Tissue Glue in Laparoscopic Inguinal Hernia Repair: A Retrospective Comparative Analysis

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Abstract

During the last two decades, there have been two revolutions in inguinal hernia repair surgery. First, the introduction of tension-free hernia repair by Liechtenstein in 1989 and then the application of laparoscopic surgery to the treatment of inguinal hernia in the early 1990s. In this context, the choice of mesh fixation methods being an integral part of this procedure remains a topic of arguments and discussions in laparoscopic inguinal hernia repair. There exist many methods of mesh fixation like polyglactin suture, titanium spiral tacks, nitinol anchors and fibrin glue.

Fixation usually uses staples that can lead to nerve injury and chronic postoperative pain. Laparoscopic repairs are associated with a risk of chronic pain of up to 22.5%. The use of fibrin glue may represent an alternative method of mesh fixation preventing the risk of nerve injury.

Keywords: Fibrin sealant, Tissue glue, Fibrin glue, Tisseel, Inguinal hernia, Laparoscopic herniorrhaphy, TAPP, TEP, Mesh fixation.

INTRODUCTION

Presently, the laparoscopic inguinal hernia repair is accomplished by two approaches: Transabdominal preperitoneal (TAPP) and totally extraperitoneal (TEP) repair with mesh prosthesis. TAPP is preferred as it manages all types of hernia (direct, oblique, femoral, and obturatory), whether unilateral or bilateral, primary or recurrent. Furthermore, because of its size the preperitoneal mesh prosthesis covers the entire musculopectineal foramen, where these hernias occur. Among various options available for mesh fixation, fibrin glue is being used increasingly as an alternative method for hernia surgery, significantly preventing the risk of nerve injury and helping to reduce the incidence of chronic pain. Reliable laparoscopic fixation of meshes prior to their fibrous incorporation minimizes recurrences following transabdominal preperitoneal hernia repair (TAPP) and totally extraperitoneal repair (TEP) repair of inguinal hernias. Various types of staples are available for reliable mesh fixation. However, their use has been associated with a certain amount of surgical trauma. Suture-tack- and staple-based fixation systems are associated with postoperative chronic inguinal pain. Reported complications include neuralgia or paresthesia because of nerve entrapments. A chronic form of pubalgia is caused by stapling of the prosthesis to Cooper’s ligament. Bleeding or hematomas in Retzius’ space (muscular, corona mortis) also may occur.

In 1997, Chevrel and Rath first proposed fibrin sealant as an alternate means of mesh fixation in hernia repair with the aim of reducing the rate of hernia recurrence. Canonico later reported the benefits of fibrin sealant in reducing bleeding complications following hernia repair in patients with impaired coagulation. Katkhouda employed a pig model using a total extraperitoneal (TEP) technique to evaluate the tensile strength of mesh fixation 12 days after the use of Tisseel®, demonstrating equal strength to staples. The results of these studies have encouraged surgeons to use fibrin sealant in daily practice as an atraumatic alternative to mechanical mesh fixation. As an atraumatic alternative, the application of fibrin glue (Tissucol/Tisseel, Baxter Healthcare, Deerfield, IL, USA) is a viable and reliable option, which keeps mesh in place without the complications associated with stapling. In terms of tensile strength and mesh dislocation, fibrin glue is equivalent to stapling.

AIMS

The aim of this review article is to evaluate the feasibility and efficacy of use of fibrin glue/tissue glue (tisseel) in laparoscopic inguinal hernia repair, short- and long-term postoperative pain, surgical complications (bleeding, seroma, hematoma, wound infection, incisional hernia, testicular complications) and recurrence rates. This article also emphasizes on evaluation of the advantages and disadvantages of fibrin glue as compared to other methods.
of mesh fixation to demonstrate whether fibrin-based mesh adhesion provides adequate biomechanical stability for repair of inguinal hernia by TAPP and TEP and to elucidate the extent to which tacks, anchor-based fixations can be replaced with fixation with fibrin glue/bioadhesives for laparoscopic inguinal hernia repair. The following parameters were also evaluated:
1. Patient selection
2. Operative technique
3. Operating time
4. Intra and postoperative complications
5. Postoperative pain
6. Hospital stay
7. Cost effectiveness.

MATERIALS AND METHODS

The literature utilized in this article were taken from search engine Google, SpringerLink library, HighWire press, Surgical endoscopy journal, World journal of surgery, Medscape. The following terms were used: Fibrin glue in laparoscopic hernia repair, TAPP, Fibrin sealant in hernia repair, role of fibrin glue in TAPP, TEP. The selected articles were screened for further references.

FIBRIN GLUE

Fibrin glue/sealant is a commercial tissue adhesive containing fibrinogen and thrombin. The commercial product is a two component system from human plasma that contains more than fibrinogen and thrombin. The first component contains highly concentrated fibrinogen, factor XIII, fibronectin, and traces of other plasma proteins. The second component contains thrombin, calcium chloride, and antifibrinolytic agent such as aprotinin (Table 1). Mixing of two components leads to activation of fibrinogen and thrombin by calcium chloride, formation and cross-linking of fibrin leading to the formation of polymerized fibrin chains, duplicating the last step of the coagulation cascade. The fibrinogen component gives tensile strength, thrombin stimulates fibroblast proliferation and aprotinin, an antifibrinolytic agent enhances the life span of the sealant.

The required dose of fibrin sealant depends on the size of the surface to be covered, as shown in Table 2.

Fibrin sealant contains the following substances in four separate vials:
1. Sealer protein concentrate (Human), vapor-heated, freeze-dried
2. Fibrinolysis inhibitor solution (Bovine)
3. Thrombin (Human), vapor-heated, freeze-dried
4. Calcium chloride solution.

Freeze-dried sealer protein concentrate and thrombin are reconstituted in fibrinolysis inhibitor solution and calcium chloride solution respectively (Flow Chart 1). The resulting sealer protein solution and thrombin solution are then combined (by using the duploject system, or equivalent delivery device) to form the fibrin sealant:

Various methods can be used to apply the two components of the sealant, the duploject and application needle being the most convenient and popular in laparoscopic surgery.

<table>
<thead>
<tr>
<th>Table 1: Composition of tissue glue</th>
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<tbody>
<tr>
<td>Package sizes</td>
</tr>
<tr>
<td>0.5 ml</td>
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<tr>
<td>------------------------------------</td>
</tr>
<tr>
<td><strong>Sealer protein concentrate</strong></td>
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<tr>
<td>Fibrinogen (mg)</td>
</tr>
<tr>
<td>Total protein (mg)</td>
</tr>
<tr>
<td>Polysorbate 80 (mg)</td>
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<tr>
<td>Sodium chloride (mg)</td>
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<tr>
<td>Trisodium citrate (mg)</td>
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<tr>
<td>Glycine (mg)</td>
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<tr>
<td>Fibrinolysis Inhibitor solution</td>
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<tr>
<td>Aprotinin (KIU)</td>
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<tr>
<td>Volume (ml)</td>
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<tr>
<td>Thrombin</td>
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<tr>
<td>Thrombin (IU)</td>
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<tr>
<td>Total protein (mg)</td>
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<tr>
<td>Sodium chloride (mg)</td>
</tr>
<tr>
<td>Glycine (mg)</td>
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<tr>
<td>Calcium chloride Solution</td>
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<tr>
<td>CaCl₂ (µmol)</td>
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<tr>
<td>Volume (ml)</td>
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<tr>
<td>Total combined</td>
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</table>
Commercially, fibrin sealant is available under the trade name of Tisseel, marketed by Baxter and is supplied in four different package sizes of 0.5, 1.0, 2.0 and 5.0 ml, containing the following components:

- Tisseel Kit 0.5 for 0.5 ml of reconstituted Tisseel solution and 0.5 ml thrombin solution.
- Tisseel Kit 1.0 for 1.0 ml of reconstituted Tisseel solution and 1.0 ml thrombin solution.
- Tisseel Kit 2.0 for 2.0 ml of reconstituted Tisseel solution and 2.0 ml thrombin solution.
- Tisseel Kit 5.0 for 5.0 ml of reconstituted Tisseel solution and 5.0 ml thrombin solution.

### OPERATIVE TECHNIQUE TAPP WITH FIBRIN GLUE MESH FIXATION

#### Patient Positioning

The procedure is performed under general anesthesia. The position of the patient is supine with a slight Trendelenburg tilt (15-20), legs together and arms alongside the body. The surgeon stands on the opposite side of the hernia and the assistant stands on the other side of the table. The scrub nurse and instrument table are beside the surgeon. The laparoscopy rack lies at the feet of the patient, in front of the surgical team.

#### Trocar Positioning

Pneumoperitoneum is achieved with a Veress needle inserted at the umbilical site. After an endoabdominal pressure of 12 to 14 mm Hg has been obtained, the first 10 mm trocar replaces the needle at the same site. A 30° scope is inserted. The other two trocars are inserted by transillumination under internal vision (with care taken to avoid the inferior epigastric vessels) at the level of the transverse umbilical line just lateral to the rectus sheath. The 5 to 12 mm operative trocar is always placed on the right hand side for both unilateral and bilateral hernias, and a 5 mm trocar is placed on the opposite side. For bilateral hernias, the two operative trocars are placed about 1 cm below the transverse umbilical line. For a unilateral hernia, the trocar would be positioned 1 cm above the line, at the intersection with the midclavicular line to create the classic triangulation of base ball diamond concept aimed at the surgical field with the trocars. The assistant operates the scope from the opposite side of the table. Evaluation of inguinal regions allows all defects of the transversalis fascia to be detected. The main landmarks are the remnants of umbilical artery, the ligament of Cooper, epigastric vessels, and the anterior superior iliac spine, all of which also allow definition of the hernia type.

#### Incision of the Peritoneum

If the hernia defect is on the right side, after the iliac spine is located by external pressure, the peritoneum is incised with the scissors at this point, and the incision is continued horizontally and medially. For a left-sided defect, the incision is performed from the lateral aspect of the umbilical artery and extended as far as the left iliac spine (Fig. 1). Peritoneal dissection follows the hernia orifice completely at about

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**Table 2: Amount of fibrin sealent required**

<table>
<thead>
<tr>
<th>Maximum size of the area to be sealed</th>
<th>Required package sizes of fibrin sealant</th>
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<tr>
<td>4 cm²</td>
<td>0.5 ml</td>
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<tr>
<td>8 cm²</td>
<td>1.0 ml</td>
</tr>
<tr>
<td>16 cm²</td>
<td>2.0 ml</td>
</tr>
<tr>
<td>40 cm²</td>
<td>5.0 ml</td>
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Flow Chart 1: Action of sealent

**Flow Chart 1:** Action of sealent

Sealer protein concentrate (human), vapor heated, freeze-dried → Thrombin (human), vapor heated, freeze dried → Calcium chloride solution → Fibrinolysis inhibitor solution → Liquid fibrin sealant → Solid fibrin sealant
Abhijit Mahanta, RK Mishra

0.5 to 1 cm from its upper margin, thus freeing the hernia sac from all visceral and parietal connections. Once the peritoneum is incised, it should be lifted bluntly with the scissors to allow seeping of the carbon dioxide (CO₂) to assist in peritoneal detachment. Forceps are used to lift the upper peritoneal margin, while scissors are used to complete peritoneal detachment and inferior epigastric vessel dissection.

**Parietalization**

The pressure of CO₂ entering between the peritoneum and abdominal wall helps dissection and basically requires two instruments: Scissors and 5 mm tissue forceps. The inferior peritoneal margin is pulled towards the surgeon with tissue forceps and bluntly dissected from the spermatic cord, which is parietalized to obtain an inverted triangle with the vas deferens running medially and the genital vessels running laterally between the iliac vessels in the so-called “triangle of disaster.” Parietalization of the spermatic cord is an important step in the procedure because it allows the free cord to be placed against the posterior wall (Fig. 2). The prosthesis then lodges perfectly on the inguinofemoral wall, thus closing the entire pectineal foramen. Next, the ligament of Cooper is identified by its characteristic consistency and greyish-white color, and dissected (Fig. 3). The dissection of Cooper’s ligament is performed bluntly with both instruments, which are divaricated to free the ligament of prevesical fat from the pubic symphysis to the external iliac vein. Once the ligament of Cooper has been prepared, the dissection proceeds towards the upper peritoneal flap, which is bluntly detached using the scissors or traction with two forceps cranially to complete the preparation of inguinofemoral wall.

**Prosthesis Positioning**

Once there is sufficient space for placement of the mesh, the next step is the positioning of prosthesis. Closing of hernia defect is achieved by means of a nonadsorbable,
wide-link mesh that enables the surgeon to recognize the structures on which it is lodged and fixed (Fig. 4). To avoid slippage and difficulty in positioning, the mesh should not be too soft. The mesh is introduced by grasping it on the medial and superior margin if the hernia is on the right, and by its superior and lateral margin if the hernia is on the left. Two forceps spread and positioned the mesh behind the peritoneal flaps against the posterior wall, so as to cover all hernia foramina, Cooper’s ligament, the “triangle of pain,” the “triangle of disaster,” the epigastric vessels, and the spermatic cord elements. It is important for the mesh to overlap the hernia foramen by at least 2 cm, and for its medial margin to be alongside the pubic symphysis.

**Fixation of the Prosthesis**

The mesh is fixed with 1 ml of Tissucol for unilateral hernias and 2 ml for bilateral hernias. The prosthesis is fixed along its upper margin, from Cooper’s ligament to the “triangle of disaster” and to the “triangle of pain,” using a 3 mm catheter (Duplotip; Baxter healthcare), which fits the Tisseel syringe. The mesh also may be fixed wherever necessary to increase its stability (Fig. 5). Tisseel may be applied in two different ways: By resting the tip of the duplotip catheter, where the mesh is to be fixed and by squeezing out a few drops of glue or the glue seeps across the mesh and fixes it. One can also separate the mesh slightly from the inguinal wall, spray the glue directly on it, and then place the mesh to the wall.

**Peritoneal Suture**

Closure of the peritoneal flap must be performed with extreme care to avoid leaving peritoneal breaches that could allow contact between the mesh and bowel loops.

Mesh prosthesis specifications and design lateral fixation near the triangle of pain Cooper’s ligament fixation with Tisseel.

**OPERATIVE TECHNIQUE TEP WITH FIBRIN GLUE MESH FIXATION**

The two components of the fibrin sealant (Tisseel, Baxter Healthcare Corporation) were reconstituted. Patient is placed in the supine position and general endotracheal anesthesia was induced. A curvilinear incision is made near the umbilicus and carried down to the anterior rectus sheath, which is doubly grasped, elevated, and incised, entering the rectus sheath. The rectus muscles are retracted laterally, exposing the posterior rectus sheath. A peritoneal dissection balloon trocar is inserted and guided by manual and videoscopic guidance down to the level of the pubis where it was inflated and left as such for several minutes for tamponade effect. It is then deflated, removed and replaced with a structural balloon trocar. Pneumoperitoneum was instituted under direct vision. Two 5 mm trocars are placed in the middle hypogastrium, one suprapubically and the second midway between pubis and umbilicus. Cooper’s ligament is identified along with cord structures and inferior epigastric vessels. The cord structures are skeletonized, and the hernia sac is reduced off the internal ring down to
the level of peritoneum. A vertical line is drawn down the long axis of the mesh by using a surgical marker to assist mesh orientation during the procedure. The mesh (8 cm × 13 cm or 10 cm × 15 cm) is placed through the preperitoneum unfurled, and placed uncut over the myopectinate orifice after soaking the mesh in saline for 10 to 15 minutes. The two solutions of the fibrin sealant were drawn into separate syringes, which are then fitted into the laparoscopic applicator (Fig. 6). Once the mesh is deployed in the position desired, it is secured to the pubic bone (Cooper's ligament) in the midline, the lacunar ligament laterally, and superiorly into the transversalis fascia with the fibrin sealant, which is allowed to set for several minutes. The posterior aspect of the matrix repair is then held in place as the pneumopreperitoneum is released under direct vision, observing the peritoneum to obtain its desired position relative to the matrix repair (Fig. 7).

In a study by S Olmi et al.1,10 on fibrin glue mesh fixation in TAPP in 230 patients with 320 hernias (unilateral and bilateral) the results had been encouraging. No perioperative complications were observed. After an average follow-up period of 26 months (range, 1-40 months), the only postoperative complications observed were six seromas (1.8%) and one trocar-site hematoma (0.3%). The mean operating time was 30 minutes for unilateral hernias and 50 minutes for bilateral hernias, whether primary or recurrent. Patients usually were discharged the day after surgery and returned to work after five days. No patient experienced an inguinal hematoma, which sometimes occurs after stapling. After a mean follow-up period of 26 months (range, 1-40 months), none of the patients reported immediate or late paresthesia or neuralgia. Evaluation of pain levels by the use of a visual analog score indicated that these patients were free of pain by day 7 after surgery. The average operating time was 30 minutes for unilateral hernias (range, 15-45 minutes) and 50 minutes (range, 30-75 minutes) for bilateral hernias, both primary and recurrent.

According to a randomized trial reported in 93rd Annual Clinical Congress of the American College of Surgeons (ACS), fibrin glue was as effective as staples in preventing hernia recurrence. At the same time, fibrin glue produces less of the postoperative pain experienced with staples, which may cause nerve entrapment. This study enrolled 100 patients scheduled to undergo totally endoscopic
preperitoneal inguinal hernia repair. Half of the subjects were randomly assigned to fibrin glue and half to staples for mesh fixation. All patients completed the protocol and were available for evaluation for one year after the procedure. Pain was reduced with the fibrin glue by several measures. On the first postoperative day, significantly fewer patients in the glue group reported mild pain (28% vs 46% of those receiving staples), and there was a nonsignificant trend toward more glue patients reporting no pain (68% for glue, 42% for staples). One week after the procedure, pain measures still showed a significant advantage for the glue. Using a 10-point visual analogue scale, patients who received glue reported pain in the 0 to 2 range, while those received staples reported pain in the range of 3 to 6. Patients in the glue group also consumed significantly less pain medication (oral diclofenac and paracetamol): A mean of 4.5 tablets per day vs 7.0 tablets per day for the staple group.

In a study by Stark et al14 the rate of nerve entrapment in laparoscopic patients was 4.2%. The genitofemoral nerve was affected with a high frequency (2%), and the ilioinguinal or lateral cutaneous nerve of the thigh was affected in 1.1% of the cases.

The morbidity in postoperative period in patients with staples and fibrin glue mesh fixation is a topic of interest. Federico Lovisetto et al15 carried out a randomized study in 197 patients with inguinal or femoral hernia. The primary outcomes were early postoperative and late neuralgia recorded using a visual analog scale (VAS). The effects of neuralgia on functional status were evaluated using the modified SF-36 questionnaire. Secondary outcomes included complications, such as nonspecific pain and recurrence. His study included 176 males (89.3%) and 21 females (10.7%) patients with a mean age of 53 years (range 18-79 years); 188 (95.4%) hernias were inguinal and nine (4.6%) were femoral. Patient characteristics were similar in the two groups. Follow-up visits were done in 1, 3, 6, and 12 months. No intraoperative complications were observed in either of the treatment groups. When compared, there was no significant difference in quality of life in the two study groups. The mean duration of intervention was 54 minutes (range 30-95 minutes) in the Tisseel/fibrin glue group vs 40 minutes (range 25-105 minutes) in the staples group. The mean postoperative hospitalization time in each group was one day. The mean recovery time to normal physical activity was 7.9 days (range 5-11 days) in the fibrin glue group vs 9.1 days (range 7-11 days) in the staples group. Early postoperative complications which included hematoma/seroma, orchitis, nonspecific pain occurred in 8% of patients in the fibrin group and in 12% of patients in the staples group. The percentage of late postoperative complications was 3 and 7% of patients in the fibrin glue and staples groups respectively. There were no cases of hernia recurrences.

Over three years, Graziano Ceccarelli et al17 compared the characteristics of mesh fixation with titanium clips and fibrin glue (Tisseel) and evaluated if the use of fibrin sealant was as safe and effective as conventional stapling and if there were differences in postoperative pain, complications and recurrences. Comparison was made between two homogeneous groups of 68 patients (83 cases) treated with fibrin glue and 68 patients (87 cases), where the mesh was fixed with staples. TAPP technique was used. Operative times were longer in the group treated with fibrin glue with a mean of 35 minutes (range 22-65 mins) compared to the group treated with staples (25 minutes, range 14-50 mins). The time of hospital stay was the same (24 hours). Postoperative complications that were more frequent in the stapled group, included trocar site pain, hematomas, intraoperative bleedings and incisional hernias. No significant difference was observed concerning seromas, chronic pain and recurrence rate.

A study by Arthur P Fine et al involved 38 adult patients with 51 inguinal hernias. 38 patients with 45 primary and six recurrent inguinal hernias were treated with laparoscopic repair by the total extraperitoneal mesh placement (TEP) technique using mesh secured with fibrin sealant. All patients could care for themselves within a day of surgery. Immediately following surgery, patients on average took 7 days off from work. Short-term pain associated with the procedure generally subsided completely within three to four days. Follow-up examination two weeks after the surgery revealed no swelling or localized abdominal pain, and the patient was allowed full activity and returned to work. Postoperative complications were minor and generally expected as a consequence of surgery. Mild cord or canal swelling or both, following the procedure being the most common complaint. The degree of swelling was not judged as severe or necessitating intervention in any case. Mild to moderate orchitis was noted in three patients (7.9%) following surgery, as were two cases of hematoma (5.3%) and one suspected case of seroma (2.6%) that resolved without intervention. Two patients (5.3%) presented with mild fever and localized pain and swelling following surgery. Ciprofloxacin was given for suspected infection, and both patients eventually resolved without further intervention.

P Topart et al3 in his retrospective analysis of Tisseel vs tack staples as mesh fixation in totally extraperitoneal laparoscopic repair of groin hernias compared the result of
66 patients with fibrin glue mesh fixation in totally extraperitoneal (TEP) laparoscopic procedure with an earlier series of 102 patients operated on according to the same procedure in which mesh fixation used tack staples. For the fibrin glue group, the operative time was 54 ± 23 minutes and no difficulty was encountered during preparation or application of the fibrin sealant. There were no reoperations or postoperative deaths. The postoperative course was uneventful for 53 patients (80.3%). Eight patients (12%) had a seroma, which did not require any dedicated treatment in the majority of cases. Three patients (4.5%) had a hematoma: One patient had to remain on calciparin at the time of the operation, and two patients (3%) had a small bowel obstruction. No fever or inflammation was reported after surgery. Overall, patients were discharged 1.5 ± 1.7 days after the operation. In the tack staple group complications occurred in 26.4% of the patients. Ten patients (9.8%) had a seroma and eight (7.8%) had a hematoma. No major complications or deaths were reported. The length of stay was a mean of 2.3 days (1.9 days, when the additional procedures were excluded). During the follow-up, three patients (2.9%) developed a recurrence at a mean of 16.3 months after surgery. It was observed that 15 patients (14.7%) complained of pain in the groin area more than three months after surgery. There were slightly more seromas but less hematomas in the fibrin glue group compared to the tack staples group, but this was not statistically significant. However, the postoperative chronic pain rate was significantly reduced in the fibrin glue group.

In an animal study in which TEP groin hernia repairs were performed, Katkhouda et al demonstrated2,4 that graft motion and tensile strength were similar in the staples and fibrin glue groups. In addition, histological examinations revealed that the fibrin glue triggered a stronger fibrous reaction and inflammatory response with more fibroblastic mesh ingrowth. The procedures were performed laparoscopically in 49 sites. Eighteen grafts were fixed with fibrin glue and 16 with staples; 15 were not fixed. There was no significant difference in graft motion between the fibrin glue and stapled groups; there was no significant difference in median tensile strength between the fibrin glue and stapled groups. Fibrin glue triggered a significantly stronger fibrous reaction and inflammatory response than in the stapled and control groups. No infection related to method of fixation was observed in any group. The experiment did not involve serial examinations at later dates because fibroblastic ingrowth has already fixed the mesh in position by postoperative day 12, after which further mesh migration or folding is unlikely. It was found out that in addition to its adhesive property, fibrin glue acts as scaffolding for fibroblastic ingrowth that is enhanced by the chemotactic action of its thrombin component.

Hung Lau et al studied in his randomized prospective study a total of 93 patients with 186 inguinal hernias, who underwent bilateral TEP and were randomized to have mesh fixation by either FS (n = 46) or mechanical stapling (n = 47). The FS group consumed significantly less analgesics compared with that of staple group. There was no significant difference in postoperative pain score at rest and on coughing from the day of operation to postoperative day 6 between the groups.

Stefano Olmi et al10 in his randomized prospective study termed “Quantification of pain in laparoscopic trans-abdominal preperi toneal (TAPP)” involving 600 patients, came to the conclusion that postoperative pain ranged from mild pain between 12 and 72 hours with Tisseel and it was higher in other methods of mesh fixation: Moderate pain with EMS (Ethicon Endo surgery, Inc.) to severe pain with Protak at 48 hours follow-up. Significant differences in length of stay occurred, no recurrence or conversion rates were observed among groups, and morbidity was generally lower with Tisseel. Patients using Tisseel also returned to work sooner than as in other mesh fixing devices. Prostheses were fixed with Protak (Tyco, Norwalk, Conn), Endoanchor (Ethicon Endosurgery, Inc., Cincinnati, Ohio), EMS (Ethicon Endosurgery, Inc.) and Tisseel (Baxter Healthcare, Milan, Italy).

RH Fortelny and R Schwab et al15 assessed the quality of life in a trial with mesh fixation with fibrin glue in TAPP. TAPP with fibrin mesh sealing was performed in 11 non-selected consecutive patients. A direct control group (e.g. TAPP with staples) was not considered as favorable change in the quality of life in patients with fibrin glue mesh fixation was the tested hypothesis and not the comparison of techniques. Quality of life and pain were assessed preoperatively and one year follow-up using the SF36 survey and the visual analogue score (VAS). Post one-year analysis of recurrences or complications was made. The analysis of the unmodified SF-36 revealed a highly significant improvement. The scale ‘social functioning’ (SOCIAL), which belongs to the mental-health-related scale, had also significantly improved. The VAS significantly reduced after one year.

In an observational prospective multicenter study involving 1,201 patients performed in France B Descottes et al16 assessed Tisseel® fibrin glue for atraumatic mesh fixation in inguinal hernia repair. Out of 1,201 patients, 526 procedures were performed using open techniques and 675 using laparoscopic repairs. Local complications occurred in 4.7% of patients: 3.0% hematoma, 1.4% seroma, 0.3%
DISCUSSIONS

During the past few years, attention has focused on the pain that may arise after groin hernia surgery. Chronic pain after hernia surgery is a complex and controversial problem that affects not only open but also laparoscopic procedures. Three pain syndromes have been identified: Somatic, neuropathic, and visceral pain. Besides nerve damage during dissection, thermal injury due to electrocautery, and inflammatory and/or mechanical reaction to the mesh, stapling of the mesh is the most frequent evoked mechanism. According to various literature sources, there is a great variation in the rate of postoperative chronic pain, ranging from 0.1 to 0.4% and 22.5% in laparoscopic repairs for which staples are used to attach the mesh. Among the explanations for such a wide discrepancy are the range of pain evaluation methods used, which include clinical examination of the patients, phone calls, and mailed questionnaires and tools to score the severity of the pain.

For laparoscopic hernia repair, the possibility of nerve injury (pain or paraesthesia) caused by entrapment from incorrect placement of staples (above all lateral cutaneous femoral nerve, but ilioinguinal, and genitofemoral are also at risk) and epigastric vessels lesion by clips application may be avoided using fibrin glue either in the TAPP technique or in the TEP. It seems that not only entrapment but also postoperative fibrous scar around the staples can lead to nerve injury. In conventional TAPP, the prosthesis is anchored using metal clips. This is a critical step that requires the utmost attention to avoid damaging the surrounding nerves and blood vessels. Lesions arising due to such intraoperative damage can lead to complications, including hemorrhage, or painful neuralgia during follow-up. The nerves in the inguinal area that are most frequently involved in postoperative pain following TAPP hernia repair include the genitofemoral, lateral cutaneous femoral, and ilioinguinal and iliohypogastric nerves. Lesions of the lateral cutaneous femoral nerve are the most frequent postoperative neurologic complications associated with laparoscopy. Such lesions are the result of damage or entrapment of the nerve during lateral fixation of the mesh to the deep inguinal ring with pain in the lateral region of thigh. TAPP hernia repair with Tisseel resulted in a low rate of postoperative pain and rapid resumption of normal activities. Postoperative complications affected only 2.2% of the hernias and were readily treated without the need to extend the hospital stay. Importantly, no recurrences have been observed. These statistics are in favor of fibrin glue mesh fixation, along with other studies, which report postoperative complication rates of 4.6% for hematomas, 2% for neuralgias, and 0.4% for chronic pain. Reported persistent neuralgia with inguinal pain attributable to stapling vary in the literature from 0.5 to 14%.

To date, the series reporting the lowest postoperative chronic pain rates have not used any means of mesh. Tamme et al19 observed 2.55 and 0% chronic pain problems respectively, after TEP repair with a recurrence rate of less than 0.6%. However, the largest of these two series did not specify the length of follow-up and the other one was a rather small series (n = 89). Although, two randomized studies with a short follow-up of nonfixed mesh in laparoscopic repairs (one in TEP and the other in TAPP) did show promising results in terms of recurrence justification for routine nonstapling of the mesh in TEP is not yet substantiated.

Studies points to the fact that mesh stapling does play a key role in generating postoperative pain after laparoscopic hernia repair. Mesh fixation with fibrin glue is preferable as it meets the requirements for both efficiency and security of fixation.

The recurrence rate in the fibrin glue was found to be slightly lower than in the tack staples group but did not differ significantly, and the case of recurrence reported in the fibrin glue group is probably related to an inadequate mesh size in a large direct hernia. Overall, the recurrence rate in the fibrin glue group remains within the range most report approve. Inadequate lateral fixation is the main cause of recurrence after both TAPP (36%) and TEP (22%). The reason for this is that most of the nerves run laterally, where no staples can be applied. Gluing a large mesh on the triangles of disaster and pain is likely to stop the prosthesis from lifting and dislocating, thereby avoiding inferomedial and inferolateral recurrence. It is not known if the enhanced inflammatory response induced by fibrin glue may explain the slightly higher rate of seromas in the fibrin glue mesh fixation. There exists no significant difference in the development of postoperative hematomas, even though the rate is slightly lower in fibrin glue mesh fixation, as compared to the tack staplers but the data available are inconclusive to give the credit to the effect of fibrin glue on local hemostasis. Although, no comparison is available between the tack staples group and the fibrin glue group in terms of operation duration, the use of fibrin glue and its application device did not seem to change the mean operative time, which is comparable to that of other series using tackers. This can be attributed primarily to the peritoneum closure using a running laparoscopic suture and the preparation of fibrin glue and its applicator during the hernia...
sac dissection. From a mechanical standpoint, fixation of the mesh was equivalent to that obtained with clips but prevented complications related to the application of staples (bleeding and hematomas in Retzius’ space, neuralgia and chronic pain).

The difference in terms of operating costs between the two fixation techniques/methods does tilt in favor of fibrin glue. Two milliliters of Tisseel is available for 149 USD, whereas the single use tacker stapler is 287 USD, 300 euros for Endoanchor (Ethicon Endosurgery) and 250 Euros for Protak (Tyco, Norwalk, CT, USA). On long-term prospective, considerable cost savings can be done if postoperative complications (neuralgias, seromas, and hematomas) are reduced and hospital recovery periods are shortened.

CONCLUSIONS

The use of fibrin glue has a distinct advantage in laparoscopic treatment of inguinal hernias compared with other conventional methods of mesh fixation. The use of fibrin sealant reduces the risk of post- and intraoperative complications, such as bleeding, seroma, chronic pain, has a lower incidence of postoperative neuralgia and provides an early faster return to social life. The recurrence rates are similar, but the operative time is slightly longer if the preparation time of the fibrin sealant is taken into consideration. Otherwise, the operative time is shorter in fibrin mesh fixation as compared to staples/tacks. Fibrin glue appears to be an effective alternative to staples, tacks and anchors for mesh fixation. Mesh fixation with fibrin glue is preferable as it meets the requirements for both efficiency and security of fixation.

REFERENCES