MODELING OF THE IMPLANT FOR VENTRAL HERNIAS


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ABSTRACT — The issue associated with hernioplasty of ventral hernias does not cease to be relevant. And even the opposite with the development of technologies and new materials for implantation, there are more and more questions, which one to choose. Our study included the ProgripTM self-gripping implant, which showed good biocompatibility properties and was also convenient to use. However, the long-term results of treatment were not satisfactory due to relapse. In this regard, a simulated implant was developed and implemented in practice.

KEYWORDS — postoperative ventral hernia, material for implants.

THE RELEVANCE OF RESEARCH

One of the first, and most common, surgical pathologies where alloplastic materials for prosthetics began to be used are hernias. Surgery for hernia of the anterior abdominal wall was initially crippling and with high mortality, therefore, in many countries of the ancient world, hernia repair was prohibited. A new stage using prosthetics began in the early twentieth century with the help of implants made of silver, gold, various metals.

But in the postoperative period, the use of implants from these materials caused severe complications. After World War 2, research began on the possibility of using polymeric synthetic materials in medicine. One of the first in 1959 F.C.Usher proposed the use of polypropylene for medical purposes [1, 2, 3]. The use of mesh implants, although reduced the risk of recurrence of the hernia, however, up to 30% of patients again seek surgical help. Therefore, the question of choosing a mesh implant and the method of its application remains open [4].

In the process of research it was proved that the implant of the cellular structure, better than monolithic or perforated, which was used initially. Through the cellular structure, connective tissue grows faster, forming the skeleton of the anterior abdominal wall, from the formation of which depends on the development of relapse in the future. The duration and quality of the ingrowth of the prosthesis into the surrounding tissues are influenced by both the physical properties of the prosthesis, such as the size and shape of the pores, the specific weight, density and structure of the material used, and the characteristics of the human body [5].

The formation of a connective tissue scar depends on the collagen metabolism at different stages of the wound process, with a change in the ratio of the amount of collagen I and type III [6, 7].

Currently, hernioplasty uses implants of three main materials:

1. Polytetrafluoroethylene (PTFE) — microporous material, the pore size is 10 microns, which ensures flexibility and elasticity. But at the same time, microporosity impairs the ingrowth of the mesh into the surrounding tissues and encapsulation occurs. As a result of the encapsulation, the elastic properties are lost, as well as the displacement of the mesh prosthesis. Therefore, these types of implants are almost not used [8, 10].

2. Polypropylene (PP) — light macroporous consisting of monofilament. This type of implant is well integrated into the surrounding soft tissue, thanks to the size of pores, the penetration of macrophages, vessels and fibroblasts is not difficult. However, a thin connective tissue capsule is again formed around the prosthesis, forming a rough scar with imperfect implant growth. And the processes of degradation of the material cause a loss of physical properties.

The reduction of the prosthesis occurs up to 50% of the original size with a displacement of the implant, which can lead to a recurrence of the hernia. A number of studies have established that the severity of the inflammatory reaction and the activity of fibrosis in the implantation zone directly depend on the amount of polypropylene [11, 12, 13].
The implant of the standard form — oval, removes the transverse load from the aponeurosis of the rectus abdominis muscles due to fixation of the rectus muscles to the back wall of the vagina and compensating for the stretching force, making the muscle-aponeurotic tissues not expandable.

Materials and methods.

The study included 66 patients operated on for postoperative ventral hernia. The first group of patients consisted of persons up to 60 years old, the second group includes patients older than 60 years old (Table 1). In each group, patients were operated on using a standard-shaped ProgripTM self-gripping implant (subgroup B) and a modeled one (subgroup A).

Self-gripping implant was used since the fixation of the implant occurs evenly in contrast to the implants which need to be fixed to the tissues with additional material around the perimeter. When changing the shape of the implant, the physical tension of the muscles of the anterior abdominal wall was taken into account.

The volume of the examinations carried out in the preoperative period met all the requirements of preparation for chronic diseases, if necessary additional examinations were prescribed in the form of ultrasound examination (USI) of the abdominal organs and kidneys, vessels (veins and arteries) and heart; spirometry, radiography, computer tomography. The risk assessment of the development of venous thromboembolic complications (VTEC) was also carried out on the Caprini scale.

In the postoperative period all patients received adequate analgesia on the first day after surgery; And the fixation points A1, A2, B1 and B2 are at the same time the points of application of tensile transverse forces and forces directed from the inside of the abdominal cavity, which leads to tearing of the tissue in a given area or tearing the implant. In the modified implant, there is a cut on the opposite sides of the implant along the longitudinal axis, due to which only the longitudinal load remains, and the lack of transverse stretching force reduces the load at points D1 and D2. This prevents tissue rupture and implant detachment, and consequently reduces the risk of recurrence (Fig. 1.).

According to the Chevrel J.P. and Rath A.M. (1999), the study included patients SmW1–4R0–2 where “Sm” means the median localization. Patients with mixed and lateral localization of the defect were excluded since it is impossible to use the simulated implant under this localization. The width of the hernial defect — “W” and the number of relapses — “R” did not matter in the selection of patients. The width of the hernial ring varied from 4 to 25 cm. The most frequent were patients with the size of the hernia gate W5 (10–15 cm) — 23 people (34.8%).

In terms of the number of relapses there were patients both after midline laparotomies and undergoing surgical interventions for postoperative hernia, including alloplasty. Recurrent hernias were present in 21 (31.8%) patients, in other cases herniolysis was performed for the first time.

The primary defect of the anterior abdominal wall was formed as a result of planned and emergency surgical interventions. The most frequent operations were hernia of the anterior abdominal wall (white hernia of the abdomen, umbilical hernia, ventral hernia) — 10 people (15.2%), which later became postoperative ventral hernias reaching the size of the hernia gate 25 cm.

The period of treatment of patients for help from the onset of the hernial defect varied greatly and ranged from 6 months to 10 years. During the year only 23 people asked for help (34.8%), which once again confirms the medical illiteracy of patients and careless treatment of their health.

The risk assessment of the development of venous thromboembolic complications (VTEC) was also carried out on the Caprini scale.

In the postoperative period all patients received adequate analgesia on the first day after surgery;
Table 1. The distribution of patients in groups

<table>
<thead>
<tr>
<th></th>
<th>1st group</th>
<th>2d group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modeled (simulated) implant</td>
<td>14</td>
<td>17</td>
<td>31</td>
</tr>
<tr>
<td>Standard implant</td>
<td>18</td>
<td>17</td>
<td>35</td>
</tr>
<tr>
<td>Total</td>
<td>32</td>
<td>34</td>
<td>66</td>
</tr>
</tbody>
</table>

Fig. 1. Simulated and standard implants

Further analgesia was performed according to indications. If necessary, antibiotic therapy and prevention of thromboembolic complications were prescribed. Used tactics of early activation of patients using a bandage.

In the postoperative period all patients received adequate analgesia on the first day after surgery; further analgesia was not according to indications. If necessary antibiotic therapy and thromboembolic complications were prescribed, same tactics on patients with the bandage.

In the immediate postoperative period postoperative wound suppuration occurred in patients with a history of secondary wound healing after surgery despite the ongoing massive antibiotic therapy and daily wound dressings. Complications associated with the formation of seroma and suppurative hematoma were also conservatively resolved; no additional surgery was required (Table 2).

In the late postoperative period the follow-up period for patients was from 2 to 8 years. Disease recurrence was recorded in 6 (9.1%) patients. However, only one patient under 60 years of age had a relapse with a simulated implant, the cause of the recurrence was the failure to follow the clinical guidelines in the first 6 months after the surgery. In the remaining 5 (7.5%) cases no reliable data on the cause of the relapse was obtained (Table 2).

Table 2. Complications in the early and late postoperative period

<table>
<thead>
<tr>
<th></th>
<th>1st group</th>
<th>2d group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>B</td>
<td>A</td>
</tr>
<tr>
<td>Postoperative wound suppuration</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Suppurative hematoma</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Seroma</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Foreign body sensation</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Relapse</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

Despite of the volume of performed examinations it was impossible to avoid the complications of chronic diseases completely. Even with the condition of early activation of patients on the 6th day after hernioplasty, one patient in the 2nd group experienced acute intestinal obstruction associated with adhesions in the abdominal cavity. On the background of the ongoing conservative therapy positive dynamics were useless. The patient underwent surgery in the volume...
of laparotomy, adhesions dissection, nasointestinal intestinal intubation. The implant was retained during surgery. Further postoperative period showed no interest.

CONCLUSION

The simulated (modeled) implant shows good results of treatment in the long term in both young and old patients, its introduction and use in a wide surgical practice can be recommended for postoperative ventral hernias of the middle localization.

REFERENCES

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