Role of the prefrontal cortex in the neonatal ventral hippocampus lesion, an animal model of schizophrenia

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Introduction

Schizophrenia is a mental disorder (DSM-5) that affects approximately 1% of the world population. This disorder is not only devastating for the patient but also affects the family. Its prognosis depends on the number of psychotic outbreaks, progression of cognitive impairments and response to neuroleptics. This psychotic disorder is characterized by the presence of symptoms classified as positive (increased locomotion in response to psychostimulants, hallucinations, delusions and thought disorders), negative (deficits in social interaction, anhedonia, and affective flattening, among others) and cognitive deficits (attention and memory deficits). In recent years, with the advancement in the image analysis and microscope systems, the neural morphological studies in postmortem tissue from schizophrenic patients have shown constant and critical changes in the prefrontal cortex (PFC), which may explain some of the symptoms observed in this complex disorder. Moreover several animal models of schizophrenia have been developed over the past 20 years. Although there is no animal model that can replicate all aspects of complexity that occurs in schizophrenia, but they can correlate changes in the animal behavior with the classification of the symptoms of schizophrenia. An increased expression of a behavior is a positive symptom, such as increased locomotor activity in a new environment. Negative symptoms involve reduction in certain behaviors, such as grooming. In addition, some behaviors may relate directly, such as deficits in social interaction, attention...
and memory. One of the most frequently used animal model of schizophrenia is the neonatal ventral hippocampus lesion (nVHL)\textsuperscript{1,3}. In the present mini-review, we discuss neurochemical and morphological changes in the PFC as reported in the nVHL rat.

**Prefrontal cortex connections in schizophrenia**

Numerous studies from different domains (clinical, postmortem, neuroanatomical and physiological) link the PFC with the pathophysiology of schizophrenia\textsuperscript{1,4}. Indeed, the PFC is a complex region of sensorimotor and emotional integration, which participates in several cognitive functions such as attention processing, spatial memory, decision making etc. This cortical region receives glutamatergic projections from thalamus (dorsomedial nucleus), hippocampus and basolateral amygdala (BLA)\textsuperscript{1}; dopaminergic projections from the ventral tegmental area (VTA); serotoninergic projections from the median raphe nucleus; noradrenergic projections from locus coeruleus and cholinergic projections from basal forebrain. To close the circuit, PFC sends glutamatergic projections to nucleus accumbens (NACC), CA1 region of the ventral hippocampus, BLA, VTA, and thalamus. Moreover, a recent report suggests that the connection among the PFC, the lateral hypothalamus (LH) and the periaqueductial gray matter (PAG) together with the brainstem nuclei form a circuit that regulates the expression of positive emotions\textsuperscript{5}. Interestingly, schizophrenic patients fail to express positive or negative emotions and are also unable to recognize emotions when presented with various faces\textsuperscript{6,7}, hence it is possible to suggest that the PFC-LH-PAG circuit is dysfunctional in patients with schizophrenia.

It has been suggested that the connections between the thalamus and PFC are made during fifth month of gestation in humans, while the connections among the ventral hippocampus, BLA and PFC start to form at seventh month of gestation. Several reports suggest that the disruption between PFC and hippocampus at an early age is involved in the etiology of schizophrenia\textsuperscript{8-10}. Rats that underwent nVHL at postnatal day (PND) 7 is an example of the early disruption between hippocampus and PFC\textsuperscript{1,3,8}. Furthermore, this animal model shows behavioral, neurochemical and morphological changes that manifest mainly after puberty\textsuperscript{1,3,11}.

**The nVHL Animal Model**

The nVHL rat is considered to be a neurodevelopmental model of schizophrenia\textsuperscript{1,3}. This model was conceived by Lipska and Weinberger in the 1990’s to address questions that pharmacological models could not answer\textsuperscript{12}. First, the nVHL rat presents a disruption in the connections between the PFC and hippocampus in a critical postnatal period. Second, as stated in O’Donnell et al.\textsuperscript{13} “the perinatal period is critical for maturation of PFC circuits” and recent research strongly suggests that adolescence is a critical period for functional organization in the PFC with high rate of synaptic pruning\textsuperscript{14}. Indeed, juvenile nVHL rats present normal behaviors when compared to sham animals\textsuperscript{12,15} suggesting that the structural changes occur at this age. Third, after puberty, a constellation of behavioral, neurochemical and neuroanatomical changes are apparent in the nVHL rat\textsuperscript{1}. The nVHL rat presents normal behaviors until the age of young adult as observed in schizophrenic patients. Therefore this animal model is suitable to further our understanding of this complex disorder.

**Neurochemical and neuromorphological changes in the nVHL rat**

The nVHL induces a constellation of neurochemical and neuronal alterations in the PFC as discussed below.

**Neurochemical alterations:** Overactivity of dopamine (DA) and glutamate in the mesolimbic system is the major working hypothesis of the etiology of schizophrenia\textsuperscript{1,3,16,17}. In this regard, while Alquicer et al. (2004)\textsuperscript{18} observed reduced DA content in the PFC, exogenous application of SKF38393, a D\textsubscript{1} agonist, increases cell firing of PFC pyramidal cells of the nVHL rat\textsuperscript{19}. SKF38393 selectively increases the release of acetylcholine in the PFC of the nVHL rat at post-pubertal age\textsuperscript{20}. Whereas ventral tegmental area (VTA) stimulation increases cell firing in the PFC of the nVHL rat\textsuperscript{21}. In a food paradigm in vivo, the nVHL rat maintains DA outflow for a longer period of time in the PFC\textsuperscript{22}. Moreover, the nVHL rat presents enhanced sensibility to the DA agonist, amphetamine\textsuperscript{1}. Regarding the glutamatergic system, the nVHL rats present hyperresponsiveness to MK-801, an n-methyl-D-Aspartate (NMDA) glutamatergic antagonist, showing no apparent changes in the expression of glutamatergic receptors\textsuperscript{15}. Moreover, bath applications of NMDA also increases cell firing in the PFC of the nVHL rat\textsuperscript{13}. The present data show hyper- and hypo-activity of DA in the PFC of the nVHL rat is observed most likely derived from contextual factors. Therefore, abnormal DA and glutamate systems alter the response to environmental or pharmacological challenges in the nVHL rat.

The inhibitory aminoacid gamma-aminobutyric acid (GABA) is consistently reduced in cortical areas in postmortem brains of schizophrenic patients\textsuperscript{23}. Moreover, an abnormal balance between excitation and inhibition is commonly observed in psychiatric disorders\textsuperscript{24}. In clear contrast, imaging studies have shown mixed results (increased, decreased or unaltered GABA levels\textsuperscript{25}). Despite of the importance of GABA in the PFC, investigations of the levels or activity of GABA in the PFC of the nVHL rat has been minimal. Recently, Ryan et al.\textsuperscript{26} showed no significant difference in vesicular GABA transporter in the PFC of adult nVHL rat. However, further research is certainly warranted.
Nitric oxide (NO) is an inter- and intra-cellular messenger involved in physiological processes including synaptic and neuronal plasticity. Moreover, schizophrenic subjects present increased level of plasma NO. The nVHL rat also presents increased levels of NO as well as increased nitric oxide synthase (NOS) immunostaining in the PFC. In fact, NO has emerged as a key player to increased nitric oxide synthase (NOS) immunostaining neurons in the PFC. In the PFC of the nVHL rats, it has been established that NO interacts with DA and glutamate neurotransmitters.

Animals that underwent the nVHL also present dysregulated proteins such as clathrin light chain B, a protein important for the expression of synaptophysin after puberty; low level of nerve growth factor inducible-B mRNA and reduced expression of brain-derived neurotrophic factor (BDNF) mRNA. In apparent agreement, patients with schizophrenia also exhibit these neurochemical alterations in the PFC including low expression of synaptophysin and decreased immunoreactivity to BDNF. Consequently, all these neurotransmitters, growth factors and proteins have an impact on the synaptic connections that ultimately reshape neurons in the PFC.

**Neuromorphological alterations**

In postmortem brains of patients with schizophrenia, Garey et al. observed reduced spine number in PFC layer III pyramidal neurons. While Koleske indicated that the shape of dendritic arbor determines the number and distribution of receptive synaptic contacts; Fiala et al., suggested that the dendritic spines are the main sites of excitatory input. Therefore alterations in spine density or dendritic arbor are associated with gain or loss of connectivity. Our group has shown a decrease in dendritic length and dendritic spines not only in the PFC layer V but also layer III in the nVHL rat. The neuronal hypotrophy observed in PFC neurons has been associated with lack of input from the ventral hippocampus at a critical stage of development.

Earlier disconnection (PD7 – PD9) of the ventral hippocampus and PFC pathway causes behavioral, neurochemical and neuromorphological changes after puberty or in early adulthood, which does not manifest itself if the damage had occurred before PD14. All this implies that there is a window of time in which the injury of this pathway may cause permanent damage. Moreover, the neonatal lesion of PFC, also causes changes in behavior, neurochemistry and morphology, however these changes are not as copious as in the case of nVHL animals. Interestingly, at adult age, lesion of the PFC in nVHL animals, ameliorate behavioral changes. Moreover, earlier lesion of VH altered the physiological response of PFC pyramidal neurons after puberty, which exhibited excessive firing in response to mesocortical stimulation. Therefore, earlier disruption of the VH and PFC pathway affected the function of PFC after puberty, such as reduced levels of the BDNF in the PFC of the nVHL rats.

**Conclusions**

The nVHL rat presents neurochemical and neuroanatomical alterations in the PFC similar to those observed in schizophrenic patients. Early on, O'Donnell stated that "disinhibited PFC could be responsible for cognitive deficits observed in schizophrenia". Moreover, PFC refinement occurs during late adolescence and the behavioral deficits in this animal model are also observed only after this period. Further investigations in this animal model are required to understand the pathology of this devastating disorder as well as to test potential novel drugs for the treatment of different deficits observed in schizophrenia.

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