Study of the density of ganglion cells in the terminal bowel of rats with anorectal malformations

Estudo da densidade das células ganglionares no intestino terminal de ratos portadores de anomalia anorretal

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ABSTRACT

Purpose: To study the ganglion cells (GC) in the terminal bowel of rats with ethylenethiourea (ETU) induced anorectal malformations (ARM). Methods: The animals were divided into three groups: Group A – normal fetuses from pregnant rats that were not administered ETU; Group B – fetuses without ARM born from pregnant rats that were administered ETU and Group C - fetuses with ARM born from pregnant rats that received ETU. ETU was administered on the 11th day of pregnancy at the dose of 125 mg/kg body weight by gastric gavage. The rats had cesarean section on the 21st day of gestation. The fetuses' terminal bowel tissue was analyzed by immunohistochemistry to demonstrate ganglion cells. Results: Statistically significant differences were found between groups A, B and C regarding ganglion cell densities. Group A had the highest cell density, followed by Group B and the lowest density was found in Group C. Conclusion: Ganglion cell densities are decreased in the terminal bowel of rats with ARM.


RESUMO

Objetivo: Estudar as células ganglionares (CG) no intestino terminal de ratos portadores de anomalia anorretal (AAR) induzida pela etilenotiouréia (ETU). Métodos: Os animais foram distribuídos em três grupos: Grupo A - fetos normais, obtidos de ratas grávidas às quais não foi administrada ETU; Grupo B - fetos não portadores de AAR obtidos de ratas grávidas às quais foi administrada ETU e Grupo C – fetos portadores de AAR obtidos de ratas grávidas às quais foi administrada ETU. A ETU foi administrada no décimo primeiro dia de gestação, na dose de 125 mg/Kg, por gavagem. As ratas foram submetidas à laparotomia e histerotomia para retirada dos fetos no vigésimo primeiro dia de gestação. O intestino terminal dos fetos foi retirado e analisado por imunohistoquímica para pesquisa de CG. Resultados: Foram encontradas diferenças estatisticamente significantes entre os grupos A, B e C quanto à densidade de CG. O grupo A apresentou a maior densidade, seguida pelo grupo B, e a menor densidade foi encontrada no Grupo C. Conclusão: Existe uma menor densidade de CG no intestino terminal de ratos portadores de AAR.

Introduction

Intestinal constipation is one of the most frequent complaints in the postoperative period of correction of anorectal malformation (ARM), occurring in 30% to 60% of patients 1-2. It is a symptom that must be aggressively treated since it may trigger a picture of pseudoincontinence 3,4. The possible causes to explain intestinal constipation in the postoperative period of ARM include: anal stenosis 5, abnormalities of extrinsic intestinal innervation 6 and rectal denervation during the surgical procedure 7. Constipation may also be attributed to regional changes in intrinsic intestinal innervation (III), and there are reports abnormalities in ganglion cells (GC) of myenteric and submucous plexuses. Such changes include aganglionosis, hypoganglionosis and intestinal neuronal dysplasia 8-12. The study of the III can be performed as follows: by conventional histology using the classic hematoxylin-eosin staining 13; enzyme histochemical methods, such as activity of acetylcholinesterase, lactate dehydrogenase, succinate dehydrogenase and nitric oxide synthetase 13; and by immunohistochemical methods. Staining for GC or for neuronal supportive cells, also known as neuroglial cells, may be used in immunohistochemistry. For neuroglial cells, S-100 protein and glial fibrillary acidic protein (GFAP) 14-15 reactions may be used; and for GC staining, neuron-specific enolase and neurofilament protein reactions 14. The aim of this research is the comparative study of GC density in the terminal intestine of normal rats and rats with ARM induced by ethylenethiourea (ETU) 16, using S-100 protein reaction.

Methods

Six female WISTAR rats, weight 241-260 grams, were used. After an one-night mating, animals that presented a vaginal smear test positive for sperm in the next morning were considered fertilized. This was considered day zero (D0) of pregnancy. Animals were kept in individual cages and received commercial feed and water ad libitum, in an environment with constant temperature and humidity and automatically controlled 12-hour daytime and night cycles. On the 11th day (D11) of pregnancy, three pregnant rats received 125 mg/Kg ETU (2-imidazolidinethione C3H6N2S 98%, batch number 24251-089, Sigma-Aldrich, Brazil) by gavage and diluted with distilled water in a concentration of 1% (12.5 ml/Kg). The other three rats received only distilled water (12.5 ml/Kg) with no ethylenethiourea. The rats underwent cesarean section on the 21st (D21) day of pregnancy with removal of fetuses. Fetuses were examined for sex determination and assessment of the presence of ARM and other malformations. The model used caused anorectal anomaly in 72% of fetuses of rats treated with ETU. No fetus from rats untreated with ETU presented malformations. The animals were then divided into three groups:

- **Group external control (A)** (N=11) – Normal fetuses, obtained from rats untreated with ETU;
- **Group internal control (B)** (N=8) – Fetuses without ARM obtained from rats treated with ETU;
- **Group anorectal malformation (C)** (N=21) – Fetuses with ARM, obtained from rats treated with ETU.

Afterwards, by means of optical microscopy, fetuses underwent exploratory laparotomy to determine the type of ARM, when present, and to excise one centimeter of terminal intestine for immunohistochemical study. In normal fetus, dissection of terminal intestine was performed up to the anal level, and in fetuses with ARM it was carried out up to the fistula site. The one-centimeter fragment of terminal intestine was fixed in 10% buffered formaldehyde, dehydrated and embedded in paraffin. The types of anorectal malformation found were: 13 cases of cloacal malformation, 4 ARM with perineal fistula and 4 ARM with urethral fistula.

**Immunohistochemistry**

From the previously parafin-embedded material, three-micron sections were performed in the intestinal longitudinal axis and the sections were processed with the streptavidin-biotin-peroxidase technique. The sequence of immunohistochemical procedures included deparaffinization in stowe, deparaffinization in xylol, rehydration in absolute alcohol, antigen recovery with 0.01M citrate buffer at pH 6.0, block of endogenous peroxidase activity with a solution of hydrogen peroxide (H2O2), block of inspecific reaction through incubation with a phosphate buffer at pH 7.2, with addition of 1% bovine serum albumin for 15 minutes, incubation in the primary antibody against S-100 protein at the dilution of 1:200 (Rabbit Polyclonal Antibody NCL-S100p-Novocastra) overnight, at approximately 8°C (46.4°F), incubation in a biotinylated secondary antibody at room temperature for 30 minutes, incubation in the streptavidin-biotin-peroxidase complex at room temperature for 30 minutes, incubation with enzyme substrate diaminobenzidine and counterstaining with Harris hematoxylin.

**Ganglion cells count**

Each fetus was represented by a slide of terminal intestine magnified 400 times, and ten digital images were...
obtained throughout the intestine with the use of a Sony® CCD-IRIS video camera attached to an Olympus (BX 50) light microscope. The images obtained were studied with a program of image analysis called “Image Tool” ® - version 3.0. For each image obtained, we calculated the area corresponding to the circular and longitudinal musculature and counted the number of GC in the isolated area. Cellular density was obtained by the total number of cells divided by the total area, thus yielding the number of cells per mm².

Analyses performed

We compared the densities of GC in the terminal intestine of fetuses from the three groups by using analysis of variance (ANOVA) and the Bonferroni method. The significance level used was 0.05 (α = 5%) and descriptive levels (p) lower than this value were considered significant.

Ethics

This experiment received approval from the Research Ethics Committee of the Federal University of São Paulo (UNIFESP-EPM - protocol n. 0406/05).

Results

The density of GC in animals of the three groups is shown in Table 1. Statistically significant differences were found in the three groups regarding GC density in the terminal intestine, when submitted to variance analysis (Table 2). Comparing the groups in pairs by the Bonferroni method, the results showed statistically significant differences among the three groups (Table 3). The external control group presented a higher density than the internal control group, which, in turn, presented higher density than ARM group (Figures 1, 2 and 3).

TABLE 1 – Density of ganglion cells in animals from Groups A, B and C (cells per mm²)

<table>
<thead>
<tr>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
</tr>
</thead>
<tbody>
<tr>
<td>28,71</td>
<td>21,64</td>
<td>18,12</td>
</tr>
<tr>
<td>27,10</td>
<td>23,02</td>
<td>9,95</td>
</tr>
<tr>
<td>26,84</td>
<td>17,49</td>
<td>17,05</td>
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<tr>
<td>23,17</td>
<td>18,93</td>
<td>17,16</td>
</tr>
<tr>
<td>26,08</td>
<td>18,43</td>
<td>13,34</td>
</tr>
<tr>
<td>24,63</td>
<td>17,69</td>
<td>14,24</td>
</tr>
<tr>
<td>24,07</td>
<td>17,16</td>
<td>11,34</td>
</tr>
<tr>
<td>21,83</td>
<td>20,56</td>
<td>12,29</td>
</tr>
<tr>
<td>33,70</td>
<td>10,10</td>
<td></td>
</tr>
<tr>
<td>29,00</td>
<td>8,77</td>
<td></td>
</tr>
<tr>
<td>26,47</td>
<td>9,78</td>
<td></td>
</tr>
</tbody>
</table>

FIGURE 1 – Photomicrography of the terminal intestine of the external control group with S-100 protein staining

FIGURE 2 – Photomicrography of the terminal intestine of the internal control group with S-100 protein staining
The occurrence of III abnormalities in ARM is a controversial issue among some authors, who have been always concerned about investigating the presence or absence of GC in biopsies or in surgical resections. In reports mentioned above 4-6, no abnormalities were found. Other authors, however, described different results. Parkkulainen et al 9 evaluated the presence of ganglia in 25 ARM individuals, and found aganglionosis in 15 (60%). Kiesewetter et al 10 conducted a study involving 32 pediatric surgery centers in the United States and Canada, in which 296 patients were assessed and 10 (3.4%) presented aganglionosis. Holschneider et al11 studied 52 patients and observed an abnormal innervation pattern in 49 (94%). Meier-Ruge et al12,17 evaluated 41 ARM patients and observed 60% of enteric nervous system anomalies. Anyway, considering that the incidence of ARM, as well as that of aganglionosis, is one in 5000 newborns, it would be expected to find an even lower frequency for the association of both lesions than for the conditions isolated. The current study evaluated the intrinsic intestinal innervation through the study of GC. This study involved the calculation of cell density per square millimeter in the area represented by the circular and longitudinal muscular layers where the myenteric plexus is located and already present in the newborn rat. The S-100-protein immunohistochemical method was chosen because the enteric glial cells and the Schwann cells in the intestine presented strong immunoreactivity for this protein with a significant visualization of the GC supportive elements. Harris hematoxylin counterstaining was associated for GC staining. Experimental studies correlating ARM with intrinsic intestinal innervation are scarce and applied different methods. Favre et al 18 used neurofilament protein and nitric oxide synthetase stainings in Danforth’s short tail mice and demonstrated an association with hypoganglionosis (41%) and aganglionosis (58%). Mandhan et al 19 used neuron-specific enolase and P-substance to assess the terminal intestine of newborn rats with ARM induced by ETU, and reported decreased immunoreactivity in both methods. The internal control group, despite not having ARM in its phenotype, presented a decreased count of GC. This fact might be related to the mechanism of action of ETU. It is believed that ETU possibly has its effects by changing the signaling of the Sonic hedgehog (SHH) gene. The molecule codified by the SHH gene is

### Table 2 - Description of the variable “Density of ganglion cells” in groups A, B and C

<table>
<thead>
<tr>
<th>Group</th>
<th>N</th>
<th>Mean ± S.D.</th>
<th>Median</th>
<th>Minimum-Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A</td>
<td>11</td>
<td>26.51 ± 3.25</td>
<td>26.47</td>
<td>21.83 – 33.70</td>
</tr>
<tr>
<td>Group B</td>
<td>8</td>
<td>19.36 ± 2.15</td>
<td>18.68</td>
<td>17.16 – 23.02</td>
</tr>
<tr>
<td>Group C</td>
<td>21</td>
<td>12.52 ± 3.39</td>
<td>11.75</td>
<td>6.55 – 19.66</td>
</tr>
</tbody>
</table>

p < 0.001  
F Test (ANOVA)

Key:  
N – Number  
SD – Standard deviation  
mm² – Square millimeter

### Table 3 - Multiple comparisons among Groups A, B and C, for the variable “Density of ganglion cells”

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean difference</th>
<th>p</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A – Group C</td>
<td>13.91</td>
<td>&lt;0.001</td>
<td>Group A &gt; Group C</td>
</tr>
<tr>
<td>Group A – Group B</td>
<td>7.12</td>
<td>&lt;0.001</td>
<td>Group A &gt; Group B</td>
</tr>
<tr>
<td>Group B – Group C</td>
<td>6.82</td>
<td>&lt;0.001</td>
<td>Group B &gt; Group C</td>
</tr>
</tbody>
</table>

Bonferroni Method  
p < 0.05

![Photomicrography of the terminal intestine of the ARM group with S-100 protein staining](image)

**FIGURE 3** – Photomicrography of the terminal intestine of the ARM group with S-100 protein staining

### Discussion

The occurrence of III abnormalities in ARM is a controversial issue among some authors, who have been always concerned about investigating the presence or absence of GC in biopsies or in surgical resections. In reports mentioned above 4-6, no abnormalities were found. Other authors, however, described different results. Parkkulainen et al 9 evaluated the presence of ganglia in 25 ARM individuals, and found aganglionosis in 15 (60%). Kiesewetter et al 10 conducted a study involving 32 pediatric surgery centers in the United States and Canada, in which 296 patients were assessed and 10 (3.4%) presented aganglionosis. Holschneider et al11 studied 52 patients and observed an abnormal innervation pattern in 49 (94%). Meier-Ruge et al12,17 evaluated 41 ARM patients and observed 60% of enteric nervous system anomalies. Anyway, considering that the incidence of ARM, as well as that of aganglionosis, is one in 5000 newborns, it would be expected to find an even lower frequency for the association of both lesions than for the conditions isolated. The current study evaluated the intrinsic intestinal innervation through the study of GC. This study involved the calculation of cell density per square millimeter in the area represented by the circular and longitudinal muscular layers where the myenteric plexus is located and already present in the newborn rat. The S-100-protein immunohistochemical method was chosen because the enteric glial cells and the Schwann cells in the intestine presented strong immunoreactivity for this protein with a significant visualization of the GC supportive elements. Harris hematoxylin counterstaining was associated for GC staining. Experimental studies correlating ARM with intrinsic intestinal innervation are scarce and applied different methods. Favre et al 18 used neurofilament protein and nitric oxide synthetase stainings in Danforth’s short tail mice and demonstrated an association with hypoganglionosis (41%) and aganglionosis (58%). Mandhan et al 19 used neuron-specific enolase and P-substance to assess the terminal intestine of newborn rats with ARM induced by ETU, and reported decreased immunoreactivity in both methods. The internal control group, despite not having ARM in its phenotype, presented a decreased count of GC. This fact might be related to the mechanism of action of ETU. It is believed that ETU possibly has its effects by changing the signaling of the Sonic hedgehog (SHH) gene. The molecule codified by the SHH gene is
involved in the development of several systems, including the nervous and skeletal systems, craniofacial structures and the gastrointestinal tract. In the gastrointestinal tract, the molecule is responsible for the regionalization of the primitive gut tube, the radial pattern of the digestive tube, the differentiation of its layers and the differentiation of enteric neurons. The abnormalities found in the internal control group might have been triggered by changes in the signaling of SHH gene, which caused some of the abnormalities found, but it did not have its full effect for not inducing ARM.

**Conclusion**

There is a lower ganglion cells density in the terminal intestine of rats with ARM induced by ethylenethiourea.

**References**