Neurodevelopment is Particular Vulnerable in Neonatal Male Rats Subjected to Maternal Separation

Chunyao Yang1*, Jing Sun2 and Changsheng Li3

1Department of Anesthesiology and Perioperative Medicine, Zhengzhou University, China
2Department of Psychiatry, Zhengzhou University, China
3Department of Anesthesiology and Perioperative Medicine, Zhengzhou University, China

*Corresponding author: Chunyao Yang, Department of Anesthesiology, the Affiliated Cancer Hospital of Zhengzhou University, Zhengzhou, China.

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Abstract

Objective: To explore the influence of maternal separation in early life on the neurodevelopment and young adult behavior of neonatal rats. Method Both male and female neonatal Sprague-Dawley rats were subjected to 2 h of maternal separation twice a day from postnatal days 2 (P2) through P9 followed by electroencephalogram (EEG) for 2 h on P10 or P11. Serum corticosteroid secretion and hippocampus expression of mGlu1R and mGlu5R were determined after EEG. Behavioral tests were examined by prepulse inhibition and elevated plus maze between P60~P70.

Results: The male group had increased corticosteroid level $t(10)=-5.458$, $P<0.001$ and decreased mGlu1R and mGlu5R expression in the hippocampi compared to the control group, with greater abnormality in EEG (This is defined by the total duration of seizure $t(10)=6.134$, $P<0.001$, number of episodes $t(10)=3.669$, $P=0.016$ and duration of a single episode $t(10)=2.916$, $P=0.009$) compared to female pups. The male rats were more severely disturbed than female rats in prepulse inhibition and performed worse in elevated plus maze treatment.

Conclusion: Maternal separation in early life induced significant abnormalities in EEG and alterations in expression of hippocampus mGlu1R and mGlu5R with greater changes in the corticosteroid level in males and extended behavior abnormalities in young adult rats.

Keywords: Neurodevelopment; Electroencephalogram; Maternal separation; Behavior

Introduction

The number of young people from rural areas that go to work in cities is increasing every year in China. In order to reduce the life burden or work convenience, the young parents usually leave their kids in native family. The children living with their grandparents and other dependents are deprived parental care and love. Children in this special child growth environment may have an increased risk of developing mood and anxiety disorders. Clinical studies showed that exposure to early adverse events such as childhood sexual abuse and trauma are associated with an increased risk of developing psychiatric disorders in adulthood [1-4]. There is no data about whether the children deprived parental care have prolonged influence in adulthood; however, in animal experiments, hypothalamus corticotrophin releasing hormone (CRH) abnormality is emerging in the neonatal rats with only one 2-hour separation from their dams [5]. After repeated separation for 3 hours per day, rats showed a greater hypothalamic-pituitary-adrenal (HPA) axis response to acute restraint stress compared to non-separated control rats [6]. Before we found a brief exposure to ET combined with a subsequent episode of stress early in life induced significant alterations in expression of amygdala CRH and some gene expression in male rats [7]. Even 2h maternal separation showed abnormal electroencephalogram activity [8].

The first two postnatal weeks is critical period, because a complex program of gene expression involved in structural and functional plasticity in neurodevelopment, forms that is responsible for subsequent brain functions and emotional behaviors during adulthood. Little is known about the ontogeny and function of the neuroendocrine system of children separated from their parents and its contribution to neurology abnormal development, growth failure, or depression after repeated maternal separation during the childhood. It has been suggested that neuronal activity modifies the chromatin complex at the crossroads of transcriptional activation...
andregulatedgeneticexpression[9,10].Thereisevidence
demonstratingthataberranttranscriptionalregulationisoneofthe
keycomponentsofthepathogenesisofsevereneschizophrenic
disorders,includingmooddisorders[11,12].ThemGluR1
andmGlu5Rgenesareessentialforsynapticencodingofspatial
experienceandimportantmediatorsforawiderangeofchronic
psychosocialstress-inducedalterationsinhumans[13,14].ThemGlu1
andmGlu5receptorsareinvolvedinhumanstudiesandmechanismsofactivity-
dependentsynapticplasticityandaretargetedbydrugdeveloped
forthetreatmentofcentralnervoussystem(CNS)disorders[15].
Inapreviousstudy,we didn’tfindmGlu1RandmGlu5Ralternation
incaudalbrainofShank3+/−/−miceinpreviousstudy,butmGlu5R
decreasedinculturedrathippocampalandcorticalneuronsinvitro
studybyVerpellietal.[16,17].ThrerateHPAaxisishypo-responsive
inthefirsttwo postnatalweeks.ThetherainHPA shares many
molecularandbiologicalcharacteristicswiththehumanHPA,so
theratmaternal separationisanappropriate model to study the
humanneurodevelopmentabnormalities.

The paradigm consisted of both male and female rats. The pups
were separated from the dams for two 2-hour separation phases
from postnatal day 2 through 9 to mimic the leftover
lidsneglectedbytheir workingparents. Wetaimedto examine the
neurodevelopmentvulnerabilityofdeprivedparentcare
inchildhood to behavioral abnormality in adulthood. Wefirst
defined the neurological mechanism underlying the development
of stress vulnerability in early maternally separated rats. Wetthen
investigated the sex difference in the neuroendocrine system
andbehavioralconsequencesofexposureto repeatedmaternal
separation in infants.

Methods and Materials
Animals
All experimental procedures were approved by the Institutional
Animal Care and Use Committee of the Affiliated Cancer Hospital of
Zhengzhou University. Male and female Sprague-Dawley rats were
studied. Animals were housed under controlled illumination (12-h
light/dark, lights on at 7:00 a.m.) and temperature (23–24 °C) with
free access to food and water. Within 24h of delivery, litters were
culled to 12 pups. At the beginning of each experiment the pups
were determined to be well nourished as judged by their stomachs
being full of milk (detectable through the transparent abdominal
wall). Delivery was verified at 12-hour intervals, and the day of
birth was considered postnatal day 0. On postnatal day 2, pups were
placed with mothers in cages and maternal separation treatment
began. The data reported in this study was collected from 120 rats
distributed evenly based on sex.

Maternal separation procedure
Twelve experimental groups were studied. On postnatal day 2
pups in experimental litters were separated from their dams twice
daily at the same time each day (9:00 a.m. - 11:00 a.m. and 15:00 p.m.
- 17:00 p.m.) in the morning and afternoon. Each subsequent day
through P9 to mimic the children separated from their parents.

Maternal separated pups were kept in a temperature-controlled
chamber (±37 °C) with a continuous supply of oxygen (1.5 L/
min). The control animals were raised undisturbed. After the last
separation pups were returned to their respective dams. Rats
were weaned on P21 and group-housed thereafter with same-sex
littermates. On P60, a subset of rats were subjected to behavioral
tests (Figure 1, A).

Electroencephalogram recording
After sequential maternal separation, electroencephalogram
(EEG) recordings were performed on P10 or P11 in a thermostated
chamber (±37 °C) with a continuous supply of oxygen (1.5 L/min) as
previous described[8]. Gas was monitored by a calibrated Datex side
stream analyzer (Datex-Ohmeda, Helsinki, Finland), which sampled
from the interior of the animal chamber. A minor operation was
performed under isoflurane anesthesia (1.6-2.0%) by the same
colleague, where four screw electrodes were implanted bilaterally
in the occipital and frontal regions of the rat pup skull with the left
frontal electrode serving as the reference electrode. The electrode
headmount was fixed by the screws and assured a good contact with
the skull. After the operation, the rat moved to the EEG recording
system. Continuous 2 hours EEG recordings were performed using
an EEG/EMG system (Pinnacle Technology, Lawrence, KS) (Figure
1, A). Acquisition of the EEG was performed using the Sirenia
software (Pinnacle Technology). The sampling interval per signal
was 200 μs. Data was filtered offline using a bandpass Bessel (8-
pole) 0.04–56-Hz filter. Sirenia CGMS analysis program assisted for
the EEG data analysis. Electroencephalogram patterns that were
characterized by the amplitude at least three times higher than
baseline and abruptly reverted to baseline were defined as seizure
electroencephalogram patterns. In most cases these patterns
start as high frequency-low amplitude activity that developed to
increased amplitude and decreased frequency and then abruptly
reverted to baseline activity. The total duration, number of episodes
and average duration of episode were calculated. The investigators
analyzing the EEGs were blinded to the experimental conditions
and all EEGs were reviewed by three independent reviewers.

Corticosterone and hippocampus mGlu1R and mGlu5R
determination
For measurements of the corticosteroid secretion, blood
samples (~300 μL) were collected using the “tail clip” method after
the EEG recording. Corticosterone concentration was determined
using a commercial ELISA kit (Cayman Chemical Company, Ann
Arbor, MI) following the manufacturer’s instructions. When blood
collecting finished, the rats were decapitated and the hippocampi
werequickly dissected from the brain and frozen in liquid
nitus. The mGlu1R and mGlu5R protein in the hippocampus
were detected as previously described[16]. Briefly, after the tissues
were homogenized and the protein was extracted, mGlu1R and
mGlu5R were resolved by sodium dodecylsulfate-polyacrylamide
gel electrophoresis and electrotransferred to nitrocellulose
membranes. The membranes were blocked in 0.1% Tween-20 in
Tris-HCl–buffered saline (TBST) containing 5% nonfat milk for
1h at room temperature and then immunoblotted with primary antibodies (anti-mGlu1R: 1:1000, Cell-signaling, Danvers, MA; anti-mGlu5R: 1:5000, β-actin:1:100,000, Sigma-Aldrich, St. Louis, MO) in TBST buffer containing 5% nonfat milk overnight at 4°C. After being washed extensively in TBST, the membranes were incubated for 1 h with horseradish peroxidase-conjugated anti-mouse or anti-rabbit immunoglobulin (Bio-Rad Laboratories, Hercules, CA) at a dilution of 1:2,000. Proteins were detected by enhanced chemiluminescence (Amersham, Piscataway, NJ). β-actin served as a loading control. The immunoblottings hands were quantified by densitometry using Image J software (National Institutes of Health, Bethesda, MD) and analyzed.

Behavioral tests

Young adult rats were sequentially evaluated in the elevated plus maze (EPM) and prepulse inhibition (PPI) of the acoustic startle response at P60~P70.

Assessment of behavior in the EPM: The elevated plus maze is a commonly used test to measure anxiety in rodents and consists of two open arms and two closed arms that are elevated above the ground. Rats that are less anxious will spend more time in the open arms compared to control rats, and rats that are more anxious will spend less time in the open arms compared to controls. This test provides a sensitive measure of anxiety, and we used the test to evaluate whether seizures on postnatal days 10–11 result in anxiety. The EPM studies were performed using the EPM apparatus and BIO-EPM 3C video tracking software (EB Instruments, Pinellas Park, FL) during the light phase of the dark-light cycle. The maze consists of two opposing open (50 × 10 × 0.5 cm) and two enclosed (50 × 10 × 45 cm) arms elevated 75 cm above the floor, with a 0.5-cm edge on the open arms. Animals were placed in the center square facing an open arm and allowed to acclimate to the maze for 5 min, at which time they were removed from the apparatus. During EPM testing, each rat’s behavior was recorded using BIO-EPM 3C video tracking software. The percentage of time spent in the open and enclosed arms, and the total distance traveled during 5 min of recording as an index of locomotor activity were compared.

Measurements of the acoustic startle response and PPI of startle: The PPI of startle tests were performed using the SR-Lab startle apparatus (San Diego Instruments, San Diego, CA) to assess sensorimotor gating. Testing occurred during the light phase of the dark-light cycle. The maze consists of two opposing open (50 × 10 × 0.5 cm) and two enclosed (50 × 10 × 45 cm) arms elevated 75 cm above the floor, with a 0.5-cm edge on the open arms. Animals were placed in the center square facing an open arm and allowed to acclimate to the maze for 5 min, at which time they were removed from the apparatus. During EPM testing, each rat’s behavior was recorded using BIO-EPM 3C video tracking software. The percentage of time spent in the open and enclosed arms, and the total distance traveled during 5 min of recording as an index of locomotor activity were compared.

Statistical Analysis

Values are reported as mean ± SEM. SigmaPlot 12.5 software (Systat Software, Inc., Point Richmond, CA) was used for statistical analyses. Single comparisons were tested using the t-test. All comparisons were run as two-tailed tests. A P< 0.05 was considered significant.

Results

The data clearly show that males appear to be more responsive to maternal separation compared to females. Maternal separation causes more seizures events in electroencephalograms of male than female P10/P11 rat pups. As explained in detail in experimental procedures, we separated the pups and dams from P2 through P9 and test the gender difference of electroencephalograms in P10 or P11. Analysis of 2 hour electroencephalographic recordings revealed statistically significant differences in seizure activities in male and female rat pups. The total duration of seizure (t(10)=6.134, P<0.001), the number of episodes (t(10)=3.669, P=0.016) and the duration of a single episode (t(10)=2.916, P=0.009) during 2 hours were higher in male (n=10) than female (n=10) rat pups. The electroencephalographic recording was normal in control pups with no seizure activity discovered (Figure 2).

Male pups showed higher vulnerability in response to maternal separation stress than female pups.

Stressors activate the hypothalamic-pituitary-adrenal (HPA) axis, triggering secretion and compensatory synthesis of hypothalamic corticotropin-releasing hormone (CRH). CRH is located in peptidergic neurons in the paraventricular hypothalamic nucleus, is secreted from nerve terminals to influence a rapid hormonal secretion from corticotrophs in the anterior pituitary [19,20]. In basal condition, male pups have greater plasma corticosterone levels than female pups. Maternal deprivation increased CORT levels in male in comparison to the control pups (t(10)=5.458, P<0.001). However, the female pups didn’t showed hyper reactivity of the HPA axis to repeated separation stress (t(10)=0.174, P=0.864). After the EEG experiments, the CORT level was higher in male pups compared to female pups (t(10)=9.245, P<0.001) (Figure 1, D).

Among all glutamatergic system components, metabotropic receptors play a main role in regulating neuronal excitability and synaptic plasticity. mGlu1R and mGlu5R present in the hippocampus are important in regulating the activity of the HPA axis after stress. mGlu1R and mGlu5R expression decrease significantly in male pups deprived mother care in comparison to the control (Figure 1, B).

Maternal separation in the neonatal period induced extended behavior abnormalities in adulthood. The long-term effects of maternal separation were evaluated by assessing sensorimotor gating. Both male (t(10)=−15.4, P<0.001) and female (t(10)=−6.583,
P<0.001) rats with neonatal maternal separation from P2 to P9 exhibited reduced PPI of the acoustic startle response at lowest prepulse intensity (5dB) when compared to rats in the control. But, male rats showed reduced PPI responses at prepulse intensities of 10dB compared to control group(\(t(10)=-10.900, P<0.001\)). PPI responses at prepulse intensities of 15dB were similar across all groups and between sexes (Figure1, E).

In the elevated plus maze, adult female rats traveled similar total distances and stayed in open arms for similar times in comparison to the control, but male rats spent a shorter time in open arms(\(t(10)=-4.871, P<0.001\)) of the EPM and covered shorter distances when compared to the control(\(t(10)=-7.928, P<0.001\)) (Figure1, C1&C2).

**Figure 1:** The influence of maternal separation exert on male and female neonatal rats. A: illustration of the experimental protocols. B: Representative Western blotting showing expression of mGlu1R and mGlu5R in hippocampus after EEG test. C: Maternal separation in early life from P2 to P9 lead to reduced time spent in open arms of the elevated plus maze (EPM) in young male, but not female rats. The EPM tests were performed at P60~P70. Shown are % of time spent in open arms of the EPM and distance traveled (C1) and (C2). Data are means ± SEM\((n=10)\). *P < 0.05 vs. Control. D: Maternal separation in early life from P2 to P9 lead to increased plasma corticosterone level after EEG recording in male, but not female rats. Data are means ± SEM\((n=10)\). *P < 0.05 vs. Control. E: Maternal separation in early life from P2 to P9 lead to impair sensorimotor gating function both in young male and female rats, but male rats showed reduced PPI responses at prepulse intensities of 10 dB compared to control group. The female in PPI responses at prepulse intensities of 10dB and 15 dB were similar. Histogram showing % prepulse inhibition (PPI) of startle in different groups\((n=10)\). *P<0.05 vs control male group. #, P<0.05 vs control female group.
Figure 2: Maternal separation caused more seizure events in electroencephalograms of male than female in a postnatal day (P) 10 or 11 rat pups. A: Examples of normal electroencephalogram patterns in P 10 or 11 rat during two hours in control rats (A1), and seizures electroencephalogram patterns in male(A2) and female(A3) rats. B: Histograms showing parameters of seizure electroencephalogram patterns during two hours in P10 or 11 (n=10). *, P<0.05 vs male rat pups.

Discussion

In the present study, we found that maternal separation exerted a vulnerable stress on male rats compared to their counterpart females. At the molecular level, maternal separation decreased the expression of mGlu1R and mGlu5R, and increased the corticosterone levels. Prolonged maternal separation during early infancy led to disturbances in prepulse inhibition and behavioral detriments in young adults especially for males. Early life chronic stress is one of the most prominent environmental factors associated with an increased risk of developing mood and anxiety disorders [21]. Early separation may act as a stressor and have adverse effects on adult neural and behavioral outcomes [7,22]. Maternal separation is an animal model of early life stress that modulates the HPA axis and studies the neurobiological underpinnings of disruption of the mother-infant relationship and loss of parental care, a highly prevalent condition in humans [23]. This paradigm may more closely approximate the human situation of children separated from their parents who are deprived parental care in infancy. The human brain requires a wide variety of experiences and environmental inputs in order to develop normally. Children who are neglected by caregivers or raised in institutional environments are deprived of numerous types of species-expectant environmental experiences that confer risk for internalizing and externalizing psychopathology [24]. Neglect disrupts the development of brain, exaggerating the activity of the HPA axis and correlating to adult depression and anxiety [25]. Depression, anxiety, and poor growth are well established consequences of chronic neglect occurring in infancy. These abnormal attachments are, in part, correlated with disregulation of the HPA axis, as shown by elevated corticosterone

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levels in separated animals. It should be noted that daily separation from dams in neonatal exerted an important impression on males. This study showed that mGlu1R and mGlu5R expression were decreased in male pups compared to female pups, but corticosterone levels were increased significantly in male pups. mGlu1R and mGlu5R present in the hippocampus are important in regulating the activity of the HPA axis after stress and within the hypothalamus to regulate rapid hormonal responses to stress [26]. Glutamate is essential for learning and memory processing, specifically, mGlu1R and mGlu5R enhance NMDA receptors excitability and play a key role in neurodevelopment positive. Other animal experiments indicated that modulation of mGlu1 and mGlu5 receptors prevented and reversed ethanol-induced memory impairment [27-29].

The interaction between glutamate and NMDA receptors is vital for synaptic plasticity and cognition. Stimulated glutamate receptors enhance NMDAR responses in the hippocampus leading to gene expression, synaptic plasticity, and memory consolidation [30]. Cognitive dysfunction may be improved by targeting mGlu5R with an appropriate therapy [31]. mGlu1R and mGlu5R alterations can lead to molecular imbalance between the excitation and inhibition underlying the emergence of a schizophrenic-like phenotype [32]. It is important to evaluate the potential of glutamate modulators in reversing the deficits characterizing the schizophrenic pathology. Together, these findings suggest that decreased mGlu1R and mGlu5R might be associated with increased HPA axis response to stress. This increased exposure to corticosterone and hyper-reactivity of the HPA axis can increase depression and anxiety-related behaviors in rodents. Among all glutamatergic system components, metabotropic receptors play a main role in regulating neuronal excitability and synaptic plasticity, specifically, mGlu1 and mGlu5 play important role in environmental modulation of schizophrenia-related behavioral impairments [33]. There is no known causal relationship between epilepsy and mGlu1R and mGlu5R; however, the male pups showed higher EEG abnormalities that are characteristic of seizure activities. Behaviors tested in adults suggest that they may share anatomical and molecular mechanisms, especially during brain development. A single 24-hour period of maternal deprivation at postnatal day 9 led to disturbances in a pup’s prepulse inhibition and latent inhibition [34]. In the PPI test, male adult rats showed disturbance in the startle response at both 5 dB and 10 dB, but female rats only suffered 5dB impairment on PPI of startle. In the EPM treatment, female rats spent more time in the open arm and travelled longer distance. While male rats moved anxiously around the open field seeking escape routes or an area of shelter; female rats seemed more bold and wise. Since spatial learning requires mGlu1R and mGlu5R signaling in the dorsal hippocampus [35], the male rats anxiety might be related to the down-regulated expression of mGlu1R and mGlu5R caused by maternal separation.

The neurodevelopment showed sex-specific relations to the aberrations in the HPA axis and neurological gene expression, suggesting that maternal separation could contribute to decreased hippocampal neurogenesis and altered neuroendocrine axis [36]. In summary, our study confirmed that maternal separation during early life is a chronic stress that could lead to behavior and neurochemical abnormalities in rats, and this model highlights the special growth environment that is influential in young adult human development.

Conclusion

Maternal separation in early life induced significant abnormalities in EEG and alterations in expression of hippocampus mGlu1R and mGlu5R with greater changes in the corticosteroid level in males and extended behavior abnormalities in young adult rats.

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Conflict of Interest

There is no conflict of interest in the study and the preliminary data was unpublished.

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