The Effect of Mesenchymal Stem Cells on Fertility in Experimental Retrocervical Endometriosis

O efeito das células-tronco mesenquimais na fertilidade em endometriose retrocervical experimental

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Abstract

Purpose To evaluate the effect of mesenchymal stem cells (MSCs) on fertility in experimental retrocervical endometriosis.

Methods A total of 27 New Zealand rabbits were divided into three groups: endometriosis, in which endometrial implants were created; mesenchymal, in which MSCs were applied in addition to the creation of endometrial implants; and control, the group without endometriosis. Fisher’s exact test was performed to compare the dichotomous qualitative variables among the groups. The quantitative variables were compared by the nonparametric Mann-Whitney and Kruskal-Wallis tests. The Mann-Whitney test was used for post-hoc multiple comparison with Bonferroni correction.

Results Regarding the beginning of the fertile period, the three groups had medians of 14 ± 7.6, 40 ± 5, and 33 ± 8.9 days respectively (p = 0.005). With regard to fertility (number of pregnancies), the endometriosis and control groups showed a rate of 77.78%, whereas the mesenchymal group showed a rate of 11.20% (p = 0.015). No differences in Keenan’s histological classification were observed among the groups (p = 0.730). With regard to the macroscopic appearance of the lesions, the mesenchymal group showed the most pelvic adhesions.

Conclusion The use of MSCs in endometriosis negatively contributed to fertility, suggesting the role of these cells in the development of this disease.

Keywords
► endometriosis
► mesenchymal stem cell
► fertility
► rabbits

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Resumo

Objetivo Avaliar o efeito das células-tronco mesenquimais sobre a fertilidade na endometriose retrocervical experimental.

Métodos Um total de 27 coelhas da raça Nova Zelândia foram divididas em três grupos: endometriose, em que os implantes endometriais foram criados; mesenquimal, em que as células-tronco mesenquimais foram aplicadas complementarmente à criação implantes endometriais; e controle, sem endometriose. O teste exato de Fisher foi realizado para comparar variáveis dichotômicas qualitativas entre os grupos. As variáveis quantitativas foram comparadas pelos testes não paramétricos de Mann-Whitney e Kruskal-Wallis. O teste de Mann-Whitney foi utilizado para a comparação múltipla pós-hoc com correção de Bonferroni.

Resultados em relação ao início do período fértil, os grupos endometriose, mesenquimal e controle tiveram medianas de 14 ± 12,7; 40 ± 5; e 33 ± 8,9 dias, respectivamente (p = 0,005). Sobre a taxa de fertilidade (número de gravidezes), os grupos endometriose e controle mostraram uma taxa de 77,78%, enquanto o grupo mesenquimal mostrou uma taxa de 11,20% (p = 0,015). Não foram observadas diferenças na classificação histológica de Keenan entre os grupos (p = 0,730). No que diz respeito à aparência macroscópica das lesões, o grupo mesenquimal mostrou maiores adesões pélvicas.

Conclusão O uso de células-tronco mesenquimais na endometriose contribuiu negativamente para a fertilidade, sugerindo o papel dessas células no desenvolvimento da doença.

Palavras-chave
► endometriose
► célula-tronco mesenquimal
► fertilidade
► coelhos

Introduction

Endometriosis is a disease characterized by the presence of glands and/or endometrial stroma outside the uterine cavity. Endometriosis can be macroscopically divided into three clinically distinct entities: superficial peritoneal endometriosis, ovarian endometriosis (endometrioma), and deep endometriosis. Deep endometriosis is estimated to occur in ~ 20% of women with the disease. The most common symptoms of endometriosis are dysmenorrhea, dyspareunia, chronic pelvic pain, and infertility, and the deep form of the disease is often related to infertility. One study reported the detection of endometriosis in 20–50% of women who underwent investigations for infertility.

Currently, groups of scientists are researching the presence of progenitor stem cells in the endometrium and the correlation of this phenomenon with endometrial regeneration and menstrual cyclicity. In 2004, a research group led by Gargett demonstrated the presence of a small group of cells exhibiting clonogenic capacity in the human endometrium. In the same year, another group led by Taylor proposed that bone marrow cells in humans could participate in the endometrial regeneration process in recipient patients.

Vidane et al. suggested that mesenchymal cells derived from the bone marrow also contribute to endometrial growth and play a role in the progression of endometriosis. Endometriosis is a disease with inflammatory characteristics. Furthermore, mesenchymal stem cells (MSCs) release factors with the following trophic properties: immunomodulatory, angiogenic, and antiapoptotic. These cells appear to need an inflammatory environment for exerting their immunomodulatory function, and such an environment exists in endometriosis.

Mesenchymal stem cells are considered non-immunogenic, but they can differentiate into various tissues and secrete immunomodulatory and regenerative/reparative factors. Therefore, MSCs are considered optimal for cell therapy. These cells are primarily used for regeneration; their effects on the immune system and the secretion of growth factors have opened fields in the clinical research.

The complexity of the pathophysiology of endometriosis, the heterogeneity of the disease in humans, and the effect of mesenchymal cells as immunomodulators are still controversial. This context, female rabbits are a good choice as experimental models. They have homogeneous lesions—that is, generally solid and hemorrhagic masses—that can be easily created by autograft of endometrial fragments or by the opening and exposure of the endometrial cavity.

The objective of this study was to evaluate the possible contribution of MSCs for the development of endometriosis and its effects on fertility through induced lesions in rabbits.

Methods

This experimental study was conducted after approval by the Animal Ethics Committee of the university (CEUA-PUCPR, project no. 710 C). A total of 29 adult female New Zealand rabbits (6–8 months of age) weighing between 3 and 4 kg were used. This total included 27 rabbits used for the sample and 2 for bone marrow collection.
The surgical procedures and the extraction and cultivation of MSCs were performed at the Operative Technique and Experimental Surgery Laboratory and the Cell Therapy Laboratory, both located at the same university in southern Brazil. All handling of the animals followed the rules of the Council for International Organization of Medical Sciences (CIOMS), the Ethical Code for Animal Experimentation and the precepts of the Colégio Brasileiro de Experimentação Animal (COBEA, www.cobea.org.br).

The rabbits were divided into three groups of nine each: control group (C), endometriosis group (E), and mesenchymal group (M). In the endometriosis group, a technique developed by do Amaral et al was performed: a median incision was initially made in the abdominal wall. Then, a 4-cm piece was removed from the right uterine horn. Patches 5 × 5 mm in size were created and sutured in the retrocervical region.

In the control group, a fragment of the uterine horn was removed following the procedure described before. However, the endometrial implant was not sutured in these animals. The mesenchymal group was submitted to the same procedure as the endometriosis group, and later, also submitted to the application of mesenchymal cells.

In all three groups there was a delay of 21 days before copulation was initiated. This is the minimum time required for disease development to occur or for the implanted endometrium to become secretory. In the Endometriosis and Control groups the copulations occurred after the first surgery. Already in the Mesenchymal group, the copulation only started after 21 days of the second surgical procedure (application of the mesenchymal cells).

Copulation occurred 4, 6, and 9 weeks after surgery, when the rabbits were fertile. To check fertility, we evaluated the coloring of the vulva daily, and females with red/violet vulva (unlike the normal whitish color) were placed in a cage with a male for mating. Abdominopelvic ultrasounds were performed, and the animals were then checked for signs of pregnancy (the presence of a gestational sac or fetus). Fig. 1 shows a flowchart with a brief description of the chronology of the experiments in the study.

Mesenchymal stem cells were obtained 12 weeks before the application by extracting bone marrow from the long bones of two rabbits. The femur and tibia were removed, and all tissue was scraped away from the bones. The bones were placed in a Petri dish containing phosphate buffered saline (PBS; Gibco Invitrogen, NY, USA). The epiphyses were cut to expose the medullary cavity. The marrow was removed from each bone with flushing medium into a tube (TPP, Trasadingen, Switzerland) using a syringe with a needle.

The collected bone marrow was processed using a Ficoll-Hypaque density gradient (d = 1.077 g/mL; Sigma Chemical, St Louis, USA). The isolated mononuclear cells were plated in 75-cm² flasks (TPP), and were cultivated using IMDM (Iscove’s Modified Dulbecco’s Media; Gibco Invitrogen) medium containing 15% fetal calf serum (FCS; Gibco Invitrogen) and 1% antibiotics. The cultures were maintained in an incubator, and the culture medium was changed every 72 hours. Upon reaching 80% confluence, the cells were trypsinized using a trypsin-EDTA solution (Gibco Invitrogen), and were subcultured to obtain the required number of cells for potency tests and transplantation.

![Fig. 1 Chronology of the experiments in the study.](image-url)
The MSCs were assessed for potency by inducing their differentiation into adipocytes, osteoblasts, and chondrocytes (Fig. 2 A-F). For adipogenic and osteogenic differentiation, cells were seeded on glass coverslips (Sarstedt, Newton, USA) in 24-well plates. A commercial differentiation medium (Gibco Invitrogen) was used for 21 days. The MSCs were stained with Oil Red O to analyze adipogenic induction. Large, rounded cells with cytoplasmic lipid-rich vacuoles were observed. Osteogenic differentiation was assessed by visualizing the mineralization of the extracellular matrix with Alizarin Red S staining. Cells were grown in a micromass culture to promote chondrogenic differentiation. In the chondrogenic differentiation assays, high-density micromass MSC cultures generated cellular nodules, which produced large amounts of cartilage-related extracellular matrix molecules, such as collagen. Paraffin sections of the aggregates stained with toluidine blue showed condensed structures with cuboidal cells and chondrocyte-like lacunae.

Untreated control cultures, which were grown in regular medium without adipogenic, osteogenic, or chondrogenic differentiation stimuli, did not exhibit spontaneous adipocyte, osteoblast, or chondrocyte formation after 21 days of MSC cultivation.

After 21 days, in the Mesenchymal group, the second surgery for the application of the mesenchymal cells was also performed. During this reoperation, the size of the lesion formed by the implant was checked. All the implants were considered viable (with an area not exceeding 25 cm²). We used 2 × 10⁶ cells in each animal (2 ml in a syringe), which were applied on the edge of the implant.

For the histological analysis, endometrial implants fixed in formalin were prepared in paraffin blocks. These samples were later sectioned at a thickness of 5 μm (4 cuts per sample), stained with hematoxylin and eosin, and evaluated with an optical microscope. The lesions were classified according to Keenan’s index. Scores were assigned as follows: score of 3: a well-preserved epithelial layer; score of 2: moderately preserved epithelium with leukocyte infiltration; score of 1: poorly preserved epithelium (occasional epithelial cells only); and score of 0: no epithelium (Fig. 3 A-B).

Fig. 2 (A-F) Differentiation of mesenchymal cells. (B) The differentiation into the adipocyte lineage was demonstrated by staining with Oil Red O; (D) Alizarin Red S staining shows mineralization of the extracellular matrix in the osteogenic differentiation; and (F) toluidine blue shows the deposition of proteoglycans and lacunae in the chondrogenic differentiation. (A, C, E) Untreated control cultures without adipogenic, osteogenic or chondrogenic differentiation stimuli are shown.
The sample size was based on similarity with other articles; to control Type II Error in the applied tests, test power was used.

Qualitative variables were described as frequencies and percentages, and quantitative variables were described as means, medians, and quartiles. Fisher’s exact test was performed to compare dichotomous qualitative variables among the groups. The quantitative variables were compared by the nonparametric Mann-Whitney and Kruskal-Wallis tests. The Mann-Whitney test was used for post-hoc multiple comparison with Bonferroni correction. Values of $p < 0.05$ were considered statistically significant. The data were analyzed with the Statistical Package for the Social Sciences (SPSS; IBM SPSS, Chicago, IL, US) software, version 20.0.

**Results**

The mesenchymal group had the worst fertility rate (11.1%; $p = 0.015$; test power: 84.29%); only one animal in this group became pregnant. In contrast, seven rabbits in the control group and seven in the endometriosis group (77.8%) became pregnant.

After having executed the surgical procedures and waited 21 days, the time to start fertility was different between the groups. The mesenchymal group had the longest delay when compared with the other groups ($p = 0.005$; test power: 92.24%) (Table 1). Considering a significant difference found among the groups, they were compared two by two.

The $p$ values and the test power of these comparisons are shown in the Table 2.

The endometriosis group had 2 fetal deaths (28.57%) and 5 live births (71.43%). The control group had 7 pregnancies, which resulted in 2 fetal deaths (28.57%), 3 abortions (42.86%), and 2 live births (28.57%).

There was no difference in implant size between the endometriosis (median = 55, 1st quartile = 15, and 3rd quartile = 90) and mesenchymal groups (median = 55, 1st quartile = 25, and 3rd quartile = 80; $p = 0.863$).

Macroscopically, the formation of a large adhesion in the cavity was noted in the mesenchymal group, even making it difficult to identify the implants (Fig. 4A). Comparatively, a minor amount of adhesion was noticed in the endometriosis group (Fig. 4B). Finally, discrete adhesion formation was observed in the control group. Despite the formation of adhesions, endometriotic lesions were only found in the sites implanted, and no dissemination was observed.

The persistence of epithelial cells in the endometrial implants was assessed semi-quantitatively with Keenan’s index (Fig. 3 A, B). The following results were observed among the nine analyzed rabbits in each group. The endometriosis group included 3 rabbits with a Keenan index of 0, and 6 with a Keenan index of 3, and the mesenchymal group included 2 rabbits with a Keenan index of 0 and 7 with a Keenan index of 3 ($p = 0.730$).

<table>
<thead>
<tr>
<th>Groups</th>
<th>n</th>
<th>Span in days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endometriosis</td>
<td>9</td>
<td>33 [18–34]</td>
</tr>
<tr>
<td>Mesenchymal</td>
<td>9</td>
<td>14 [14–16]</td>
</tr>
<tr>
<td>Control</td>
<td>9</td>
<td>40 [33–40]</td>
</tr>
</tbody>
</table>

Note: Nonparametric test of Kruskal-Wallis; $p < 0.05$; median [1st - 3rd quartiles]; Test power = 92.24.

**Table 2** Test power and $p$ values in the inter-group comparison of time until the onset of the fertile period

<table>
<thead>
<tr>
<th>Groups</th>
<th>$p$</th>
<th>Test power</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control x Endometriosis</td>
<td>0.061</td>
<td>0.3314747</td>
</tr>
<tr>
<td>Control x Mesenchymal</td>
<td>0.043</td>
<td>0.7495601</td>
</tr>
<tr>
<td>Endometriosis x Mesenchymal</td>
<td>&lt; 0.001</td>
<td>0.9583571</td>
</tr>
</tbody>
</table>

Note: Nonparametric test of Kruskal-Wallis; $p < 0.05$.

![Fig. 3](image-url) (A) Histological image of the endometrial implant with a Keenan index of 3. (B) Histological picture of the endometrial implant with a Keenan index of 0.
Discussion

The selection of rabbits as an experimental model was based on their efficacy for developing ectopic endometrial foci in the abdominal wall. It is known that the production of various cytokines occurs in endometriosis, which generally stimulates endometrial proliferation and contributes to the production of acute-phase inflammatory proteins, creating a pro-inflammatory environment. It is also known that in endometriosis, the endometrial cells exhibit the characteristics of chronicity and expansion, showing capacities for migration, invasion, and apoptosis resistance.

This scenario has led to new theories exploring the involvement of the immune system, such as immunological mechanisms facilitating the development of endometriosis, or the immune system being activated by it. The involvement of stem cells (particularly of the mesenchymal lineage) in the origin of the disease has also been proposed. In contrast with the reports on endometriosis leading to infertility, the development of endometriosis in the rabbits did not worsen the fertility rate. However, the rabbits were not a good model for endometriosis-related infertility. The use of the obtained MSCs caused the relative risk of infertility to increase 4-fold in these animals.

One mechanism that may explain the aforementioned result involves the pathophysiology of the disease. The evidence of stem cells in the human endometrium and the suggestion of the presence of stem cells—specifically the mesenchymal lineage—may be related to the worsening of the fertility rate in the group that received mesenchymal cells, because we used cells that may be involved in the origin and progression of the disease.

Another mechanism that may be involved is the epithelial–mesenchymal transition (a process comprising a series of changes in epithelial cell phenotype that causes epithelial cells to assume the characteristics of MSCs, such as having a greater capacity for migration, invasion, and apoptosis resistance). Because endometriotic cells in the context of endometriosis also take on this function, they represent a sum of the effects of both mechanisms.

A question remains regarding whether the mesenchymal group’s delay in entering the fertile period was related to the involvement of MSCs in the origin and progression of endometriosis.

With regard to obstetric outcomes, the endometriosis and control groups showed no statistically significant difference. This result may be because of the small sample size. The mesenchymal group requires further analysis because of its low fertility rate.

The macroscopic evaluation of the pelvic cavity of each animal (during euthanasia) revealed that the mesenchymal group had a greater rate of adhesions than the other groups, in an analogy with the widely used Revised Classification of the American Society for Reproductive Medicine (ASRM). Thus, we could consider the mesenchymal group a carrier of stage-IV disease (severe endometriosis with dense and firm adhesions).

It is known that anatomical distortion and adhesions are some of the mechanisms that generate infertility in the context of endometriosis. This mechanism may explain why the group that received MSCs on the endometrial implants may have had a low fertility rate: its higher rate of pelvic adhesions.

Finally, a histological analysis of the implants using Keenan’s index was performed. This index measures the persistence of epithelial cells in endometrial implants. A comparison of the two groups revealed no statistically significant difference in the scores of Keenan’s index.

Although it may be possible that the worsening of fertility in the mesenchymal group can be explained by two other mechanisms (the longer time the animals were exposed to the implants and the occurrence of two surgical procedures in this group), in this study design, according to our results, we can conclude that applying MSCs on female rabbits via endometrial implants provides a substrate for the development of endometriosis and thus harms fertility in these animals.
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