Abstract

Hintergrund und Ziele: Die laparoskopische Reparation ventraler Narbenbrüche umfasst das Implantieren eines synthetischen Netzes in das Abdomen. Besorgniserregend ist hierbei die mögliche Bildung schwerwiegender viszeraler Adhäsionen an die Prothese. Insgesamt gibt es nur sehr wenig klinische Informationen über Adhäsionen an Biomaterial, welches intraabdominell platziert wurde, die auf Ergebnissen bei reoperierten Patienten beruhen. Wir führten eine Multizenterstudie zu Adhäsionen an implantierten Netzen aus expandiertem Polytetrafluorethylen (ePTFE) durch, bei der wir Patienten untersuchten, die sich zuvor einer laparoskopischen Narbenhochbehandlung unterzogen, wobei die gleiche Technik zum Implantieren des Netzes verwendet wurde.

Methoden: Bei insgesamt 65 Patienten beurteilten 9 Chirurgen retrospektiv den Schweregrad der Adhäsionen an das ePTFE-Netz bei der Nachoperation. Für jeden Fall wurde der Schweregrad der Adhäsion nach einer Skala von 0–3 bewertet, wobei 0 keine Adhäsion und 3 schwere Adhäsionen bedeutete.

Ergebnisse: Die durchschnittliche Zeitspanne von der Netzimplantation bis zur Nachoperation betrug 420 Tage (bei einer Spanne von 2–1739 Tagen). In 15 Fällen wurden keine Adhäsionen beobachtet. In 44 Fällen wurden Adhäsionen des Schweregrades 1 und in 6 Fällen des Schweregrades 2 festgestellt. Es wurden keine Adhäsionen mit Schweregrad 3 beurteilt. Somit waren 45 Patienten (69%) entweder keine oder ganz leichte avaskuläre Adhäsionen auf. Während der Débridements traten keine Enterotomien auf.

Background and Objectives: Laparoscopic ventral incisional hernia repair involves intraabdominal placement of a synthetic mesh, and the possibility of formation of severe visceral adhesions to the prosthesis is a principal concern. Little clinical information based on reoperative findings is available about adhesions to biomaterials placed intraabdominally. We conducted a multi-institutional study of adhesions to implanted expanded polytetrafluoroethylene (ePTFE) mesh at reoperation in patients who had previously undergone laparoscopic incisional hernia repair done with the same mesh implantation technique.

Methods: Nine surgeons retrospectively assessed the severity of adhesions to ePTFE mesh at reoperation in 65 patients. For each case, adhesions were assigned a score of 0 to 3, with 0 indicating no adhesions and 3 severe adhesions.

Results: The mean time from mesh implantation to reoperation was 420 days (range, 2–1739 days). No adhesions were observed in 15 cases. Forty-four cases received an adhesion score of 1, and 6 cases a score of 2; no scores of 3 were assigned. Thus, 59 patients (91%) had either no or filmy, avascular adhesions. No enterotomies occurred during adhesiolysis.

Conclusions: In this large series of reoperations after laparoscopic incisional hernia repair, no or minimal formation of adhesions to implanted ePTFE mesh was observed in 91% of cases, and no severe cohesive adhesions were found. Comparative analyses of newer materials based on clinical reoperative fin-
Schlussfolgerung: Bei dieser großen Anzahl von nachoperierten Patienten nach laparoskopischer Reparation von Narbenbrüchen konnte bei 91% der untersuchten Fälle keine oder nur minimale Adhäsionsbildung an das implantierte ePTFE-Netz beobachtet werden und es wurden keine schweren kohäsiven Adhäsionen festgestellt. Vergleichende Analysen von aktuellere Ergebnissen klinischer Nachoperationen sind gerechtfertigt, um die Sicherheit von intraabdominell implantierten Netzen beurteilen zu können.

Schlüsselwörter
Ventraler Narbenbruch · Adhäsionen · Laparoskopie · Polytetrafluorethylen

The need for reinforcement of the abdominal wall by implantation of synthetic materials during repair of abdominal incisional hernias is well established [1–4]. Moreover, because early use of various biomaterials in ventral incisional hernia repair resulted in some serious complications, including fistulization due to adhesions [5–7], it is clear that avoiding contact between the mesh and viscera is desirable [2, 3, 8, 9]. In laparoscopic incisional herniorrhaphy, however, first described by Leblanc and Booth in 1993 [10], placement of the mesh in the intraabdominal position is necessary. Thus, this procedure requires a biomaterial that provides both adequate strength and a minimal risk of severe adhesion formation that could result in bowel obstruction or fistula development. Numerous studies of the formation of adhesions to biomaterials placed intraabdominally have been conducted in animals, but few investigations of this issue have been based on information obtained at reoperation in patients. Moreover, no large study has examined reoperative observations pertaining to adhesions to a biomaterial after laparoscopic incisional hernia repair specifically. Therefore, we conducted a retrospective review of findings at reoperation in patients who had undergone such a repair and in whom a 1-mm-thick expanded polytetrafluoroethylene (ePTFE) dual–surface mesh (Dual-Mesh, W.L. Gore & Associates, Flagstaff, AZ) had been implanted during the repair.

Methods and materials
Records of adult patients who had undergone a laparoscopic repair of a ventral incisional hernia including placement of Dual-Mesh intraabdominally between April 1993 and April 2001 were reviewed after institutional IRB approval. Patients who subsequently underwent abdominal reoperation for various indications, during the same time period, were selected for retrospective review. The technique used for initial mesh placement was essentially the same in all cases and was described in detail previously [11]. Briefly, the mesh was placed laparoscopically in the intraabdominal onlay position and secured with sutures as well as circumferentially placed 5-mm titanium tacks. In some cases, titanium staples were used around the periphery. For the review, the nine surgeons (authors) participating in the study completed a questionnaire about findings at reoperation in each case meeting the criteria described above. The questionnaire included a section asking the surgeons to score, retrospectively on the basis of the operative reports, the adhesions encountered according to the severity scale described by Diamond [12]. Thus, a score of 0 was to be assigned if no adhesions were present; a score of 1 if there were filmy, avascular adhesions; a score of 2 for vascular and/or dense adhesions; and a score of 3 for cohesive adhesions. In all cases, the same surgeon who performed the original repair completed the questionnaire on findings at reoperation. Data from the questionnaires were compiled according to adhesion score and surgeon. In addition, patient demographics, time from initial mesh implantation, and reason for reoperation were included.

Results
Sixty-five patients were included in the study. There were 35 females and 30 males, with ages ranging from 27–86 (average of 55) years old. The mean time between the laparoscopic repair including mesh implantation and reoperation was 420 days (range, 2–1739 days). Seventeen of the patients underwent reoperation because of hernia recurrence as a result of either mesh lodgment due to inadequate fixation or infection necessitating removal of the mesh. Ten patients required reoperation because of complications, including early infection, delayed enterotomy, and bowel obstruction; none of the cases of obstruction were due to adhesions to the prosthesis. The remaining 38 patients had a new diagnosis requiring surgery. The adhesion scores reported by the surgeons are shown in Table 1. No adhesions to the mesh were observed in 15 cases (23%). In 44 cases (68%), the adhesions observed were filmy and avascular (score of 1). Vascular and/or dense adhesions (score of 2) were found in 6 cases (9%). and no adhesions scores of 3 were assigned. There was no significant difference in adhesion scores with respect to implant duration, in particular with respect to patients with a score of 0 compared with those with a score of 2 (Table 2–4). Therefore, at reoperation in 91% of patients who had undergone laparoscopic abdominal hernia repair, adhesions were either not present or were filmy and avascular. All the dense adhesions (adhesion score 2) involved omentum only, and most involved only the exposed portions of the titanium tacks. Dissection of adhesions to the mesh required little effort, and no enterotomies occurred during adhesiolysis. Representative photographs obtained at reoperation are shown in Fig. 1–4. In early reoperations (fewer than 14 days after the original repair), no adhesions to the mesh were observed (Fig. 1), although in several cases, adhesions had already formed to the exposed titanium tacks (Fig. 2). At reoperations...
Table 1 Adhesion scores for 65 patients at reoperation, according to surgeon

| Surgeon | Adhesion Score * |       |       |       |       |
|---------|------------------|-------|-------|-------|-------|
|         | 0           | 1     | 2     | Total |
| 1       | 1           | 6     | 3     | 10    |
| 2       | 4           | 4     | 0     | 8     |
| 3       | 1           | 7     | 1     | 9     |
| 4       | 4           | 5     | 0     | 9     |
| 5       | 0           | 9     | 0     | 9     |
| 6       | 3           | 5     | 0     | 8     |
| 7       | 2           | 3     | 1     | 6     |
| 8       | 0           | 5     | 1     | 6     |
| Total   | 15          | 44    | 6     | 65    |

* Adhesions were scored according to the severity criteria of Diamond24; thus, 0 indicated no adhesions; 1, filmy, avascular adhesions; 2, vascular and/or dense adhesions; and 3, cohesive adhesions. No adhesions received a score of 3. Values are numbers of patients with each score.

† Nine surgeons participated in the study, but 2 surgeons in the same practice reported their results together.

Table 2 Patients with adhesion scores of 0 versus mesh implant duration

Table 3 Patients with adhesion scores of 1 versus mesh implant duration

Table 4 Patients with adhesion scores of 2 versus mesh implant duration

Fig. 1 Appearance of DualMesh 5 days after implantation in a patient in whom an incarcerated inguinal hernia developed 5 days after repair of a previously incarcerated flank hernia in a nephrectomy incision. There are no adhesions to the mesh.

Fig. 2 Adhesion of viscera to titanium tack 12 days after incisional hernia repair in a patient in whom a small-bowel obstruction developed as a result of enteroenteric adhesions unrelated to the repair. There are no adhesions to the DualMesh done after 2 weeks or longer after the laparoscopic repair, a “neoperitoneum” across the mesh was frequently observed (Fig. 3). This membrane could be removed intact from the underlying mesh surface (Fig. 4).

Discussion

In a multi-institutional series of 65 reoperations after laparoscopic ventral incisional hernia repair in which the same technique for mesh implantation had been used in all patients, we observed either no adhesions to DualMesh or filmy, avascular adhesions in 59 (91%) cases. These findings are in agreement with the few other clinical reports of reoperations after implantation of ePTFE in laparoscopic incisional hernia repair. Gillion et al observed no adhesions during 6 reoperations after ePTFE had been placed in the intraperitoneal or extraperitoneal position for re-
pair of incisional hernias of the anterolateral abdominal wall [13], Koller et al. found minimal adhesions to ePTFE during 2 reoperations [14]. We also observed, in patients in whom the DualMesh had been implanted for greater than two weeks, a well-vascularized “neoperitoneum” across the entire mesh surface, similar to that described by Bellon et al in experimental and clinical studies of ePTFE [15, 16], by Gillion et al. in their series of clinical reoperations [13], and by Carbajo et al., who observed complete reperitonealization of DualMesh in the 2 reoperations they described [17]. Avoiding contact between prosthetic mesh and viscera to minimize formation of adhesions after ventral incisional hernia repair was a central concern even before development of the laparoscopic approach. In reports describing open repair, Stoppa [2] and Wantz [3] advised that contact between an intraperitoneally placed prosthesis (polyester) and the viscera should be prevented by covering the pros-

thesis with omentum (if possible) or by inserting an absorbable synthetic prosthesis. Condon noted that because of the intense inflammatory reaction caused by polypropylene mesh, the bowel should be protected from that material [9]. In a review of their experience with the Stoppa technique, Temudom et al emphasized the importance of positioning autogenous tissue between the bowel and the surface of polypropylene mesh [18]. Some authors, however, still recommend direct intraabdominal placement of polyester mesh [19]. There have been numerous reports of complications resulting from placement of mesh in contact with the viscera. Many of these described erosion or fistula formation occurring with use of polypropylene mesh in trauma cases in which extensive contamination was present, the mesh was in contact with the viscera by necessity, and no tissue covered the mesh externally [5, 6, 20 – 22]. However, fistula formation after implantation of various biomaterials in the abdominal wall has also been observed in several nontrauma cases [7, 22]. With the laparoscopic approach to abdominal incisional hernia repair, peritoneal dissection and covering of the mesh is often impossible with true incisional hernias. Holzman et al observed that efforts to separate the peritoneum of the hernia sac may produce a large peritoneal defect and leave the mesh exposed [23]. They suggested omental interposition, as did Franklin et al. [24], and others have reported that this technique can reduce or eliminate adhesion of viscera to polypropylene [22, 22]. However, in the series of 19 reoperative cases described by Franklin et al. [24], in which this omental protection technique had presumably been used during the previous hernia repair, one third of patients had severe adhesions to the polypropylene mesh. Moreover, covering mesh with omentum can be difficult because of the large defects being repaired laparoscopically. Park et al., in their early experience with laparoscopic repair of large incisional hernias [25], used mesh sizes as large as 530 cm². Koehler and Voeller [26] reported that 38% of their cases involved mesh of at least 18 cm × 24 cm and that 9% required 2 pieces of mesh sewn together. Other reports of laparoscopic repairs reveal similar experiences [11, 17, 27]. If an absorbable mesh is inserted along with a nonabsorbable prosthesis to prevent temporary contact between the nonabsorbable material and the viscera – a method recommended by Wantz [3] and others [28] – the time during which peritoneal coverage is provided is a concern. One study of adhesion formation in a rat model suggested that most adhesions were fully developed by 7 days after implantation of mesh [29]; however, Amid expressed caution against such results that suggest a temporary barrier is needed for only a short time [30]. There have been no clinical studies based on findings at reoperation indicating that absorbable barriers are effective, and Luijendijk et al. [4] stated that neither experimental nor clinical studies have provided conclusive information on the efficacy of interposition of polyglactin mesh in preventing adhesions, bowel obstruction, and fistulas. We believe that placement of an absorbable barrier to a large mesh that is subsequently fixed to the moving environment of the anterior abdominal wall represents an improbable solution. In contrast to the experience with some other materials, early experiences with ePTFE showed that it was apparently effective in limiting formation of adhesions on the visceral side, but limited fascial surface ingrowth was a concern. DualMesh was developed to address this concern. This mesh is composed of 2 layers of ePTFE with different surface characteristics; one layer has a “smooth,” nonporous surface to

Fig. 3 Appearance of DualMesh at reoperation 20 months after implantation. A neoperitoneum and extensive vascularization are present across the surface of a large piece of mesh (18 cm × 24 cm).

Fig. 4 Removal of neoperitoneum from DualMesh surface in the same patient as in Fig. 3. DualMesh resulted in no adhesion formation.
be placed against the viscera to limit adhesions and the other has a “rough” surface with large interstices in the material (22 μm) to encourage tissue ingrowth. DualMesh has so far been used in more than 1000 laparoscopic ventral and incisional hernia repairs [11, 26, 27, 31, 32] with good results with respect to recurrences and complications. The current study of 65 reoperations supports the assumption that no or minimal adhesions result from the use of DualMesh in such procedures. Interestingly, we observed no adhesions to this type of mesh in patients in whom adhesions to titanium tacks developed, a finding indicating that even in patients with a possible tendency to form adhesions, DualMesh serves to limit such formation. Although this study has the limitation of being a retrospective analysis of operative findings, all the surgeons involved had extensive experience with laparoscopic repair of incisional hernias and with clinical studies of this procedure, including assessments of the adhesiogenic properties of biomaterials. Any reoperations necessary after incisional hernia repairs were therefore analyzed and described carefully in operative reports, and particular attention was paid to findings regarding adhesions. This study, like several others, used “Diamond scores” to describe the severity of adhesions; we did not use the quantitative portion of the Diamond criteria (ie, determination of the percentage of surface area involved with adhesions), since our primary interest was to assess how difficult it was to separate the omentum or viscera during adhesiolyis. We think, however, that Diamond scores have important limitations in representing operative situations. For example, easily sweeping aside several hundred square centimeters of filmy adhesions may take less than a minute, without incident, whereas removing a few square centimeters of dense adhesions can lead to an enterotomy. In addition, making comparisons of studies using Diamond scores is difficult because of varying assumptions about what constitutes mild and severe adhesions. Therefore, we favor an adhesion scale that reflects the ease or difficulty with which adhesiolyis is accomplished. That described by Zuhlke et al. [33] is close to achieving this goal. On this scale, 1 denotes adhesions that are filmy and easy to separate; 2, adhesions that may be removed by blunt dissection, but partial sharp dissection may be required; 3, adhesions that are strongly attached and can be removed by sharp dissection only; and 4, adhesions that are so severe that injury to an organ is likely during dissection. Application of this scale to the reoperative findings in our study would result in all cases scored as 0 or 1 according to the Diamond scale being assigned a “Zuhlke score” of 1.

Conclusions

Laparoscopic ventral incisional hernia repair requires placement of nonabsorbable mesh in an intraabdominal location. Many studies of intraabdominal placement of synthetic meshes in animals have been done to assess the risk of visceral adhesion to these biomaterials, but little information has been available on reoperative findings after mesh implantation in patients. In this study, we investigated the use of one type of ePTFE mesh in a retrospective review of reoperations in patients in whom the technique used for mesh placement during laparoscopic repair of ventral incisional hernia was identical. Our findings suggest that DualMesh is effective in minimizing, and often eliminating, visceral adhesion formation after this repair. Research is continuing on new prostheses for abdominal wall hernia repair, especially meshes composed of a combination of absorbable and nonabsorbable materials. Clinical reoperative findings evaluated with use of relevant adhesion-scoring systems will be important in determining the comparative effectiveness of these new materials in preventing formation of visceral adhesions.

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Alte bayerische Krankenhäuser

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