NEW ALTERNATIVE FOR WOUND PROTECTION IN LAPAROSCOPIC COLECTOMY

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ABSTRACT - Background: Large number of surgical services use laparoscopy to approach the colon. One of the concerns on the resection using this way is the high rate of cancer relapse at in- and outlet site of the surgical instruments. Aim: To describe a protective device for surgical isolation in laparoscopic colectomy. Methods: The device is made of sterile polyethylene plastic cover used to protect the fiber optic cable in laparoscopic surgery and one 20 Fr. urethral catheter working as a conduit. Results: The device was used in six laparoscopic colectomies, three for adenocarcinoma of the colon and three for intestinal endometriosis. It was effective to avoid contact of the specimen with the abdominal wall, in order to reduce the risk of implantation of cancer or endometriotic cells and surgical site infection. The device was made intraoperative at all surgeries and allowed good visualization in laparoscopy and maintenance of the pneumoperitoneum. It cost R$ 22,00 (approximately US$ 10), R$14.50 related to the plastic cover and R$7.50, the urethral tube. The production time of the device and its installation in the abdominal cavity was measured in each procedure and was, on average, respectively, of 66 s and 25 s. Conclusion: The device proved to be feasible, not requiring any special training and can be performed by the surgical team itself, even at institutions with limited resources.

INTRODUCTION

In recent years many surgical services have addressed the colon laparoscopically, both for the treatment of benign diseases as well as for tumors, with similar results to laparotomy24. Initially, the laparoscopic surgical treatment of colon diseases was restricted to malignant cases. The concern was in relation to tumor resection, ie if it assures appropriate oncologic attendance and, also, with the high rate of cancer relapse in the incisions used by laparoscopic surgical devices1. Studies have shown alarming recurrence of colonic neoplasia in these incisions in up to 21%12. This led, in mid-90s, the American Society of Colon and Rectal Surgeons to recommend that the treatment of colon cancer by laparoscopy should be performed only in controlled trials3.

Several studies were conducted to elucidate the reason for this so high incidence1. In 2007 Fleshman et al, based on the COST study (tested with follow-up of five years), showed that there is no statistical difference in tumor recurrence in the surgical wound when comparing laparotomic or laparoscopic colectomy6. Other studies corroborate this finding and found that most responsible for this high frequency of recurrence were technical devices related to laparoscopy - under development at that time6.
The surgical wound provides rich environment factors that support the growth of cancer cells. The large enough contact of these cells in the incision allows them to be implanted and developed. It is therefore crucial, in order to avoid this complication, to prevent contact of tumor cells with the abdominal wall.

Endometriosis, although considered benign, has typically neoplastic characteristics, with invasion capacity to adjacent tissue and also causing distant implantation, and similarly to cancer, can be implanted in the abdominal wall. The pathophysiological explanation for this behavior is controversial, but the metastatic theory is advocated by some authors, who maintain that the handling and the contact of endometrial tissue with the surgical incision would be a condition for the development of the disease in this site.

Routine use of protective devices to isolate the intestine during its withdrawal from the abdominal cavity, can be useful in the prevention of cells implantation into the abdominal wall. In addition, the decrease of the exchange of surgical instruments through the portals, careful handling to avoid the trauma of surgical specimen and control of pneumoperitoneum loss, are other important care that should also be observed.

In addition, some studies have shown that the use of specimen protector on withdrawal from cavity in digestive operations reduces the occurrence of site infection. This complication is very common in colorectal surgeries, reaching ratio of 26% in some studies. According to published data, the proportion drops from 22.7% without the protective use to 4.7% when the device is used. In laparoscopy, the risk of infection is lower, but the numbers are still relevant (6.6%).

Thus, it is advisable to standardize medical practice using devices that allow the isolation of the surgical specimens during its withdrawal in laparoscopic procedures in patients with colon cancer or intestinal endometriosis. Considering the high prevalence of disease in focus, it is important to facilitate the effective production of protective devices at low cost, to enable the wide use of this type of device.

The objective of this study is to describe a sterile device for specimen isolation in laparoscopic colectomy.

**METHODS**

The study was conducted at Hospital Santa Rita and University Hospital Cassiano Antônio de Moraes, Vitória, ES, Brazil, after approval by the Ethics in Research Committee of Santa Casa de Misericórdia under number 121/2011.

The use of the device occurred in six patients, who were informed about the study and signed a consent form. Were selected by convenience - three patients with adenocarcinoma and three of intestinal endometriosis - and underwent laparoscopic colectomy. In surgical procedures were used automatic staplers for anastomosis of intestinal segments. The surgical wound site was evaluated postoperatively in all patients.

**Preparation technique of the protective device**

The materials used were: 1) polyethylene sterile plastic used in laparoscopic surgery to surround the fiber optic cable, and 2) urethral catheter 20Fr in diameter. The catheter was shaped as a ring. The plastic was then inserted through the ring and everted completely to form a hollow cylinder - as a conduit - with about 20 cm long (Figure 1).

After this step, the plastic edge of the conduit was sealed by a Kelly clamp (Figure 2). The ring formed by the catheter was then collapsed between the surgeon's fingers before insertion into the abdominal cavity, which occurred through a transverse 5 cm incision above the pubis. Inside the cavity, the counterforce of urethral catheter caused the ring to open up, positioning itself in the parietal peritoneum around the incision (Figure 3).
(at the incision), allowing the first clamp be released and the end of the conduit be freed. After dissection of the affected intestinal segment, was introduced a grasping forceps in the free end of the conduit, and the surgeon compressing the plastic cover with his hand against this clamp, maintained pneumoperitoneum after detaching the second Kelly clamp. The steps described before since insertion of the device into the abdominal cavity, are shown in Figure 4.

FIGURE 4 – Device handling

Finally, it was possible to achieve the segment that was externalized with the grasping forceps; at this time, the pulled gut occupied the device lumen and prevented air leakage, allowing the surgeon to do the external resection of the specimen (Figure 5). After the manipulation, the intestine was reintroduced into the cavity and the edge of the orifice was sealed again with a Kelly clamp, proceeding with the laparoscopic operation.

FIGURE 5 – Intestine was exteriorized through the orifice (A) and resected outside the abdominal cavity (B)

RESULTS

In the three cases of colon adenocarcinoma right colon was involved at early stage of the disease, with tumors restricted to the intestinal wall. The affected bowel segment was externalized through the device, being resected with laterolateral anastomosis - terminal ileum with transverse colon - using linear stapler with 80 mm cartridge.

The three cases of bowel endometriosis had involvement of the rectosigmoid junction. In such cases, after dissection of the affected intestine, was used an articulated linear endoscopic stapler of 45 mm in upper rectal disease free, sectioning the intestinal segment proximal to the lesion to be resected, and stapling the distal portion. The proximal part was then externalized through the orifice and resected. In the remaining segment was inserted circular stapler warhead, being contained by purse suture. Then the specimen was returned to the abdominal cavity. Rectally, was then introduced the second part of the circular stapler, which has been coupled to the warhead in order to complete the anastomosis between the remaining portion of the colon and rectum. The integrity of the anastomosis was confirmed by blowing up air through Foley catheter via rectal route, known as “tire repairman maneuver”.

In one of the cases of endometriosis, following the procedure described above and during the finalization of the abdominal cavity, was found an implantation of 3 cm of the disease in the cecum. Due to lack of laparoscopic material for further operation it was converted to laparotomy, being enlarged the incision and performed right colectomy without use of the protective device.

Each device cost approximately US$ 10, referring to the plastic cover and the urethral catheter. The average time for device manufacturing was 66 seconds, and conduit insertion into the abdominal cavity was 25 seconds - from the parietal peritoneum incision to insertion of the ring with plastic cover filled by pneumoperitoneum (Table 1).

TABLE 1 – Time for device manufacture and control of pneumoperitoneum after the incision of the parietal peritoneum in the chronological order of operations

| Procedure | Device Manufacturing (seconds) | Pneumoperitoneum Control (seconds) |
|-----------|-------------------------------|----------------------------------|
| 1st       | 0:50                          | 28                               |
| 2nd       | 0:08                          | 22                               |
| 3rd       | 0:58                          | 30                               |
| 4th       | 0:54                          | 25                               |
| 5th       | 0:00                          | 24                               |
| 6th       | 0:48                          | 21                               |
| Average   | 0:06                          | 25                               |

The part of the device that was inserted into the abdominal cavity allowed good visualization in laparoscopy.

The patient with endometriosis with operation converted to laparotomy developed an abscess on the 7th postoperative day. Drainage and antibiotics, solved the situation. The remaining cases did not present complications during the follow-up period of 30 days.

DISCUSSION

The presented device proved to be an alternative of low cost, easy to manufacture, effective to prevent contact of the specimen with the abdominal wall and maintaining pneumoperitoneum in laparoscopic operation.

The removal of the colon without contact with the surgical wound by laparoscopy is still a challenge for most surgeons in our country. Companies that provide laparoscopic material - to the date of this article -, do not routinely provide any kind of conduit device that allows the viscera externalization, as described herein. The device offers the possibility of maintaining the pneumoperitoneum - paramount in laparoscopic procedures - allowing careful review of the abdominal cavity even after intestinal resection and externalization. In all cases of endometriosis in this series, was possible to maintain the pneumoperitoneum to test the integrity of the anastomosis performed laparoscopically and, in one instance, the review showed other unidentified lesions in the initial inventory.

The low cost of manufacture of this product is an unquestionable advantage in the context of a national health system, with lack of resources and the high prevalence of the diseases under discussion. Seeking in the market other protective devices for colectomy, it was found that there are many options. The name Alexis® device of Applied Medical has an average value of US$ 70 - minus import costs, taxes and intermediaries - , so, much more expensive than device presented here. Parts of the device are materials often used in

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health institutions, making the use of a viable tool in centers with fewer resources. Its easy making with no need of special training, is perfectly feasible to be made by the surgical team during surgery. The short conduit assembly time probably does not interfere significantly in the total length of surgery, but when in advance is certain that intestinal resection is be performed, the preparation can be carried out before the operation.

Although the main objective of this article was to describe the device, it is believed that, later, would be interesting to realize a controlled study with representative sample of patients in order to assess its effectiveness in preventing implantation of endometriotic and neoplastic cells in the surgical wound and infection control.

CONCLUSION

The preparation of the device presented here proved to be feasible and requires no special training and can be performed by the surgical team at reduced costs.

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