PERSPECTIVE

Cell genesis and dendritic plasticity: a neuroplastic pas de deux in the onset and remission from depression

A Mateus-Pinheiro¹,², P Patrício¹,², JM Bessa¹,², N Sousa¹,² and L Pinto¹,²

Brain neuroplasticity is increasingly considered to be an important component of both the pathology and treatment of depressive spectrum disorders. Recent studies shed light on the relevance of hippocampal cell genesis and cortico-limbic dendritic plasticity for the development and remission from depressive-like behavior. However, the neurobiological significance of neuroplastic phenomena in this context is still controversial. Here we summarize recent developments in this topic and propose an integrative interpretation of data gathered so far.

Keywords: antidepressants; cell genesis; depression; neuroplasticity; stress

INTRODUCTION

The potential to respond to environmental stimuli through dynamic rearrangements of synapto-dendritic networks, as well as by regulating the generation of new neuronal and glial cells, renders the brain highly mutable. These phenomena, collectively known as neuroplasticity, are critical to promote neuronal adaptations; its failure is now increasingly considered to be a major component in many neuropsychiatric conditions. Among these, depressive spectrum disorders are a paradigmatic example of the importance of neuroplastic alterations in the adult brain. Recent studies provide a comprehensive picture of the effects of stress, a major trigger factor in depression, in the (de)regulation of molecular and physiological mechanisms regulating neuroplastic processes are relevant for the onset of depressive symptoms, they have also been implicated in the action of antidepressants (ADs). So far, and although there is still much to be elucidated, it is becoming evident that the triad stress-neuroplasticity-depression constitutes fertile ground for new findings.

NEW CELLS AND DENDRITES: IMPORTANCE FOR THE TREATMENT AND REMISSION FROM DEPRESSION

Although different forms of neuroplasticity are affected in depression, a debate endures concerning the exact neurobiological significance of postnatal hippocampal cell genesis, both for the development of depressive pathology and for the therapeutic action of ADs. From the bulk of evidence gathered so far, it is increasingly appreciated that alterations in cell genesis are involved in the pathology and treatment of depression; however, there are several conflicting reports regarding its relevance. Three factors are likely to contribute to these controversies. First, there is a ‘necessary’ difficulty to approach this question in humans suffering from depression; postmortem studies in humans and animal models of depression have, nevertheless, provided important insights. Second, it seems to exist a major prevalence of studies focusing on the functional implications of neurogenesis, in disregard of gliogenesis, a parallel cell-genesis process likely to be of relevance in this context. Lastly, because these events are highly dynamic, the adoption of different experimental models and time frames when analyzing the participation of cell genesis in the pathology and treatment of depression is critical to have a complete perspective of the topic. On account of these experimental dissimilarities, an integrative, and careful, interpretation of data published in the last years is required when attempting to put the pieces of the puzzle together.

Suppression of hippocampal cell proliferation in naive animals through irradiation, pharmacological approaches or through the use of transgenic models of cytogenesis ablation has been shown to be associated with the development of deficits in different behavioral dimensions commonly affected in depression. Strikingly, most of the studies in which analyses were performed shortly after cytogenesis ablation did not reveal significant deficits in most behavioral domains normally assessed in the characterization of animal models of depression (Figure 1). However, recent reports in which abrogation of cytogenesis is maintained for long periods (over 4 weeks) or in which the behavioral analysis was conducted only 4 weeks after the cessation of cytogenesis suppression, reported multidimensional behavioral deficits that emerged only weeks after the antiproliferative insult. Importantly, the specific late manifestation of depressive-like behavior and cognitive disabilities in animals in which cytogenesis had been suppressed illustrates how manipulating lengthy neuroplastic phenomena is associated with the non-immediate development of behavioral impairments, which are only fully manifested once newborn cells are expected to be incorporated in local neuroglial circuits. This view has been recently supported by the demonstration that the specific inhibition of 4-week old new hippocampal neurons causes deficits in memory retrieval in mice; remarkably, inhibiting the activity of either younger or less-plastic older neurons does not produce effects in this cognitive domain.

¹Life and Health Sciences Research Institute (ICVS), School of Health Sciences, University of Minho, Braga, Portugal and ²ICVS/3B’s—PT Government Associate Laboratory, Braga/Guimarães, Portugal. Correspondence: Professor N Sousa or Dr L Pinto, Life and Health Sciences Research Institute (ICVS), School of Health Sciences, University of Minho, Braga 4710-057, Portugal.
E-mail: njcsousa@ecsaude.uminho.pt or luisapinto@ecsaude.uminho.pt
Received 10 January 2013; revised 1 April 2013; accepted 8 April 2013; published online 28 May 2013
The participation of dendritic plasticity and hippocampal cell genesis in the development and treatment of depression. (a) In animal models of depression, chronic exposure to stress, one major trigger factor of depression, leads to the dendritic atrophy of hippocampal and prefrontal cortex neurons and compromises the generation of new cells. These neuroplastic impairments are associated with the development of multidimensional deficits manifested in behavioral dimensions such as mood, anxiety behavior and cognitive performance. Treatment with typical SSRIs allows for the fast reestablishment of dendritic neuronal architecture, which is associated with the fast remission from depression-like behavior. The slow generation of new neuronal and glial cells, potentiated by antidepressant (AD) treatment will be fundamental to maintain this remission and to promote a sustained long-term recovery from depression. (b) Suppression of hippocampal cell genesis, using X-ray irradiation, the cytostatic agent methylazoxymethanol or transgenic animal models of cytogenesis ablation, does not seem to have an impact on the mood-improving effects of ADs, but rather to preclude AD’s efficacy in reversing anxiety behavior and impairments in some cognitive modalities (blue line); other cognitive functions remain unaffected (dashed blue line). (c) Ablation of cytogenesis in naïve animals, while maintaining dendritic morphology intact, leads to the short-term manifestation of hyperanxiety behavior; deficits in mood and cognitive modalities are only evident later on, when newborn cells are expected to be fully matured and incorporated in the local neurocircuitry. SSRIs, selective serotonin reuptake inhibitors.
An exception must be made in respect to anxiety behavior, because disruption of hippocampal neurogenesis is associated to the immediate development of heightened anxiety, which is commonly comorbid in depressed patients. In fact, it has been demonstrated that immature newborn neurons display a major role in anxiety behavior control.10–12

In contrast with the slow reconfiguration of neural networks promoted by the addition of new cells in the adult brain, are the rapid synaptic and dendritic morphological changes. These underlie the short-term impacts on distinct emotional and cognitive processes observed in the onset of depressive symptoms. Indeed, defects on neuronal cytoarchitecture have been documented in postmortem analysis of brain tissue from depressive patients, which seem to be ameliorated by chronic AD treatment.5,11,14 These structural defects include cortico-limbic dendritic atrophy in depressed subjects,15 accompanied by abnormalities in glial cell structure. Importantly, in animal models of depression, these changes are associated with hallmarks of depressive behavior, such as anhedonic behavior, behavioral despair and cognitive disabilities.2

Besides this common participation of slow cytogenesis processes and rapid dendritic rearrangements in the pathology of depressive spectrum disorders, these two neuroplastic phenomena, with very different temporal dynamics, constitute the substrate by which ADs (partially) exert their therapeutic actions. Thus, the reestablishment of normal neural networks seems to be achieved in a biphasic manner (Figure 1): in a short-term context, AD actions rely on rapid modulatory effects upon genes involved in the restructuring of the synaptic network; later on, the generation of new fully matured neuronal and glial cells will have an impact on the long-term remission from emotional and cognitive disabilities manifested during a depressive episode. In fact, despite triggering an immediate pro-proliferative response, this early effect corresponds only to the onset of a slow neuroadaptation whose neurobiological importance can only be fully appreciated later on, once new cells attain complete maturation and functionality, and are integrated in the local neurocircuity. Taken together, and because mammalian neurogenesis is described to take 4–6 weeks, the overall outcome of AD’s therapeutic action upon neuroplasticity may only be entirely manifested after this period. Remarkably, this period correlates with the time latency that typically prescribed ADs take to fully manifest their action in depressive patients.

Although the neuroplastic alterations occurring during the onset, treatment and remission from depression are being increasingly characterized, comprehension of the processes (namely genetic and epigenetic programs) that orchestrate these alterations is still limited. Future research focusing on these processes should also be extended to the still underexplored glioplastic component of this disorder. Furthermore, local neuroplastic adaptations are likely to occur in articulation with systemic neuroendocrine and immunological alterations, which are still to be integrated in the complex puzzle of mechanisms implicated in depression.

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