Histogenesis of nucleus proprius of lumbar spinal cord of fullterm human foetus

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ABSTRACT

Dorsal horn of grey matter of spinal cord is key centre for perception of not only nociceptive but also other sensory modalities. Earlier authors designated six laminae for the dorsal horn in different mammals including human. The literature available on the cytoarchitecture of nucleus proprius of human foetal spinal cord is meager. The present study is an attempt to observe histomorphometry of neurons of nucleus proprius of lumbar segment of spinal cord numerical density and population study of nucleus proprius. The spinal cord of a destitute full term foetus has been dissected and processed for light microscopic study. Holm’s silver nitrate staining was done for all sections. Nucleus proprius was found to be bulbous. There is a difference between right and left nucleus proprius in as far as morphometry, numerical density and population study of right and left nucleus proprius.

1. Introduction

Dorsal horn of the spinal cord is the key centre for the perception of not only nociceptive but also other sensory modalities. Substantia gelatinosa of Rolando was first identified to have been associated with nociception by Rolando. Later in 1888 Waldeyer identified a similar group deeper to substantia gelatinosa. Rexed [1] hypothesised a six horizontal laminar pattern arranged dorsoventrally and designated the laminae in roman numerals. Lamina II corresponds to substantia gelatinosa while the nucleus proprius corresponds to lamina III and IV [2]. The literature reveals the availability of tract cell neurons and interneurons in the nucleus proprius. A full term destitute foetus after perfusion with 10% formalin was subjected to laminectomy. Lumbar segment of the spinal cord has been dissected and processed for study under light microscopy. Histomorphometry of nucleus proprius in rat lumbar spinal cord by cresyl fast violet stained preparations were used for recording various measurements by Abercrombie’s method [3] by Bharadwaj et al [4]. Two zones have been identified in the nucleus proprius namely dorsal and ventral by the above authors. The authors observed dominance of spindle cells in dorsal as well as ventral divisions of nucleus proprius. Round cells were also encountered the fine architecture of dorsal horn and were studied by Ralston H [5] in macaque, in cat spinal cord by Ralston J [6], by Leonard, R.B and Cohan, D.H [7] in the spinal cord of pigeon. However the cytoarchitecture of human foetal dorsal horn of spinal cord was not traced in literature, an attempt is made to study the cytoarchitecture of human foetal spinal cord lumbar segment with reference to histomorphometry, numerical density & population study.

2. Materials and methods

One full term destitute male foetus available at the department of anatomy MIMS medical collage is the study material. The foetus has been perfused with 10% formalin has per the protocol. After 72 hours laminectomy was done and the spinal cord was exposed. Lumbar segment was identified and 1 mm thick lumbar spinal cord segment was processed for light microscopic study. Five microns thick sections were taken and stained with H and E, and Holue’s silver nitrate stain for identification of the tract cells of nucleus proprius. Morphometric studies as well as population studies have been done. The nucleus proprius constituted rounded and spindle shaped neurons. After going through all the sections, the length and breadth of nucleus proprius was calculated by using stage and eye piece micrometer. The volume was calculated by using the physics formula. The formula advocated by Saleem and Krishna Murthy [8] i.e. number of cells per cubic mm X total volume of the nucleus = total number of cells in the nucleus proprius was used.

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3. Results

The dorsal horn consisting of tract cells and interneurons arranged in nuclear zones. Substantia gelatinosa of Rolando present at the apex was recognized by densely packed small rounded neurons, ventral to this nucleus proprius was identified as a bulbous expansion having an average length of 662.5 microns on right side and 650 microns on left side, an average breadth of 868.75 microns on right side and 881.25 microns on the left side (fig 1). The nucleus has large and small rounded neurons, not having a restricted area for either of them as the large and small cells are present throughout the nucleus alternatively (fig 2). The length and breadth of the neuron is 6.12 microns on right side, whereas 5.575 microns on left side. The diameters of cells differ from right side to left side. The numerical density has been calculated by using the formula ND = NA / D + T, where ND is the number of cells per millimetre, NA is mean number of cells per reticule, D is the mean diameter and T is the thickness of the section [9, 10]. Average of both length and breadth gives mean diameter. Thickness of section was 5 microns. The numerical density of nucleus proprius on right side is 2920750 and on left side 26470638 (table 1). The population study of nucleus proprius revealed 759307 on right side, 694827 on left side (table 2).

Fig 1: Showing nucleus proprius Holme’s silver nitrate stain 10x10

Fig 2: Showing round cells of nuclei proprius Holme’s silver nitrate stain 10x40.

Table 1 showing morphometry of neurons of nucleus proprius

<table>
<thead>
<tr>
<th>Nucleus proprius</th>
<th>Average length</th>
<th>Average Breadth</th>
<th>Average Volume</th>
<th>Average No of cells per reticule</th>
<th>Numerical density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right</td>
<td>6.12 microns</td>
<td>6.12 microns</td>
<td>119.1 cubic microns</td>
<td>17.875</td>
<td>2920750</td>
</tr>
<tr>
<td>Left</td>
<td>5.575 microns</td>
<td>5.575 microns</td>
<td>90.10 cubic microns</td>
<td>14.625</td>
<td>2647063</td>
</tr>
</tbody>
</table>

Table 2 showing population study of nucleus proprius

<table>
<thead>
<tr>
<th>Nucleus proprius</th>
<th>Average length</th>
<th>Average Breadth</th>
<th>Volume</th>
<th>Population = Numerical Density x volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right</td>
<td>662.5 microns</td>
<td>868.75 microns</td>
<td>0.25997 millimeter cube</td>
<td>759307</td>
</tr>
<tr>
<td>Left</td>
<td>650 microns</td>
<td>881.25 microns</td>
<td>0.26249 millimeter cube</td>
<td>694827</td>
</tr>
</tbody>
</table>

4. Discussion

Histomorphometry of nucleus proprius has been studied by Bhardwaj et al [4], in rat lumbar dorsal horn of spinal cord. In rat the nucleus proprius could be divided into a dorsal part having smaller neurons and broader ventral part containing loosely dispersed neurons. Rexed [1] and Ralston [5] designated the apex of dorsal horn as substantia gelatinosa and designated it as lamina II. Schizogonthai [11] and Scheibel, M.E & Scheibel, A.B[12] could not differentiate lamina II & lamina III in rats and consolidated the 2 zones. Then neurons of dorsal division are small and densely packed. Spindle cells also dominated lamina II. Rounded cells constituted the second cell type having 23.18% of total cell population. In the ventral part of nucleus proprius, spindle cells dominated constituting 57.15% of total population, whereas rounded cells contributed 42.5%. Beal, JA & Cooper; M.H [13] and Ralston,H.J [5] revealed similar finding in the monkey spinal cord. In the present study nucleus proprius of full term of human foetal spinal cord, we could not observe a clear division of dorsal and ventral parts and nucleus is dominated by rounded cells of various dimensions as mentioned in table 1. There is a difference in cell dimension of nucleus proprius of right and left sides. Similarly nuclear density and population of cells differ from right to left.

On the right side the numerical density is 2920750, whereas on the left side the numerical density is 2647063. The population study revealed a difference between right & left side as shown in table 2. The present work revealed that the observations recorded as far as cell dimension, numerical density & population study are of significant value when compared with findings of other workers in mammals. Nucleus proprius is well defined and confined to III, IV, V laminae of grey matter of dorsal horn. No similar study has been made earlier as far as the search of the available literature reveals.
5. References


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