INTRODUCTION

Childhood is a critical period for brain development. Early-life neglect has been a social trend during global urbanization, leading to negative effects in both urban and rural areas. Exposure to early-life neglect induces early-life stress (ELS), which has extensive influences on the central nervous system, including behavior, emotion, and cognition, and could increase the risk of long-lasting effects on brain development and changes in personal characteristic, and may cause symptoms including cognitive impairment, anxiety, depression, addiction, and impulsivity. These effects are the result of gene × environment interactions.

Literature reports that multiple brain areas contribute to the pathogenesis of clinical manifestations, such as the amygdala,
fusiform gyrus, insula, striatum, and prefrontal cortex. Different brain areas and neural circuits participate in behavioral or pathological manifestations. It has been reported that dopamine and receptors in caudate-putamen of rodent striatum might contribute to object recognition behavior. Orbital frontal cortex (OFC) and medial prefrontal cortex (mPFC) play important roles in impulsivity. mPFC is an essential cortex area for complex cognition and emotion modulation, connected to diverse brain areas and forming afferent and efferent neural pathways. Liu et al. found that oligodendrocytes and myelination impairment of mouse mPFC played a role in social life stress. In previous work, we demonstrated changes in rat medial prefrontal cortex (mPFC) of an early-life neglect model. Chen et al. reported that the excitation/inhibition balance of mPFC layer V also contributed to depression and anxiety-like emotional disorders of rats. Intrinsic excitability of mPFC neurons acts as a neural basis of physiological functions, as well as their information integration and processing. In pathological states, the functional changes of neural excitability are considered neural mechanisms of symptom and behavioral phenotype.

In this research, we take advantage of maternal separation with early weaning (MSEW) as a rat model for early-life neglect, and decipher different neural excitability of GABAergic and glutamatergic neurons in rat mPFC by whole-cell patch clamp. Our findings suggest different neuronal excitability alterations of rat mPFC GABAergic and glutamatergic neurons in an early-life neglect model, which may be considered as potential therapeutic targets in the future.

2 MATERIAL AND METHODS

2.1 Animals

Four adult specific-pathogen-free (SPF) Sprague–Dawley female rats and 2 male rats were acquired from Beijing Huafukang Bioscience Co. Ltd. For rat breeding, every cage contained 1 male rat and 2 female rats. All the animals were housed at room temperature –24–26°C, 12-h/12-h light/dark cycle, with food and drink ad libitum. This project was approved by the Institutional Animal Care and Use Committee of ILAS (ILAS-QC-2017-002). All the experiment procedures were conducted in accordance with the ethical guidelines, 3R rules, and national standard.

2.2 Early-life neglect model of rats

Maternal separation with early weaning (MSEW) was regarded as an early-life neglect model of rats. Methods were described previously by George et al. Twenty litters, half male and half female, were obtained after breeding. From postnatal day (PND) 0, litters were randomly divided into 4 groups (each group n = 5): female MSEW group, female control group, male MSEW group, and male control group. Control mouse groups were undisturbed and weaned normally until PND 21. MSEW were separated from the dam every day as follows: PND 2–PND 5, separated for 4 h each day; PND 6–PND 16, separated for 8 hours each day; PND 17, weaned. Separated litters were kept in separate rat cages, without bedding. Temperature was kept at 32°C with heating pads. After weaning, litters were group-housed with the same sex and group.

2.3 Patch-clamp electrophysiology

2.3.1 Brain slice preparation

Whole-cell patch clamp of rat acute brain slices was used to examine the excitation of mPFC neurons. Five rats and 10 cells were recorded for each group. For patch-clamp slice preparation, rats were anesthetized by pentobarbital sodium intraperitoneally. The whole brain was rapidly dissected and submerged in oxygenated (95% O2/5% CO2), ice-cold high-sucrose cutting solution, containing (in mM): KCl 2.5, NaH2PO4 1.25, CaCl2 0.5, MgSO4 10, NaHCO3 26, glucose 10, sucrose 230 (pH 7.4, 300–310 mOsm). With a Leica VT1000S vibrating microtome (Leica Instruments, Germany), 250 μm coronal slices of medial prefrontal cortex (mPFC) were made. Brain slices were transferred to an incubation chamber with 32°C oxygenated artificial cerebrospinal fluid (ACSF), containing (in mM): NaCl 126, KCl 2.5, MgCl2 1.3, NaH2PO4 1.2, CaCl2 2.4, NaHCO3 18, glucose 10 (pH 7.4, 300–310 mOsm). Slices were equilibrated for more than 0.5 h at room temperature before recording.

2.3.2 Whole-cell patch-clamp recording

The excitation of mPFC neurons was examined by whole-cell patch-clamp recording. Brain slices were transferred to a recording chamber under an Olympus BX51 microscope (Olympus, Germany), while cell images were taken with an infrared differential interference contrast camera. During recording, slices were perfused 2 ml/min with regular ACSF. Borosilicate glass capillary tubes (Sutter 150-86-10, USA) were pulled by PC-10 pipette puller (Narishige, Japan). The resistance of glass pipettes was ~5–8 MΩ, filled with K+–Met-sulfonate internal solution containing (in mM): K+–Met-sulfonate 140.5, NaCl 7.5, HEPES hemisodium salt 10, Mg-ATP 2, Na-GTP 0.2 (pH 7.33, 300–310 mOsm). Data were recorded by a Multiclamp 700 B amplifier (Molecular Devices, Sunnyvale, CA), filtered at 10 kHz and sampled at 50 kHz, and later analyzed with pClamp 10.6 software (Molecular Devices, USA). Recordings were discarded if the series resistance increased by more than 20% during the course of the recordings. ACSF and solutions were applied to the slice via a peristaltic pump (Longer Pump, China).

To distinguish different cell types of rat mPFC, we run the firing test protocol under current-clamp mode. A 100-pA, 1-second current was injected in the neuron, and the number and frequency of firing was measured. To examine the excitability of different cells, a single and fast depolarizing current was injected to trigger a single
action potential (AP) (30 ms, 100 pA). To trigger a series of APs, a series of 600-ms current pulses was injected, with intensities between −200 and 400 pA in 100-pA increments. The rest membrane potential, single action potential, and number of multiple action potentials were analyzed and plotted as done previously.15,16

2.4 | Statistics

All data are presented as mean ± standard deviation (SD). Data were analyzed with unpaired t-test between 2 groups, and 2-way analysis of variation (ANOVA) for multiple groups. Graphpad Prism 8 software was used for data analysis, and \( p < 0.05 \) was considered significant.

3 | RESULTS

3.1 | Electrophysiological properties of different cell types in mPFC

To study different mechanisms of mPFC neurons during MSEW, we used whole-cell patch-clamp test to analyze mPFC neural functions in layer II/III (Figure 1A). The medial prefrontal cortex consists of glutamatergic and GABAergic neurons, which compose the 2 most basic circuits of mPFC, respectively the excitatory and inhibitory synapses.17 To separate these 2 types of mPFC neurons, before whole-cell states, these 2 cell types can be preliminarily distinguished by cell shape under phase-contrast microscopy. Glutamatergic neurons were almost pyramidal neurons in mPFC,

**FIGURE 1** Electrophysiological properties of putative glutamatergic and GABAergic neurons in medial prefrontal cortex. (A) Schematic coronal section of a rat brain, showing the medial prefrontal cortex (mPFC). Recording electrodes were placed in layer II/III of mPFC. Excitation of neurons were recorded by whole-cell patch clamp. (B–D) Summary histograms demonstrating different electrophysiological properties of mPFC glutamatergic and GABAergic neurons, regarding rest membrane potentials (B), action potential firing frequency (C), and halfwidth (D). (E) Typical example of a putative glutamatergic neuron from mPFC. Left, pyramidal shape of a mPFC glutamatergic neuron. Middle, multiple action potentials triggered by 1-ms depolarizing current of 100 pA. Right, a typical single action potential from the neuron in left panel. (F) Typical example of a putative GABAergic neuron from mPFC. Left, oval shape of a mPFC GABAergic neuron. Middle, multiple action potentials triggered by 1 ms depolarizing current of 100 pA. Right, a single action potential from the neuron in left panel.
appearing as a triangle-like shape with one main neurite (Figure 1E, left panel). GABAergic neurons in mPFC have an oval shape, and are smaller than glutamatergic ones (Figure 1F, left panel). Under whole-cell patch-clamp recording, these 2 cell types have distinct intrinsic electrophysiological properties in terms of neural rest membrane potential, innate action potential firing frequency, and halfwidth. Our results show that the rest membrane potential of glutamatergic neurons was $-72.84 \pm 3.003 \text{ mV}$, while that of GABAergic neurons was $-67.65 \pm 3.066 \text{ mV}$ (Figure 1B). With injection of depolarizing current during whole-cell recording, the intrinsic firing frequencies were typically $2.600 \pm 2.702 \text{ Hz}$ for glutamatergic neurons and $18.80 \pm 4.817 \text{ Hz}$ for GABAergic neurons (Figure 1C,E,F, middle panel). When triggering a single action potential, the halfwidth of glutamatergic neurons was $2.800 \pm 0.3000 \text{ ms}$, and $1.580 \pm 0.04472 \text{ ms}$ for GABAergic neurons (Figure 1D–F, right panel). On the basis of the above 3 characteristics, glutamatergic neurons and GABAergic neurons were distinguished from the mPFC brain slices for further patch-clamp recordings.

### 3.2 | MSEW generated functional changes in mPFC glutamatergic neurons

Under whole-cell current clamp, the rest membrane potential (RMP) of mPFC glutamatergic neurons was decreased in male rats but not female rats, from $-70.34 \pm 5.212 \text{ mV}$ in the male control group to $-63.37 \pm 6.295 \text{ mV}$ in the male MSEW group, facilitating the activation of these neurons (Figure 2A, Table 1). Similarly, when triggering a single action potential (AP) in these mPFC glutamatergic neurons, the threshold of the male MSEW group was increased compared with the male control group, from $-42.85 \pm 1.878$ to $-47.04 \pm 2.157 \text{ mV}$, which also facilitated the triggering of an action potential (Figure 2B). Taken together, the alterations in RMP and AP threshold showed that male MSEW rats produced an action potential more easily.

To comprehensively analyze the intrinsic excitation, we also injected a longer depolarizing current of 600 ms to trigger multiple action potentials. Results demonstrated that in male, but not female, rats, MSEW modeling altered the number of APs, causing alterations in firing frequencies and refractory periods (Figure 2D–F). This result is consistent with our previous research, where we demonstrated that selectively in male MSEW rats, but not female mice, mPFC glutamatergic neurons exhibited a decrement in the number of multiple action potentials triggered by a series of 600-ms depolarizing current (0, 100, 200, 300, 400 pA). This decreased excitability between male MSEW rats and male control rats contributed to the anxiety-like behavior of male MSEW rats.

For female rats, MSEW modeling only increased the amplitude of a single AP (43.40 ± 6.991 mV in the female control group,

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**FIGURE 2** Distinct alterations in glutamatergic neuronal excitation between male and female rats. (A) Altered rest membrane potentials of mPFC glutamatergic neurons between male control and male MSEW group. (B) Altered threshold of a single action potential of mPFC glutamatergic neurons between male control and male MSEW group. (C) Increased amplitude of a single action potential of mPFC glutamatergic neurons in female MSEW group. (D) Multiple action potentials of 4 groups. Decreased number of APs between male groups. (E–H) Representative multiple action potentials of the 4 groups (*p < .05; **p < .01, MSEW group compared with control group. N = 10 cells for each group).
and 56.75 ± 12.06 mV in the female MSEW group), without other changes in electrophysiological properties (Figure 2C,G,H).

3.3 Different alterations in excitability of mPFC GABAergic neurons

Different from the above-mentioned functional changes in mPFC glutamatergic neurons, MSEW modeling led to multiple changes in mPFC GABAergic neurons.

Under whole-cell current clamp, the rest membrane potential (RMP) of mPFC GABAergic neurons was decreased in female rats but not male rats, from −69.35 ± 2.669 mV in the female control group to −64.17 ± 4.602 mV in the female MSEW group, facilitating the activation of these neurons (Figure 3A, Table 2).

For the thresholds of a single action potential, MSEW modeling decreased the threshold in both male and female groups: in male groups, −52.02 ± 0.6987 mV in the male control group and −47.11 ± 1.474 mV in the male MSEW group; in female groups, −55.97 ± 1.786 mV in the female control group and −46.51 ± 2.803 mV in the female MSEW group (Figure 3B).

For the amplitude of a single action potential, MSEW modeling only increased the amplitude in the male group, from 33.20 ± 0.5492 mV in the male control group to 39.40 ± 5.648 mV in the male MSEW group (Figure 3C). To comprehensively analyze intrinsic excitation, we also injected a longer depolarizing current of 600 ms to trigger multiple action potentials. Results demonstrated that in male, but not female, rats, MSEW modeling altered the number of APs, which caused alterations in firing frequency and refractory periods (Figure 3D-F).

**TABLE 1** Summary of intrinsic physiological properties of mPFC glutamatergic neurons

|                          | Male control | Male MSEW | Female control | Female MSEW |
|--------------------------|--------------|-----------|----------------|-------------|
| Rest membrane potential (mV) | −70.34 ± 5.212 | −63.37 ± 6.295** | −73.70 ± 11.65 | −70.61 ± 7.90 |
| Voltage threshold (mV)    | −42.85 ± 1.878 | −47.04 ± 2.157b | −44.76 ± 8.785 | −45.02 ± 6.365 |
| Action potential amplitude (mV) | 43.62 ± 3.016 | 42.77 ± 4.928 | 43.40 ± 6.991 | 56.75 ± 12.06^a |

Note: Data are shown as mean ± SD from 10 mPFC cells from 5 rats.
aControl group compared with MSEW group.  
\( p < 0.05. \)

**Control group compared with MSEW group.  
\( p < 0.01. \)

**FIGURE 3** Distinct alterations in GABAergic neuronal excitation between male and female rats. (A) Altered rest membrane potentials of mPFC GABAergic neurons between female control and female MSEW group. (B) Altered threshold of a single action potential of mPFC GABAergic neurons between control and MSEW groups. (C) Increased amplitude of a single action potential of mPFC glutamatergic neurons in male MSEW group. (D) Multiple action potentials of 4 groups. Decreased number of APs in between male groups. (E–H) Representative multiple action potentials of the 4 groups (* \( p < .05; \) ** \( p < .01; \) **** \( p < .0001, \) MSEW group compared with control group. \( N = 10 \) cells for each group).
In our research, we found that, in an MSEW rat model, female rats has been demonstrated to facilitate the excitation of mPFC neurons. The firing response of the mPFC neurons to injected depolarizing currents can also be analyzed. Besides, for the first AP in a series of multiple APs, the amplitude, halfwidth, and afterhyperpolarization (AHP) rates was examined. Rheobase refers to the minimum injected current that can trigger AP. Other membrane properties of AP can also be measured, such as rest membrane potential and input resistance. The excitability of neurons can be verified by different aspects. Under the current mode of whole-cell patch clamp, the firing response of the mPFC neurons to injected depolarizing currents was examined. Rheobase refers to the minimum injected current that can trigger AP. Other membrane properties of AP can also be measured, such as rest membrane potential and input resistance. 

Medial prefrontal cortex (mPFC) consists of excitatory glutamatergic neurons and inhibitory GABAergic neurons. Different types of neurons compose excitatory or inhibitory synapses. In our results, male MSEW rats had facilitated glutamatergic neurons, while the GABAergic neurons were relatively stable. In female rats, glutamatergic neurons were stable (except for higher AP amplitude), while triggering of AP in GABAergic neurons was not easy as the threshold changed. Taken together, we found that, although the results looked controversial between male/female and glutamatergic/GABAergic, the overall excitatory/inhibitory (E/I) ratio had similar trends in both male and female rats.

Regarding the E/I ratio, the results were similar, but by different implementation: male rats by more excited glutamatergic neurons, female rats by less excited GABAergic neurons. In the future, the exact E/I ratio can be examined by ratio of (m/s) EPSC frequency to (m/s) IPSC frequency.

Sex differences in the nervous system due to estrous cycle, sex hormones, and their receptors have gained increasing attention in recent years. Yousuf et al. demonstrated that infralimbic estradiol could enhance neuronal excitability of female rats, which could be a putative reason for the sexually dimorphic properties. Potent estrogen, 17β-estradiol (E2), can be directly infused into the mPFC area, and has been demonstrated to facilitate the excitation of mPFC neurons. In our research, we found that, in an MSEW rat model, female rats showed decreased excitation in GABAergic neurons and increased AP amplitude of glutamatergic neurons. Combining these 2 alterations, it can be inferred that the excitation of mPFC as a whole can be increased by estrogen. In terms of E/I ratio, estrogen might result in increased excitatory synapse and decreased inhibitory synapse. In summary, our research found that an early-life neglect model of rats had different alterations of electrophysiological properties in mPFC glutamatergic and GABAergic neurons: male MSEW rats mainly by glutamatergic neurons, and female MSEW rats by GABAergic neurons.

4 | DISCUSSION

In this work, we chose maternal separation with early weaning (MSEW) as a rodent model of early-life neglect. Since early-life stress has shown increasing impact on brain development and physiological functions, we tried to find related brain areas, cell types, and neural electrophysiological mechanisms.

The excitability of neurons can be verified by different aspects. Under the current mode of whole-cell patch clamp, the firing response of the mPFC neurons to injected depolarizing currents was examined. Rheobase refers to the minimum injected current that can trigger AP. Other membrane properties of AP can also be measured, such as rest membrane potential and input resistance. Besides, for the first AP in a series of multiple APs, the threshold, amplitude, halfwidth, and afterhyperpolarization (AHP) can also be analyzed.

Medial prefrontal cortex (mPFC) consists of excitatory glutamatergic neurons and inhibitory GABAergic neurons. Different types of neurons compose excitatory or inhibitory synapses. In our results, male MSEW rats had facilitated glutamatergic neurons, while the GABAergic neurons were relatively stable. In female rats, glutamatergic neurons were stable (except for higher AP amplitude), while triggering of AP in GABAergic neurons was not easy as the threshold changed. Taken together, we found that, although the results looked controversial between male/female and glutamatergic/GABAergic, the overall excitatory/inhibitory (E/I) ratio had similar trends in both male and female rats. Regarding the E/I ratio, the results were similar, but by different implementation: male rats by more excited glutamatergic neurons, female rats by less excited GABAergic neurons. In the future, the exact E/I ratio can be examined by ratio of (m/s) EPSC frequency to (m/s) IPSC frequency.

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