental group: control (Co) and maternal deprivation (MD). All pups were weaned at pnd 22 and housed in pairs of sibling rats separated by sex. All animals were maintained in our animal facilities at the Faculty of Biological Sciences in the Complutense University of Madrid (EX08-UCS) at a constant temperature (21 ± 1°C) and humidity (60 ± 10%) in a reverse 12-hour dark-light cycle (lights on at 20.00 h), with free access to food (commercial diet for rodents A04/A03; SAFE, Augy, France) and water.

2.2. Early Maternal Deprivation (MD). Early MD was performed as previously described [15]. In brief, on pnd 9, litters were submitted to 24 hours of maternal deprivation; that is, dams were removed from their home cages at 09.00 h, pups were weighted and remained undisturbed in the same room until the next day (pnd 10, 09.00 h) when pups were weighted again, and dams were placed back in their corresponding home cages. Pups from the control group (Co) were not manipulated but for body weight control on pnd 9 and pnd 10. Body weight was registered every six days until weaning (on pnd 16 and pnd 22). A total of 10 litters were used in the present experiment (5 Co litters and 5 MD litters); however, due to a technical mistake, 2 cages from the MD group had to be excluded from the study, and thus the total number of animals used was 76 (20 Co males and 20 Co females and 18 MD males and 18 MD females).

2.3. Voluntary Alcohol Consumption: Two-Bottle Choice. Once the rats became adolescents (pnd 28), rats were given free access to a bottle containing an ethanol solution (20% v/v) in addition to the water bottle. The ethanol solution (v/v) was prepared using 96° ethanol (Alcoholes Aroca S.L., Madrid, Spain) diluted in tap water. The positions of the bottles were changed daily to avoid position preference. Ethanol solution and water were changed every day. Measurements of alcohol and water intake were made daily during the recording periods, as well as body weight. Voluntary alcohol intake was recorded (1) during adolescence, from pnd 28 to pnd 50; (2) during a four-day period after one week of alcohol cessation, from pnd 57 to pnd 60; and (3) during a four-day period after a second period of alcohol cessation combined with exposure to restraint immobilization stress (30 min. per day under white light conditions) on the last three days of alcohol cessation, from pnd 67 to pnd 70. The amount of alcohol intake was expressed as grams (g.) and alcohol intake per body weight (kg.). Daily voluntary alcohol consumption was calculated during the three recording periods, and alcohol intake during the first 2 h. of alcohol exposure was also evaluated for the two last periods following exposure to additional stress (see experimental schedule for details, Figure 1). Body weight and food intake were controlled throughout the experimental period. During the evaluation of alcohol consumption animals remained paired housed to avoid possible interferences with the MD protocol due to isolation [28, 29].

2.4. Food Intake, Body Weight, and Growth Rate. Food intake was evaluated by placing a known amount of food in each cage and measuring the remaining amount after 24 hours. This procedure was performed during the adolescent period (voluntary alcohol consumption) and after the first and second alcohol cessation periods. Body weight was registered at pnd 9, pnd 16, and pnd 22 (before weaning) and daily there-
after during the periods of voluntary alcohol consumption, that is, adolescence, and following alcohol cessation. Body weight (BW) gain (in grams) was analyzed and calculated as the difference in BW referenced to pnd 9 during the preweaning period, pnd 28 during adolescence, and pnd 50 during the two periods of alcohol cessation. Additionally, growth rate (GR) was calculated as the percentage of BW gain divided by the reference BW employed during each period: GR (%) = (BWgain/BWref.) × 100.
Neural Plasticity 3

Adolescence
Postnatal
Weaning
Adulthood
pnd 0 9–10 22 28 50 56 61 67 70
Voluntary alcohol intake
pnd 28–50
Control males
MD males
Control females
MD females
Water
Alcohol 20%
Maternal deprivation
MD
Two-bottle paradigm

Figure 1: Schedule of experimental design. A total of ten litters were assigned to either of these two experimental groups: exposure to a single episode of early maternal deprivation (24 h on pnd 9, MD) or not deprived (control group, Co). All animals were given free access to alcohol (20%, v/v) and water in the two-bottle choice paradigm for (1) the whole adolescent period (from pnd 28 to pnd 50), for (2) four days (pnd 57 to pnd 60) after one week of alcohol cessation, and for (3) four days (pnd 67 to pnd 70) after a second period of alcohol cessation combined with 30 min. of restraint stress under white light conditions on the last three days of alcohol cessation. n = 18–20 animals per experimental group.

2.5. Statistical Analyses. Data were analyzed using a two-way analysis of variance (ANOVA), considering sex (male or female) and neonatal condition (Co or MD) as independent factors. Shapiro-Wilk and Levene tests were used to confirm normality and homoscedasticity of the data. Repeated measures ANOVA was employed when appropriate, that is, BW gain and growth rate. Tukey post hoc comparisons were performed only for significant interactions, as well as additional one-way ANOVAs. Significance level was set at P < 0.05 for all comparisons. Statistical analyses were performed by the SPSS 19.0 software package (SPSS Inc., Chicago, IL, USA).

3. Results

3.1. Voluntary Alcohol Intake

3.1.1. Voluntary Alcohol Intake during Adolescence. No significant differences in alcohol intake were found during voluntary alcohol consumption at adolescence (control males: 16.6 ± 1.9 g/kg/day; control females: 17.2 ± 1.9 g/kg/day, MD males: 17.2 ± 2.0 g/kg/day; and MD females 20.2 ± 2.0 g/kg/day). Neither the repeated measures ANOVA nor the ANOVA analysis of the averaged alcohol intake during adolescence revealed any significant difference (see Figure 2). However, it is worth mentioning that MD females showed a trend for increased levels of alcohol intake during the first days of alcohol exposure.

3.1.2. Voluntary Alcohol Intake following One Week of Alcohol Cessation. Following one-week cessation, no significant differences were found on daily voluntary alcohol consumption (data not shown). However, a significant effect of the neonatal condition was found during the first 2 h of alcohol exposure (F(1, 34) = 4.48; P < 0.05). Both male and female animals consumed more alcohol than their corresponding Co animals during the first 2 h of alcohol exposure (Figure 3(a)). Although not significant, a trend for higher levels of alcohol consumption was found for female animals when compared to males during these first 2 h of alcohol exposure (Figure 3(a)). No significant interaction between sex and neonatal condition was found.

3.1.3. Voluntary Alcohol Intake following a Second Week of Alcohol Cessation Combined with Restraint Stress. Following a second period of alcohol cessation combined with restraint stress during the last three days, no significant differences were found on daily voluntary alcohol consumption (data not shown). However, a significant effect of the neonatal condition was found during the first 2 h of alcohol exposure (F(1, 34) = 5.47; P < 0.05). MD animals exhibited a higher alcohol intake than Co animals during the period of alcohol exposure (Figure 3(b)). During this time period, a significant effect of sex appeared (F(1, 34) = 5.30; P < 0.05). As for the previous time period, females showed higher levels of alcohol consumption compared to males during the first 2 h of alcohol exposure. No significant interaction between sex and neonatal condition was found.
3.2. Body Weight Gain and Growth Rate

3.2.1. Body Weight Gain and Growth Rate before Weaning.
During the preweaning period, a significant effect of the neonatal condition was found when analyzing BW gain ($F(1,16) = 89.88; P < 0.001$) and growth rate ($F(1,16) = 20.58; P < 0.001$). Neither a significant effect of the sex condition nor a significant interaction between factors was found. The MD episode induced a significant reduction in BW gain and growth rate during the preweaning period (Table 1).

3.2.2. Body Weight Gain and Food Intake during Adolescence.
During adolescence, when animals had free access to alcohol, BW gain (Figure 4(a)) and growth rate (Figure 4(b)) were...
significantly affected by sex \((F(1,72) = 281.62; P \leq 0.001; F(1,72) = 97.26; P < 0.001, \text{resp.})\) and neonatal condition \((F(1,72) = 6.47; P < 0.05; F(1,72) = 28.88; P < 0.001, \text{resp.})\). No significant interaction between sex and neonatal condition was found. As expected, female animals showed lower BW gain and growth rate than male animals, while MD animals showed a lower BW gain but a higher growth rate than control animals. The analysis of accumulated food intake rendered a significant effect of sex \((F(1,34) = 104.78; P < 0.001)\) with females eating less than males and a significant effect of the neonatal condition \((F(1,34) = 11.57; P < 0.005)\) with MD animals consuming less food than their corresponding controls (Figure 4(c)). No significant interaction was found in this case.

3.2.3. Body Weight (BW) Gain and Food Intake following One Week Alcohol Cessation and a Second Week of Alcohol Cessation Combined with Restraint Stress. A significant effect of sex on BW gain was found after the first \((F(1,72) = 501.87, P < 0.001)\) and second \((F(1,72) = 683.69; P < 0.001)\) periods of alcohol cessation, with females showing a lower BW gain than males across periods (data not shown). Similarly, the analysis of growth rate (Figures 5(a) and 5(c)) revealed a significant effect of sex after the first \((F(1,72) = 242.03, P < 0.001)\) and second \((F(1,72) = 341.55; P < 0.001)\) periods of alcohol cessation. Moreover, the analysis of growth rate also revealed a significant effect of the neonatal condition after the two alcohol cessation periods (first period: \(F(1,72) = 7.30; P < 0.01\) and second period: \(F(1,72) = 14.65; P < 0.001\)). For the growth rate analyses, a significant interaction between sex and neonatal condition was also found (first period: \(F(1,72) = 8.14; P < 0.01\) and second period: \(F(1,72) = 8.62; P < 0.005\)). During the two periods of alcohol exposure, female animals showed a lower growth rate than male animals, but MD only increased growth rate among male animals (no differences were observed among females). Regarding the analysis of the accumulated food intake (Figures 5(b) and 5(d)), a significant effect of sex was revealed (first period: \(F(1,34) = 210.18; P < 0.001)\) and second period: \(F(1,34) = 162.40; P < 0.001)\); male animals consumed more food than females during these two time periods. A significant effect of the neonatal condition was also found after the second period of alcohol cessation \((F(1,34) = 6.29; P < 0.05)\); during this period MD animals consumed less food than control animals. No significant interaction between sex and neonatal condition was found for any of the time periods analyzed.

### 4. Discussion

Early life stress, in the form of a single 24-hour episode of MD at pnd 9, did not affect baseline voluntary alcohol intake during adolescence in rats of both sexes. However, MD seemed to exacerbate the effects of subsequent stress exposure on alcohol intake. Animals exposed to neonatal MD, compared to control non-MD animals, showed higher levels of alcohol intake following a short period of alcohol cessation as well as after the combination of an additional period of alcohol cessation concomitant to restraint stress exposure.

Under the present experimental conditions, MD, *per se*, did not appear to predict prospective alcohol intake during adolescence in either sex. Previous studies have investigated the impact of early life stress on voluntary alcohol intake, but different protocols of maternal separation were employed, that is, repeated and shorter episodes of maternal separation [8], and it is well known that different experimental procedures of this model of early life stress render different results. Moreover, most of the previous studies have investigated the effects of early life stress on adult animals, leaving adolescents out of the focus. Following repeated brief episodes of neonatal maternal separation, some studies reported an increase in alcohol intake [9–12], although other studies found no changes in alcohol consumption [13–15]. Discrepancies with previous reports may result from the timeframe selected for alcohol exposure, that is, adolescence; the protocol of early life stress employed, that is, MD; and differences in the protocol of alcohol exposure, that is, previous alcohol habituation, alcohol concentration, and/or choice between two or more bottles. However, present findings suggest MD animals to be more vulnerable to the acute effects of stress in terms of later alcohol consumption. In fact, MD animals showed higher levels of alcohol intake than control animals following a short period of alcohol cessation, and an additional short period of alcohol cessation combined with restraint stress. A previous study reported an increase in voluntary alcohol intake in adolescent animals exposed to a protocol of repeated maternal separation by using a paradigm of intermittent free access to alcohol. Since the paradigm of alcohol exposure employed in that study included brief periods of alcohol withdrawal [30], also considered as stressful stimuli (see below), those results could be considered similar to our present findings. Alcohol withdrawal has been considered as a potent stressor capable of disrupting several

| Table 1: Metabolic parameters during the preweaning period. |
|------------------------------------------------------------------|
| Control Males | BW gain | Growth rate | BW gain | Growth rate | Control Females | BW gain | Growth rate | BW gain | Growth rate |
|----------------|---------|-------------|---------|-------------|----------------|---------|-------------|---------|-------------|
| pnd 10         | 1.9 ± 0.1 | 9.0 ± 0.8   | –0.8 ± 0.1 \(^B\) | –4.3 ± 0.3 \(^B\) | 1.7 ± 0.2  | 8.4 ± 1.2   | –0.9 ± 0.1 \(^B\) | –4.6 ± 0.4 \(^B\) |
| pnd 16         | 15.0 ± 0.3 | 70.3 ± 4.2  | 9.0 ± 0.6 \(^B\) | 46.5 ± 4.8 \(^B\) | 13.6 ± 1.4  | 66.5 ± 8.0  | 8.8 ± 0.7 \(^B\) | 46.8 ± 5.7 \(^B\) |
| pnd 22         | 37.3 ± 0.9 | 175.0 ± 9.7 | 27.9 ± 11.8 \(^B\) | 143.0 ± 5.4 \(^B\) | 34.0 ± 1.6  | 165.3 ± 10.4| 26.7 ± 0.8 \(^B\) | 140.5 ± 8.3 \(^B\) |

Body weight gain (g) and growth rate (%) on different days during the preweaning period. Reference day was considered postnatal day, pnd, 9. Rats were exposed to a single episode of early maternal deprivation (24h on pnd 9, MD) or not deprived (control group, Co). Data are expressed as mean ± S.E.M and were analyzed by a repeated measures two-way ANOVA, \(P < 0.05\). \(^B\) General effect of the neonatal condition. \(n = 5\) litters per experimental group.
stress circuitries including the HPA axis [24] and has been proposed as a suitable model to study the impact of alcohol reexposure on relapse [31]. Moreover, a previous study that investigated the vulnerability of animals to the consequences of stress on alcohol intake revealed that alcohol deprivations in combination with more severe stress protocols, that is, electric foot shock, were able to increase alcohol consumption if animals had initiated alcohol intake during adolescence [25]. Similarly, Roman et al. [14] reported an increase in alcohol intake during the exposure in adult life to a restraint protocol and throughout the postrestraint period in male and female animals exposed to repeated periods of brief maternal separation. The relationship between stress and alcohol consumption is complex and poorly understood (see [23, 24] for review); however, present data suggest that early life stress and in particular MD may render animals more vulnerable to the consequences of future stressful situations regarding alcohol intake.

Early life stress has been associated with alterations in the development and establishment of several neurotransmitter circuitries involved in the stress response. As an example, MD has been reported to alter the development of the dopaminergic, serotonergic, and glutamatergic systems, known to be involved in alcohol effects and vulnerability [19, 32, 33]. A disruption of the endogenous cannabinoid system has also been reported as a consequence of MD [17, 34–38]. The endocannabinoid (eCB) system appears to be of special interest in the context of the present study since it has been

Figure 4: Metabolic parameters during adolescence in response to voluntary alcohol intake. (a) Body weight (BW) gain (g), (b) growth rate (%), and (c) accumulated food intake (g). Rats were exposed to a single episode of early maternal deprivation (24 h on pnd 9, MD) or not deprived (control group, Co) and exposed during adolescence, from postnatal day (pnd) 28 to pnd 50, to voluntary alcohol intake (20% v/v, in a two-bottle choice paradigm). Data are presented as mean ± S.E.M and were analyzed by a repeated measures two-way ANOVA (BW gain and growth rate) or by a two-way ANOVA (accumulated food intake). $P < 0.05$; (A) general effect of sex and (B) general effect of the neonatal condition; $n = 18–20$ animals per experimental group.
Figure 5: Metabolic parameters following stressful events in young adult rats in response to voluntary alcohol intake. ((a), (c)) Growth rate (%) and ((b), (d)) accumulated food intake (g). Rats were exposed to a single episode of early maternal deprivation (24 h on pnd 9, MD) or not deprived (control group, Co) and exposed to voluntary alcohol intake (20% v/v, in a two-bottle choice paradigm) during adolescence and following one week of alcohol cessation (from pnd 57 to pnd 60, upper panels) and after a second period of alcohol cessation combined with exposure to restraint immobilization stress (30 min. per day under white light conditions) on the last three days of alcohol cessation (from pnd 67 to pnd 70, lower panels). Data are presented as mean ± S.E.M and were analyzed by a repeated measures two-way ANOVA (BW gain). Tukey post hoc comparisons: ***P < 0.005, MD males versus Co males; ###P < 0.005, Co females versus Co males; or by a two-way ANOVA (accumulated food intake). P < 0.05; (A) general effect of sex, (B) general effect of the neonatal condition; n = 18–20 animals per experimental group.

proposed as one of the underlying mechanisms of alcohol use and abuse [39] and, more recently, as a putative candidate for the proclivity to alcohol intake observed in animals submitted to certain protocols of maternal separation [40]. Therefore, the changes induced by the MD episode on the eCB system may possibly underlie the increased alcohol intake reported after the exposure to stressful challenges later in life.

In humans, epidemiological studies indicate higher rates of alcohol abuse and dependence in men compared to woman [41]. However, data from animal models show the opposite; female animals seem more prone to consume alcohol than males [42, 43]. Discrepancies between human and animal studies may result from sociocultural factors, but the investigation of the underlying neurobiological substrates may help us to better understand these sex differences. In the present study, female animals consumed more alcohol than males in a two-bottle choice paradigm following an episode of alcohol cessation combined with restraint stress. Since sex differences in the response to stressors have been extensively described [44, 45], a differential response to stress in male and female animals might underlie, at least in part, the greater influence of stress on alcohol intake in females.
As in previous studies from our group, MD animals showed a significantly diminished BW gain until weaning when compared to Co animals [36, 46, 47], as well as a significantly decreased growth rate. However, these growing profiles changed after weaning. During adolescence, when animals had free access to alcohol, MD animals still showed lower BW gain than control animals, but they also showed a higher growth rate than controls. These results suggest that MD animals gain less body weight but they did it faster than control animals. The higher growth rate was accompanied by a decrease in food intake, suggesting an altered body weight regulation in MD animals that is evidenced in adolescence, coincident with alcohol exposure. It is worth mentioning that later in life, as evaluated following the exposure to stressful events, that is, alcohol cessation and alcohol cessation combined with restraint stress, only among male animals did the effects of MD on growth rate persist. Though a specific effect of weaning on the different growth rate of control and MD animals cannot be ruled out, it is important to point out that MD modifies the metabolic response to a metabolic challenge such as a high fat diet [48]. From this point of view, the present data may indicate that MD alters the metabolic response to alcohol, apparently in a sex dependent manner, though a direct confirmation of this hypothesis would require further experiments. We have previously reported that MD rats show a long lasting leptin deficiency (observed in neonates and adult rats) as well as an altered hypothalamic development [47, 49, 50]. Recently, de Timary et al. [51] have proposed a model for disruption of energy balance in alcohol dependent subjects in which altered leptin levels play a central role. The interactions between metabolic alterations induced by early life stress and those induced by alcohol abuse deserve further investigation, since adaptations of energy balance may contribute (aside from the effects of alcohol on the brain reward system) to the modulatory effects of MD on stress-induced alcohol intake.

5. Conclusions

Early maternal deprivation did not affect baseline voluntary alcohol consumption during adolescence but enhanced the levels of voluntary alcohol intake following exposure to stressful events later in life. Moreover, this episode of early life stress seemed to alter energy balance and metabolism since somatic growth and food intake were persistently modified in these animals. Future research is needed in order to clarify the impact of metabolic changes, such as altered leptin levels, on the susceptibility to increase alcohol intake in response to stress.

Conflicts of Interest

The authors have no conflict of interests.

Authors’ Contribution

María-Paz Viveros, Eva M. Marco, and Jose Antonio López-Moreno were responsible for the design of the experiments performed. Sara Peñasco and Eva M. Marco contributed to the acquisition of animal data. Sara Peñasco, Virginia Mela, Eva M. Marco, and María-Paz Viveros assisted with data analysis and interpretation of findings. María-Paz Viveros and Eva M. Marco drafted the paper. All authors critically reviewed content and approved final version for publication.

Acknowledgments

The authors would like to thank María del Rincón for her assistance in the acquisition of animal data. They acknowledge the Instituto de Salud “Carlos III” (FIS), RETICS, Red de Trastornos Adictivos (RD2012/0028/0021; RD12/0028/0015), GRUPOS UCM-BSCH: Reference, UCM 95179, and the European Foundation for Alcohol Research (EA 12 21) for funding.

References

[1] C. Heim and C. B. Nemeroff, “The role of childhood trauma in the neurobiology of mood and anxiety disorders: preclinical and clinical studies,” Biological Psychiatry, vol. 49, no. 12, pp. 1023–1039, 2001.
[2] B. S. McEwen, “Early life influences on life-long patterns of behavior and health,” Mental Retardation and Developmental Disabilities Research Reviews, vol. 9, no. 3, pp. 149–154, 2003.
[3] M.-A. Enoch, “The role of early life stress as a predictor for alcohol and drug dependence,” Psychopharmacology, vol. 214, no. 1, pp. 17–31, 2011.
[4] World Health Organization, Global Status Report on Alcohol and Health, WHO Library Cataloguing-in-Publication Data, World Health Organization, 2011.
[5] F. Cirulli, A. Berry, and E. Alleva, “Early disruption of the mother-infant relationship: effects on brain plasticity and implications for psychopathology,” Neuroscience and Biobehavioral Reviews, vol. 27, no. 1-2, pp. 73–82, 2003.
[6] C. B. Faturi, P. A. Tiba, S. E. Kawakami, B. Catallani, M. Kerstens, and D. Suchecki, “Disruptions of the mother-infant relationship and stress-related behaviours: altered corticosterone secretion does not explain everything,” Neuroscience and Biobehavioral Reviews, vol. 34, no. 6, pp. 821–834, 2010.
[7] E. M. Marco, W. Adriani, R. Llorrente, G. Laviola, and M.- P. Viveros, “Detrimental psychophysiological effects of early maternal deprivation in adolescent and adult rodents: altered responses to cannabinoid exposure,” Neuroscience and Biobehavioral Reviews, vol. 33, no. 4, pp. 498–507, 2009.
[8] I. Nylander and E. Roman, “Is the rodent maternal separation model a valid and effective model for studies on the early-life impact on ethanol consumption?” Psychopharmacology, vol. 229, no. 4, pp. 555–569, 2013.
[9] R. L. Huot, K. V. Thrivikraman, M. J. Meaney, and P. M. Plotsky, “Development of adult ethanol preference and anxiety as a consequence of neonatal maternal separation in Long Evans rats and reversal with antidepressant treatment,” Psychopharmacology, vol. 158, no. 4, pp. 366–373, 2001.
[10] K. Ploj, E. Roman, and I. Nylander, “Long-term effects of maternal separation on ethanol intake and brain opioid and dopamine receptors in male wistar rats,” Neuroscience, vol. 121, no. 3, pp. 787–799, 2003.
[11] E. Roman, P. Hyytiä, and I. Nylander, “Maternal separation alters acquisition of ethanol intake in male ethanol-prefering
AA rats,” Alcoholism: Clinical and Experimental Research, vol. 27, no. 1, pp. 31–37, 2003.

[12] E. Roman, L. Gustafsson, P. Hytiiä, and I. Nylander, “Short and prolonged periods of maternal separation and voluntary ethanol intake in male and female ethanol-prefering AA and ethanol-avoiding ANA rats,” Alcoholism: Clinical and Experimental Research, vol. 29, no. 4, pp. 591–601, 2005.

[13] J. N. Jaworski, D. D. Francis, C. L. Broman, E. T. Morgan, and M. J. Kuhar, “Effects of early maternal separation on ethanol intake, GABA receptors and metabolizing enzymes in adult rats,” Psychopharmacology, vol. 181, no. 1, pp. 8–15, 2005.

[14] E. Roman, K. Ploj, and I. Nylander, “Maternal separation has no effect on voluntary ethanol intake in female Wistar rats,” Alcohol, vol. 33, no. 1, pp. 31–39, 2004.

[15] M. Marmendal, E. Roman, C. J. P. Eriksson, I. Nylander, and C. Fahlike, “Maternal separation alters maternal care, but has minor effects on behavior and brain opioid peptides in adult offspring,” Developmental Psychobiology, vol. 45, no. 3, pp. 140–152, 2004.

[16] E. M. Marco, W. Adriani, R. Canese, F. Podo, M. P. Viveros, and G. Laviola, “Enhancement of endocannabinoid signalling during adolescence: modulation of impulsivity and long-term consequences on metabolic brain parameters in early maternally deprived rats,” Pharmacology Biochemistry and Behavior, vol. 86, no. 2, pp. 334–345, 2007.

[17] E. M. Marco, M. Valero, O. de la Serna et al., “Maternal deprivation effects on brain plasticity and recognition memory in adolescent male and female rats,” Neupharmacology, vol. 68, pp. 223–231, 2013.

[18] R. Llorente, L. Arranz, E.-M. Marco et al., “Early maternal deprivation and neonatal single administration with a cannabinoid agonist induce long-term sex-dependent psychoimunoenocrine effects in adolescent rats,” Psychoneuroendocrinology, vol. 32, no. 6, pp. 636–650, 2007.

[19] R. Llorente, E. O’Shea, M. D. Gutierrez-Lopez, A. Llorente-Berzal, M. I. Colado, and M.-P. Viveros, “Sex-dependent maternal deprivation effects on brain monoamine content in adolescent rats,” Neuroscience Letters, vol. 479, no. 2, pp. 112–117, 2010.

[20] A. Llorente-Berzal, S. Fuentes, H. Gagliano et al., “Sex-dependent effects of maternal deprivation and adolescent cannabinoid treatment on adult rat behaviour,” Addiction Biology, vol. 16, no. 4, pp. 624–637, 2011.

[21] A. Llorente-Berzal, C. Manzanedo, M. Daza-Losada et al., “Sex-dependent effects of early maternal deprivation on MDMA-induced conditioned place preference in adolescent rats: possible neurochemical correlates,” Toxicology, vol. 311, no. 1-2, pp. 78–86, 2013.

[22] A. Llorente-Berzal, M. A. Assis, T. Rubino et al., “Sex-dependent changes in brain CB1R expression and functionality and immune CB2R expression as a consequence of maternal deprivation and adolescent cocaine exposure,” Pharmacological Research, vol. 74, pp. 23–33, 2013.

[23] S. Macri, S. Spinelli, W. Adriani, J. Dee Higley, and G. Laviola, “Early adversity and alcohol availability persistently modify serotonin and hypothalamic-pituitary-adrenal-axis metabolism and related behavior: what experimental research on rodents and primates can tell us,” Neuroscience and Biobehavioral Reviews, vol. 31, no. 2, pp. 172–180, 2007.

[24] H. C. Becker, M. F. Lopez, and T. L. Doremus-Fitzwater, “Effects of stress on alcohol drinking: a review of animal studies,” Psychopharmacology, vol. 218, no. 1, pp. 131–156, 2011.

[25] S. Siegmund, V. Vengeliene, M. V. Singer, and R. Spanagel, “Influence of age at drinking onset on long-term ethanol self-administration with deprivation and stress phases,” Alcoholism: Clinical and Experimental Research, vol. 29, no. 7, pp. 1139–1145, 2005.

[26] F. Alen, G. Moreno-Sanz, A. Isabel de Tena et al., “Pharmacological activation of CB1 and D2 receptors in rats: predominant role of CB1 in the increase of alcohol relapse,” European Journal of Neuroscience, vol. 27, no. 12, pp. 3292–3298, 2008.

[27] F. Alen, A. Mouret, M.-P. Viveros et al., “Converging action of alcohol consumption and cannabinoid receptor activation on adult hippocampal neurogenesis,” International Journal of Neuropsychopharmacology, vol. 13, no. 2, pp. 191–205, 2010.

[28] F. Biggio, M. G. Pisu, A. Garau et al., “Maternal separation attenuates the effect of adolescent social isolation on HPA axis responsiveness in adult rats,” European Neuropsychopharmacology, 2014.

[29] J. Zlatković, N. Todorović, M. Bošković, S. B. Pajović, M. Demajo, and D. Filipović, “Different susceptibility of prefrontal cortex and hippocampus to oxidative stress following chronic social isolation stress,” Molecular and Cellular Biochemistry, vol. 393, no. 1-2, pp. 43–57, 2014.

[30] L. Daoura, J. Haaker, and I. Nylander, “Early environmental factors differentially affect voluntary ethanol consumption in adolescent and adult male rats,” Alcoholism: Clinical and Experimental Research, vol. 35, no. 3, pp. 506–515, 2011.

[31] A. D. Lê and Y. Shaham, “Neurobiology of relapse to alcohol in rats,” Pharmacology and Therapeutics, vol. 94, no. 1-2, pp. 137–156, 2002.

[32] B. A. Ellenbroek, N. Derks, and H.-J. Park, “Early maternal deprivation retards neurodevelopment in Wistar rats,” Stress, vol. 8, no. 4, pp. 247–257, 2005.

[33] M. Roceri, W. Hendriks, G. Racagni, B. A. Ellenbroek, and M. A. Riva, “Early maternal deprivation reduces the expression of BDNF and NMDA receptor subunits in rat hippocampus,” Molecular Psychiatry, vol. 7, no. 6, pp. 609–616, 2002.

[34] J. Suárez, R. Llorente, S. Y. Romero-Zerbo et al., “Early maternal deprivation induces gender-dependent changes on the expression of hippocampal CB1 and CB2 cannabinoid receptors of neonatal rats,” Hippocampus, vol. 19, no. 7, pp. 623–632, 2009.

[35] J. Suárez, P. Rivera, R. Llorente et al., “Early maternal deprivation induces changes on the expression of 2-AG biosynthesis and degradation enzymes in neonatal rat hippocampus,” Brain Research, vol. 1349, pp. 162–173, 2010.

[36] A. Llorente-Berzal, V. Mela, E. Borcel et al., “Neurobehavioral and metabolic long-term consequences of neonatal maternal deprivation stress and adolescent olanzapine treatment in male and female rats,” Neupharmacology, vol. 62, no. 3, pp. 1332–1341, 2012.

[37] M. López-Gallardo, R. Llorente, A. Llorente-Berzal et al., “Neuronal and glial alterations in the cerebellar cortex of maternally deprived rats: gender differences and modulatory effects of two inhibitors of endocannabinoid inactivation,” Developmental Neurobiology, vol. 68, no. 12, pp. 1429–1440, 2008.

[38] M. López-Gallardo, A. B. López-Rodríguez, A. Llorente-Berzal et al., “Maternal deprivation and adolescent cannabinoid exposure impact hippocampal astrocytes, CB1 receptors and brain-derived neurotrophic factor in a sexually dimorphic fashion,” Neuroscience, vol. 204, pp. 90–103, 2012.

[39] M. J. Pava and J. J. Woodward, “A review of the interactions between alcohol and the endocannabinoid system: implications
for alcohol dependence and future directions for research,” *Alcohol*, vol. 46, no. 3, pp. 185–204, 2012.

[40] A. Romano-López, M. Méndez-Díaz, A. E. Ruiz-Contreras, R. Carrizosa, and O. Prospéro-García, “Maternal separation and proclivity for ethanol intake: a potential role of the endocannabinoid system in rats,” *Neuroscience*, vol. 223, pp. 296–304, 2012.

[41] A. Kalaydjian, J. Swendsen, W.-T. Chiu et al., “Sociodemographic predictors of transitions across stages of alcohol use, disorders, and remission in the National Comorbidity Survey Replication,” *Comprehensive Psychiatry*, vol. 50, no. 4, pp. 299–306, 2009.

[42] J. M. Barker, M. M. Torregrossa, A. P. Arnold, and J. R. Taylor, “Dissociation of genetic and hormonal influences on sex differences in alcoholism-related behaviors,” *The Journal of Neuroscience*, vol. 30, no. 27, pp. 9140–9144, 2010.

[43] J. Simpson and J. P. Kelly, “An investigation of whether there are sex differences in certain behavioural and neurochemical parameters in the rat,” *Behavioural Brain Research*, vol. 229, no. 1, pp. 289–300, 2012.

[44] B. S. McEwen and J. H. Morrison, “The brain on stress: vulnerability and plasticity of the prefrontal cortex over the life course,” *Neuron*, vol. 79, no. 1, pp. 16–29, 2013.

[45] C. Dalla, P. M. Pitychoutis, N. Kokras, and Z. Papadopoulou-Daifoti, “Sex differences in response to stress and expression of depressive-like behaviours in the rat,” *Current Topics in Behavioral Neurosciences*, vol. 2011, no. 8, pp. 97–118, 2011.

[46] R. Llorente, C. Miguel-Blanco, B. Aisa et al., “Long term sex-dependent psychoneuroendocrine effects of maternal deprivation and juvenile unpredictable stress in rats,” *Journal of Neuroendocrinology*, vol. 23, no. 4, pp. 329–344, 2011.

[47] M. P. Viveros, R. Llorente, M. López-Gallardo et al., “Sex-dependent alterations in response to maternal deprivation in rats,” *Psychoneuroendocrinology*, vol. 34, supplement 1, pp. S217–S226, 2009.

[48] V. Mela, A. Llorente-Berzal, F. Díaz, J. Argente, M.-P. Viveros, and J. A. Chowen, “Maternal deprivation exacerbates the response to a high fat diet in a sexually dimorphic manner,” *PLoS ONE*, vol. 7, no. 11, Article ID e48915, 2012.

[49] M. P. Viveros, F. Díaz, B. Mateos, N. Rodriguez, and J. A. Chowen, “Maternal deprivation induces a rapid decline in circulating leptin levels and sexually dimorphic modifications in hypothalamic trophic factors and cell turnover,” *Hormones and Behavior*, vol. 57, no. 4–5, pp. 405–414, 2010.

[50] M. P. Viveros, R. Llorente, F. Díaz et al., “Maternal deprivation has sexually dimorphic long-term effects on hypothalamic cell-turnover, body weight and circulating hormone levels,” *Hormones and Behavior*, vol. 58, no. 5, pp. 808–819, 2010.

[51] P. de Timary, P. D. Cani, J. Duchemin et al., “The loss of metabolic control on alcohol drinking in heavy drinking alcohol-dependent subjects,” *PLoS ONE*, vol. 7, no. 7, Article ID e38682, 2012.