A morphometric study of the progressive changes on NADPH diaphorase activity in the developing rat’s barrel field

Marco Aurélio M. Freire a, Walace Gomes-Leal a, Walther A. Carvalho a, Joanilson S. Guimarães a, João G. Franca c, Cristovam W. Picanço-Diniz a,∗, Antonio Pereira Jr. b,1

a Laboratory of Functional Neuroanatomy, Department of Morphology, Federal University of Pará, 66075-900 Belém, PA, Brazil
b Department of Physiology, Federal University of Pará, 66075-900 Belém, PA, Brazil
c Institute of Biophysics Carlos Chagas Filho, Federal University of Rio de Janeiro, Rio de Janeiro, RJ, Brazil

Received 25 March 2004; accepted 27 May 2004

Available online 2 July 2004

Abstract

The distribution of NADPH diaphorase (NADPH-d)/nitric oxide synthase (NOS) neurons was evaluated during the postnatal development of the primary somatosensory cortex (SI) of the rat. Both cell counts and area measurements of barrel fields were carried out throughout cortical maturation. In addition, NADPH-d and cytochrome oxidase (CO) activities were also compared in both coronal and tangential sections of rat SI between postnatal days (P) 10 and 90. Throughout this period, the neuropil distributions of both enzymes presented a remarkable similarity and have not changed noticeably. Their distribution pattern show the PMBSF as a two-compartmented structure, displaying a highly reactive region (barrel hollows) flanked by less reactive regions (barrel septa). The number of NADPH-d neurons increased significantly in the barrel fields between P10 and P23, with peak at P23. The dendritic arborization of NADPH-d neurons became more elaborated during barrel development. In all ages evaluated, the number of NADPH-d cells was always higher in septa than in the barrel hollows. Both high neuropil reactivity and differential distribution of NADPH-d neurons during SI development suggest a role for nitric oxide throughout barrel field maturation.

© 2004 Elsevier Ireland Ltd and The Japan Neuroscience Society. All rights reserved.

Keywords: Somatosensory cortex; Barrel field; PMBSF; NADPH diaphorase; Nitric oxide; Cytochrome oxidase; Cortical development; Rat

1. Introduction

In mammals, specializations of the sensory periphery influence the organization of the central nervous system (CNS) during development, sculpting sensory maps which reflect the distribution and diversity of receptors (Killackey et al., 1995). In the isocortex, cells in sensory areas are normally organized in a columnar fashion from pial surface to white matter, effectively breaking up the peripheral representation into a mosaic of processing modules (Mountcastle, 1997).

In some small rodents, this modular organization is most evident in layer IV of the primary somatosensory cortex (SI), where cell aggregates, aptly called barrels, are found (Woolsey and van der Loos, 1970; Welker, 1971; van der Loos and Woolsey, 1973). Individual barrels aggregate to form regular arrays in the somatosensory cortex representing the entire contralateral periphery (see Rice, 1995 for review). The largest barrels are located at the posteromedial barrel sub-field (PMBSF) and are neatly organized in rows which reproduce the organization of the mystacial vibrissae, a highly specialized tactile organ (Glassman, 1994).

Since barrels are regions with high metabolic rates (Mölter and Smith, 1998), they can be easily identified by simple histochemical techniques that reveal the activity of metabolic enzymes such as succinic dehydrogenase (SDH) and cytochrome oxidase (CO) (Wallace, 1987). CO is a key enzyme in the final steps of oxidative metabolism...
The analysis of its distribution throughout the nervous system has allowed the discrimination of individual brain areas and the identification of modules within some of them (Wong-Riley, 1989). In the rodent barrel field, CO activity is first detected around postnatal day 4 (P4), with a pattern very similar to the classical Nissl stain in adult animals and has been used as a trustworthy method to reveal the organization and morphology of barrel field (Wong-Riley and Welt, 1980; Vercelli et al., 1999).

Since the early 1960's it has been known that the enzyme nicotinamide adenine dinucleotide phosphate diaphorase (NADPH-d) reveals a selective neuronal sub-population of interneurons (Thomas and Pearse, 1961). Lately, it has been described a co-localization between NADPH-d and nitric oxide synthase (NOS) in this neuronal sub-population (Hope et al., 1991; Dawson et al., 1991; Luth et al., 1995). NOS is the neuronal enzyme responsible for the synthesis of nitric oxide (NO), a highly diffusible molecule involved in several physiological and pathological processes in the CNS (Iadecola, 1993; Wallace et al., 1996; Estevez et al., 1998; see Contestabile, 2000 for review). Accordingly, several groups have effectively used NADPH-d histochemistry as an indirect method for revealing the distribution of NO in the CNS, especially in fixed tissue (Matsumoto et al., 1993).

NADPH-d histochemistry/NOS immunohistochemistry reveals two subtypes of neurons (Luth et al., 1994). Type I neurons are intensely stained. In addition, they possess large cell bodies and a rich pattern of dendritic arborization. These cells comprise about 2% of the entirety of cortical neurons (Gabbott and Bacon, 1995). Type II neurons, on the other hand, are generally more numerous and faintly stained. Contrary to Type I cells, they have small cell bodies and few or no labelled processes (Sandell, 1986; Luth et al., 1994; Yan and Garey, 1997). The presence of Type I neurons was described in virtually all species studied so far, but Type II cells seems to be absent in lower vertebrates (Luebke et al., 1992).

Numerous studies have investigated the distribution of NADPH-d neurons in different regions of the developing and adult rat CNS (Vincent and Kimura, 1992; González-Hernandez et al., 1993; Tomic et al., 1994; Table 1).

Table 1
Summary of animals and procedures used in the present study

<table>
<thead>
<tr>
<th>Postnatal age</th>
<th>Number of animals</th>
<th>Section plane</th>
<th>Thickness (μm)</th>
<th>Histological technique</th>
</tr>
</thead>
<tbody>
<tr>
<td>P10</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P14</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P18</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P23</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P25</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P29</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P31</td>
<td>04</td>
<td>Tangential/coronal</td>
<td>100</td>
<td>NADPH-d/CO</td>
</tr>
<tr>
<td>P90</td>
<td>02</td>
<td>Coronal</td>
<td>50</td>
<td>NADPH-d/NOS</td>
</tr>
</tbody>
</table>

*In the tangential sections, CO histochemistry was carried out in contralateral hemispheres. In the coronal plane, CO was made in alternate sections reacted for NADPH-d.
flattened between two glass slides and immersed in PB overnight. The flattened brains were cut tangentially in 100 μm thick sections using a vibratome (Pekko International, Series 1000). In some P90 hemispheres, coro-
nal sections were cut 50 μm thick for double staining with NADPH-d and NOS immunocytochemistry (see Table 1).

2.3. NADPH-d histochemistry

To reveal NADPH-d activity, brain sections were collected and washed three times in PB and incubated, free-floating, in a solution containing 0.6% malic acid, 0.05% nitroblue tetrazolium, 0.03% manganese chloride, 0.5% β-NADP and 1.5–3% Triton X-100 in 0.1 M TRIS buffer, pH 8.0 (modified from Scherer-Singler et al., 1979). The histochemical reaction was monitored every 30 min to avoid overstaining. Sections were incubated at the same time and in the same solution for all animals. The incubation time ranged from 7 to 8 h for all ages evaluated and was usually interrupted by rinsing sections five times in 0.1 M PB (pH 7.4). All reagents were purchased from Sigma Company, USA. Sections were mounted in gelatin-coated glass slides, left to air-dry overnight, dehydrated and coverslipped with Entellan (Merck, Germany).

2.4. Immunocytochemistry

In order to evaluate the relationship between NADPH-d and NO in the barrel field, NOS immunocytochemistry was performed in some sections previously reacted for NADPH-d. These sections were washed three times in 0.1% PB saline (PBS)/Tween and incubated in 10% normal goat serum in PBS for 1 h. After that, sections were incubated in mouse anti-NOS primary antibody (dilution at 1:150 in PBS, Serotec, UK) for about 48 h at 10°C, washed three times in PBS/Tween and incubated in biotynilated goat anti-mouse secondary antibody (dilution at 1:150 in PBS, Serotec, UK) for about 1 h. Free-floating sections were then washed three times in PBS/Tween, and incubated at room temperature under constant agitation in a solution containing two drops of A and B solutions of Elite ABC Kit (Vector Laboratories) per 10ml of histochemical incubation media for 1 h. In this report, we used the DAB/Nickel method for revealing the antibody presence in the tissue, as described elsewhere (Shu et al., 1988; Gomes-Leal et al., 2002).

2.5. Cytochrome oxidase histochemistry

The CO histochemistry was used as a benchmark of the relative position and organization of the barrel field in all ages evaluated. In order to reveal CO activity (Wong-Riley, 1979) in both coronal (adjacent sections containing the barrel fields) and tangential brain sections (contralateral hemispheres) were incubated free-floating in a solution containing 0.05% diaminobenzidine (DAB), 0.05% cytochrome c and 0.02% catalase in 0.1 M PB. The CO histochemical reaction was monitored every 30 minutes in order to avoid overstaining. Similar to NADPH-d histochemistry, sections were incubated at the same time and in the same solution for all animals. The duration of the incubation ranged from 7 to 8 h for all ages evaluated and was usually interrupted by rinsing sections five times in 0.1 M PB (pH 7.4). All reagents were purchased from Sigma Company, USA. Sections were mounted in gelatin-coated glass slides, left to air-dry overnight, dehydrated and coverslipped with Entellan (Merck, Germany).

2.6. Qualitative and quantitative analyses

A complete reconstruction of the PMBSF, revealed by NADPH-d histochemistry, with the relative position of NADPH-d type I neurons in the barrel field was made using the software NeuroLucida (MicroBrightField Inc., USA). For each age evaluated (Table 1), three 100 mm thick tangential sections at the level of the barrel fields per animal (n = 3 animals per postnatal age) were reconstructed and superimposed using the blood vessels as landmarks. This procedure allowed complete reconstructions of the PMBSF. Since reactive neuropil forms barrels across approximately 300 μm vertically in sections from the intermediate levels of the gray matter (Zilles and Wree, 1985), we were able to unfold the entire barrel height in three 100 μm thick sections. The PMBSF areas of rats of different ages were measured using an image processing software (Scion Image for Windows, version Beta 4.0.1, Scion Corporation, USA) based on the complete reconstructions made in NeuroLucida. First, the outermost limits for the PMBSF were delineated in order to calculate the total PMBSF area. The area for each individual barrel was also measured. The area occupied by septa was obtained for the difference between the total PMBSF area and the sum of all individual barrel areas.

The relative position of the NADPH-d neurons in the PMBSF was plotted in the same nine tangential sections per age (three per animal), in which the PMBSF has been previously reconstructed to further quantitative analysis. This approach enabled us, based in the complete reconstructions of the PMBSF through layer IV, evaluate the distribution of the total number of NADPH-d type I neurons in the entire barrel field.

We performed counts of the number of these plotted cells in all reconstructed sections for the different developmental stages (P10, P23, P31 and P90) in order to quantitatively evaluate their distribution during cortical maturation. Comparisons between different groups were assessed by analysis of variance (ANOVA) with Newman–Keuls post hoc test at 95% (P < 0.05) confidence level. Comparisons between the numbers of NADPH-d neurons (septa versus barrels) in each group were assessed by Student’s t-test at 95% confidence level (P < 0.05).
Fig. 1. NADPH-d (right column) and CO (left column) reactivities during PMBSF development. Both enzymes are non-uniformly distributed in the PMBSF, being more reactive inside barrels, which are flanked by regions of weaker reactivity (septa). Note that the distribution of both enzymes is markedly similar and seems not to change during barrel field maturation. In all ages, sections reacted to CO were obtained from contralateral hemisphere of the same animal. Scale bar: 400 μm. Orientation: up, lateral; right, posterior.
3. Results

3.1. Postnatal development of the barrel field dimensions revealed by CO and NADPH-d histochemistries

The barrel field and all its compartments (especially the PMBSF) could be easily identified using both NADPH-d and CO histochemistries (Fig. 1) in all ages analyzed. Both enzymes are unevenly distributed in the PMBSF (Fig. 1), being more intense inside barrels, which are flanked by regions of weaker reactivity (septa).

The quantitative analysis performed on the Neurolucida reconstructions (Fig. 2) revealed a progressive increase in the total PMBSF area from P10 to P31 (Fig. 3A). From P31, the total PMBSF area remained stable and was not significantly smaller than that in P90 (Fig. 3A). The quantitative analysis of total individual barrel area and septal area revealed a trend similar to that previously reported for the total PMBSF area (Fig. 3B). Additionally, total individual barrel area did not differ \((P > 0.05)\) from septal area in any of the ages analyzed (Fig. 3B). Our results regarding the increase in the rat PMBSF area are in agreement with previous reports in the mouse barrel field (Rice and van der Loos, 1977).

3.2. Laminar distribution and morphology of NADPH-d neurons during postnatal barrel field development

The laminar distribution of both CO and NADPH-d did not change significantly during the postnatal development (data not illustrated). In a coronal view, both CO- and NADPH-d-reactive neuropil had virtually the same laminar pattern (Fig. 4): Layer I appears as a band of moderate reaction. In layers II–III, the neuropil displayed a higher reactivity and it was rather difficult to determine the limit between the two layers. In layer IV, both NADPH-d and CO revealed the presence of barrels and septa, as described elsewhere (Franca and Volchan, 1995; Vercelli et al., 1999). Layer V was characterized as a region of

Fig. 2. Neurolucida reconstructions of PMBSF during development. The number of NADPH-d cells in the septal region is higher than inside barrels for all ages evaluated (black dots). Arrows point to the four straddlers next to the five PMBSF barrel rows. Scale bars: 2 mm (lower magnification drawings); 300 μm (enlargements).
Fig. 4. Coronal view showing NADPH-d (A) and CO (B) distribution in adjacent sections of SI (P90). Note the similar distribution of both enzymes, which permit to define the six cortical layers. In A it is possible to see some type I NADPH-d neurons throughout layers (arrows). Arrowheads: white: example of barrel; black: septa. Asterisks: blood vessels. Scale bar: 200 μm.

low reactivity, while layer VI presented a darker staining, making it rather easy to identify the limit between these two layers and with the underneath white matter (Fig. 4). In the white matter, both enzymes were distributed more homogeneously with considerably less reaction product than that seen in any other layer of the cortical gray matter (Fig. 4).

The laminar distribution of NADPH-d neurons was similar to that previously described in the literature (Leigh et al., 1990; Bravo et al., 1997; Vercelli et al., 1999). Bipolar and multipolar NADPH-d aspiny neurons could be seen scattered throughout the cortical layers (data not illustrated). Nevertheless, these cells were rarely seen in layer I, but a higher number of these cells were present in layers II–III, where their dendritic trees were oriented in both horizontal and vertical planes. In layer IV, the majority of NADPH-d neurons were found in the less reactive septa.

The NADPH-d cells were especially numerous around the limit between layer VI and white matter. Interestingly, some white matter cells, especially those present at the border with layer VI, projected their dendrites into the gray matter and their cell bodies were elongated, with their principal axes oriented in parallel to the limit between gray and white matters (Fig. 5). Similar findings have been described in the primary visual cortex of other rodent species (Costa et al., 1994). In addition, it was possible to see many small neurons weakly reactive for NADPH-d in all ages evaluated (see examples in the Fig. 6), especially in the layers II and III. These neurons presented poorly labeled dendritic arbors. They are similar to type II NADPH-d neurons previously described in other species (Sandell, 1986; Franca et al., 1997; Franca et al., 2000; Wiencken and Casagrande, 2000).

There is a progressive modification in the complexity of the dendritic arborization of NADPH-d neurons during barrel field development (Fig. 7). In the earlier ages evaluated (P10–P14), only the cell bodies appeared highly reactive, without a prominent dendritic arbor (Fig. 7). In P23, the dendritic arbor of NADPH-d cells became more complex, displaying dendrites with both secondary and tertiary branches (Fig. 7). The adult pattern of dendritic ramification seemed to be fully established on P31 (Fig. 7). After this age it was not possible to see conspicuous modifications in cell morphology compared to that seen in mature animals (Fig. 7).

In the present study, type I NADPH-d neurons in septal regions seem to possess a more complex dendritic arborization than cells inside barrels, especially in more aged animals (Fig. 7).

In the sections submitted to double-labeling NADPH-d/NOS, the NADPH-d type I neurons could be labeled with the anti-neuronal NOS antibody (Fig. 8). This confirms that type I NADPH-d neurons correspond to a cell population responsible for the synthesis of NO in the brain (see also: Hope et al., 1991; Dawson et al., 1991; Valschanoff et al., 1995; Luth et al., 1995; Picanço-Diniz et al., 2004). For type II neurons, it was difficult to discern any pattern of double-labeling (Fig. 8).
3.3. Morphometric analysis of the NADPH-d positive neurons

NADPH-d neurons were present in both septa and barrels in all ages analyzed (see Fig. 7). Quantitative analysis of these neurons over the postnatal development (P10, P23, P31, P90) revealed a significant increase ($P < 0.05$) in the number of these cells from P10 to P23 (highest number of cells) in the PMBSF (Fig. 9). On the other hand, from P23 to P31 there was a significant decrease in the number of cells ($P < 0.05$). Nevertheless, from P31, the number of NADPH-d cells remained stable until P90 with no statistical differences between groups ($P > 0.05$). However, the number of NADPH-d neurons inside barrels was always lower than in septa (Fig. 9).
4. Discussion

The main goal of this study was to use qualitative and quantitative methods to investigate the distribution of NADPH-d neurons in rat SI and the maturation of their morphology, as seen by the distribution of the NADPH-d reaction product, during the barrel field development. In addition, we evaluated the postnatal development of NADPH-d activity in neuropil during development of this CNS region.

4.1. NADPH-d reactivity in the rat barrel field during development

In all ages evaluated in the present study (from P10 to P90) the barrel field could be easily visualized by using either NADPH-d or CO histochemistries. These results disagree with previous studies which report a virtual disappearance of the NADPH-d activity in the rat barrels from P21 to adulthood (Vercelli et al., 1999) and also in the mouse barrel field, which became undetectable by the end of the second postnatal week (Mitrovic and Schachner, 1996). Such conflicting results may be explained by a range of methodological differences, from fixation procedures to the specific NADPH-d histochemistry method used. For instance, according to our experience, postfixative procedures trick NADPH-d histochemistry (see Pereira et al., 2000). It is well known that NADPH-d activity in the brain can be altered by fixatives (Matsumoto et al., 1993). One can speculate that, during certain stages of postnatal development, NADPH-d activity is produced by enzymatic isoforms that are more sensitive to fixation. Nevertheless, the positive result obtained in this study reveals that NADPH-d activity in barrel cortex is unequivocally present during the whole postnatal development. In respect to the adult pattern, our
The pattern of NADPH-d reactivity in the barrel field was markedly similar in all ages evaluated in the present investigation. The enzymatic activity is segregated, though, so that the central regions of the barrels (hollows) appear as intensely stained patches, flanked by less reactive regions (septa). The barrel hollow is occupied by axon terminals arriving from thalamus (Kim and Ebner, 1999). The septa, on the other hand, are the target for callosal fibers coming from the opposite hemisphere (Koralek et al., 1990). It is possible that the NADPH-d/NO activity typical of barrel hollows can reflect the presence of NADPH-d-positive axon terminals in this region. This would be similar to thalamo-recipient layer of monkey primary visual cortex (layer IVC), where electron microscopy studies revealed that NADPH-d/NO reactivity is massively present in axonal terminals (Aoki et al., 1993).

4.2. Distribution of the NADPH-d neurons during barrel field maturation

The number of NADPH-d cells in PMBSF increased from P10 to P23, with a peak in the absolute number around the end of the third week/beginning of fourth week (P23). This result is similar to that described for other cortical regions, such as the prefrontal cortex (Tomic et al., 1994) and also for some subcortical regions, like hippocampus, lateral geniculate nucleus, and striatum (González-Hernandez et al., 1993; Moritz et al., 1999; Murata and Masuko, 2003). These findings suggest that, during the rat’s postnatal development the same quantitative changes in the number of NADPH-d neurons in PMBSF occur simultaneously in other brain regions. Nevertheless, for some brain areas, such as caudate-putamen, Yan et al. (1994) have reported that the highest number of NADPH-d cells is found in an earlier time point (around P14), but they did not evaluate the cortical area which includes the somatosensory cortex.

Whether or not the evolution of NADPH-d activity in individual neurons is similar in all cortical areas, and what are regulating it are still open questions. Based in previous reports, which describe a heterogeneous distribution of results are in agreement with previous reports of our group, which have shown the presence of a NADPH-d-positive barrel field in rats and mice (Franca and Volchan, 1995; Franca et al., 2000; Pereira et al., 2000).
NADPH-d in some parts of the rodent brain (Vincent and Kimura, 1992; Derer and Derer, 1993; Bidmon et al., 1997; Oermann et al., 1999; Franca et al., 2000), it is possible to hypothesize that there is an ontogenetic origin for the heterogeneous distribution of this cortical neuronal sub-population.

The pattern of distribution of NADPH-d neurons in the sub-compartments of the barrel field (septa and barrels) was markedly similar in all ages evaluated. The number of cells was always higher in septa than barrels. Such distribution is virtually similar to that described for other subgroups of interneurons, such as those immunoreactive to glutamic acid decarboxylase (GAD) and γ-aminobutyric acid (GABA) in the barrel field of small rodents (Lin et al., 1985; Chmielowska et al., 1986; Spreatco et al., 1988; Válchanovitch et al., 1993). Further studies are necessary to elucidate the physiological meaning of this differential distribution of the major neuronal groups in the barrel field.

In addition to their distribution, the morphology of NADPH-d neurons also appears to change noticeably during cortical maturation and seems to be well established around P23. The dendritic arbor of the neurons observed in P10–P14 is less complex than those seen in more aged animals (around P23 and until P90), for instance. These normal maturing events seem to be influenced by mechanisms dependent on NO (Harsanyi and Friedlander, 1997; Zhang et al., 2002; Fernández et al., 2003).

4.4. Conclusions

We present evidence that the enzymatic activity of NADPH-d remains high in the rat somatosensory system since P10 until adulthood, forming barrel fields that are well correlated with CO activity. In addition, our results show that the progressive changes in both morphology and number of NADPH-d neurons seem to occur at the same time window during barrel field maturation. Besides, distribution of NADPH-d cells in the barrel field seems to correlate with the distribution of these cells in some other regions of the developing rat brain. Further studies, using NOS-specific inhibitors or NOS knockout models to evaluate the expression of the NADPH-dNO during the period of barrel field maturation, should investigate whether or not NO plays a role on the mechanisms of barrel field formation.

Acknowledgements

This study was part of a dissertation submitted to UFPA Biological Science Post-Graduation Department as one of the requirements for the Master’s Degree of Marco Aurélio M. Freire. Study supported by grants from Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) and Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq). Thanks go to Mr. Dennyson de Faria for help in some experiments.

References


Tomic, D., Zobinjariu, M., Mezagoac, M., 1994. Postnatal development of nicotinamide adenine dinucleotide phosphate diaphorase...


