Microsurgical Ligation for Painful Varicocele: Effectiveness and Predictors of Pain Resolution

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Purpose: We evaluated the effectiveness of microsurgical ligation for painful varicocele and predictive factors of pain resolution. Materials and Methods: Between January 2006 and March 2009, a total of 114 patients (mean age, 30.2±8.9 years), who underwent microsurgical inguinal varicocelectomy for painful varicocele, were included and followed up for 1 year after the surgery. The quantity of preoperative and postoperative pain was assessed by means of 11-point numeric rating scale (NRS). We retrospectively analyzed the outcome of surgical ligation and predictive factors of pain resolution using patient age, height, weight, body mass index, grade and location of varicocele, duration, quantity and quality (dull, dragging, aching) of pain, and postoperative pain resolution.

Results: In 104 patients (91.2%), complete or marked resolution of pain was reported at follow-up 1 year after surgery. Only 10 patients (8.8%) had recurrent or persistent pain (≥3 points in NRS scores). On multivariate analysis, low quantity (≤6 points in NRS scores) and dull or dragging natured preoperative pain were independent factors associated with surgical success rates (p=0.004; odds ratio=1.62, p=0.012; odds ratio=1.76, respectively).

Conclusion: Microsurgical ligation is an effective treatment of painful varicocele. The quantity and quality of preoperative pain are independent predictive factors of pain resolution after surgery.

Key Words: Varicocele, ligation, pain

INTRODUCTION

Varicocele is the most commonly diagnosed peripubertal andrologic disease and the most commonly treatable cause of male-related impaired fertility potential. Also, varicocele is associated with chronic scrotal pain, but this relationship has been less widely investigated. It is evaluated that the prevalence of painful varicoceles is 2-10%. This pain was described as a dull, throbbing pain, worsening with exertion and strain.

Varicocele ligation has been assessed as an effective treatment of painful varicocele. Other comparative reports have assessed different techniques, but at present there is no consensus as to which surgical method is best suited for varicocele treatment, although microsurgical techniques of varicocelectomy have gained popularity with minimal complication rates and good outcomes. Also, little data ex-
ists regarding the outcome of surgical treatment of painful varicoceles and factors able to predict success. This retrospective study, in a selected homogeneous population evaluated the effectiveness of microsurgical ligation against painful varicocele, including the predictive factors affecting pain resolution.

MATERIALS AND METHODS

Patients
Between January 2006 and March 2009, a total of 114 patients (mean age, 30.2±8.9 years) who underwent microsurgical inguinal varicocelectomy for pain were included and followed up for at least 1 year (mean; 15.8 months: range, 13-49) after surgery. Written consent from each patient was obtained. Diagnosis of varicocele was made using self-diagnosed symptoms, such as scrotal discomfort or pain, and on clinical grounds of both physical examination and Doppler ultrasound scan (USS). Patients whose cause of scrotal pain was ambiguous and who had a previous history of scrotal trauma or surgery were excluded from this study. Men with lower urinary tract symptoms were also excluded, as were men with a documented psychological disturbance.

Preoperative evaluations
Medical records were retrospectively investigated to document patient demographics [age, height, weight, body mass index (BMI), etc.], grade (I, II, III) and, location of varicocele (right, left or bilateral), duration (months) and quality (dull, dragging, aching) of pain before surgery. To evaluate the quantity of pain, we used a 11-point numeric rating scale (NRS) where 0 indicated no pain and 10 indicated the worst pain imaginable (0, 1-3, 4-6, 7-10: none, mild, moderate, severe; respectively).

Procedure
The microsurgical inguinal varicocelectomy was performed under general or spinal anesthesia by a single experienced surgeon using a similar technique previously described, except for we did not deliver the testicle or examine the gubernaculum. An operative microscope (10-25X) was used during spermatic cord dissection to identify and preserve the testicular artery and lymphatics.

Follow-up evaluation
All patients were asked to visit the clinic at 1 month, 3 months and 1 year after surgery, at which time they were evaluated with a careful symptom review, scrotal examination at supine and standing positions, and 11-point NRS to assess the quantity of postoperative pain. Also, Doppler USS was performed when deemed clinically necessary for discovery of varicocele recurrence.

Definition
Varicocele was classified into the following grades: grade I, varicocele palpable during a Valsalva maneuver when upright; grade II, varicocele palpable when upright without Valsalva maneuver; and grade III, varicocele palpable and visually detectable. Surgical success was defined as the lack of palpable scrotal veins on physical examination and reporting of scores in the lowest range on the 11-point NRS scale (≤2 points) after surgery.

Analysis
Our primary objective was to analyze overall surgical success rates after microsurgical inguinal varicocelectomy. Our secondary objective was to determine which preoperative factors influenced the pain resolution after surgery by evaluation clinical factors. Normally distributed variables were compared using ANOVA and Student’s t-tests. The Fisher’s exact and chi-square tests were used for dichotomous variables. To determine the predictive factors affecting the resolution of pain, univariate analysis was performed using a logistic regression analysis. A 5% level of significance was used for all statistical testing and all statistical tests were two-sided. Analyse was performed using the statistical software SPSS (14.0KO for Windows, Release 14.0.2).

RESULTS
One hundred and fourteen patients with painful varicocele underwent microsurgical inguinal varicocelectomy with complete follow-up, and all patients who underwent surgery failed an initial conservative treatment (scrotal support, non-steroidal anti-inflammatory treatment and exercise limitation) for 4-24 weeks. Baseline characteristics of the 114 men are illustrated in Table 1.

Of the 114 patients, 104 (91.2%) reported complete or marked resolution of pain at follow-up 1 year after surgery. Only 10 (8.8%) had recurrent or persistent pain (≥3 points in 11-point NRS scores).

Four patients (3.5%) had recurrence of pain (mean points...
in 11-point NRS scores, 4.0±0.8 points) after an initial period of resolution (mean period, 1.8±0.5 months). All 4 patients with recurrent pain were subjected to Doppler USS and no patient had varicocele recurrence.

Six patients (5.3%) reported persistent pain (mean points in 11-point NRS scores, 4.7±1.4 points) at follow-up 1 year after surgery. Of those, one complained that his symptoms were worse. All 6 patients with persistent pain were also subjected to Doppler USS, and only one (0.8%) had varicocele recurrence. However, although successful radiographic embolization was performed in this patient, his pain persisted.

Table 2 shows the preoperative characteristics in the failure group of pain resolution. From the viewpoint of pain quality, all patients with recurrent or persistent pain had dragging or aching preoperative pain at follow-up 1 year after surgery.

Univariate analysis of the preoperative factors demonstrated that the quality and quantity of preoperative pain were associated with surgical success rates (Table 3). When we divided the patients into three groups according to preoperative pain qualities, we found that the success rates were 78.4% in men with aching pain, 96.2% with dragging pain and 100% with dull pain (p=0.003, OR 1.91). There was also a significant difference in success rates among patients’ groups according to preoperative NRS scores (1-3, 4-6, 7-10: mild, moderate, severe group) (p=0.002, OR 2.24).

Multivariate analysis indicated that quality and quantity of preoperative pain were independent factors related to success rates (Table 3). However, when the relationships between pain resolution and patient age, height, weight, BMI, duration of pain before surgery, grade and location of varicocele were examined, none of these factors significantly predicted the resolution of pain (p>0.05).

There were no intraoperative or early postoperative complications, such as orchitis, infection, or scrotal hematoma. At 1 year follow-up, as aforementioned, 10 patients had recurrent or persistent scrotal pain and 1 had varicocele recurrence; but no patients had hydrocele formation, evidence of testicular loss or progressive hypotrophy.

### Table 1. Characteristics of 114 Patients

| Characteristic                        | Value          |
|--------------------------------------|----------------|
| Mean age, yrs (range)                | 30.2 (13-52)   |
| Mean body mass index, kg/m² (range)  | 22.2 (17.1-26.6) |
| No. with varicocele location (%)     |                |
| Left                                 | 109 (95.6)     |
| Right                                | 0 (0.0)        |
| Both                                 | 5 (4.4)        |
| Mean pain duration before surgery, months (range) | 13.7 (2-68) |
| No. with varicocele grade (%)        |                |
| I                                    | 8 (7.0)        |
| II                                   | 36 (31.6)      |
| III                                  | 70 (61.4)      |
| Mean preoperative NRS scale (range)  | 6.1 (3-10)     |
| No. with preoperative NRS scale (%)  |                |
| Mild (1-3)                           | 3 (2.6)        |
| Moderate (4-6)                       | 70 (61.4)      |
| Severe (7-10)                        | 41 (36.0)      |
| No. with preoperative pain quality (%) |            |
| Dull nature pain                     | 24 (21.0)      |
| Dragging nature pain                 | 53 (46.5)      |
| Aching nature pain                   | 37 (32.5)      |
| Mean operation time, minutes (range) | 76.2 (50-125) |
| Mean hospitalization time, days (range) | 1.8 (1.0-4.0) |
| No. with anesthesia (%)              |                |
| Spinal                               | 74 (64.9)      |
| General                              | 40 (35.1)      |

NRS, numeric rating scale.

### Table 2. Preoperative Characteristics in the Failure Group of Pain Resolution (n=10)

| Characteristic                        | Value          |
|--------------------------------------|----------------|
| Mean age, yrs (range)                | 21.5 (13-29)   |
| Mean body mass index, kg/m² (range)  | 22.5 (18.9-26.6) |
| No. with varicocele location (%)     |                |
| Left                                 | 8 (80.0)       |
| Right                                | 0 (0.0)        |
| Both                                 | 2 (20.0)       |
| Mean pain duration before surgery, months (range) | 4.2 (2-7) |
| No. with varicocele grade (%)        |                |
| I                                    | 1 (10.0)       |
| II                                   | 4 (40.0)       |
| III                                  | 5 (50.0)       |
| Mean preoperative NRS scale (range)  | 9.0 (7-10)     |
| No. with preoperative NRS scale (%)  |                |
| Mild (1-3)                           | 0 (0.0)        |
| Moderate (4-6)                       | 0 (0.0)        |
| Severe (7-10)                        | 10 (100.0)     |
| No. with preoperative pain quality (%) |            |
| Dull nature pain                     | 0 (0.0%)       |
| Dragging nature pain                 | 2 (20.0%)      |
| Aching nature pain                   | 8 (80.0%)      |
| Mean operation time, minutes (range) | 80.5 (70-100) |
| Mean hospitalization duration, days (range) | 2.1 (1.0-4.0) |
| No. with anesthesia (%)              |                |
| Spinal                               | 7 (70.0)       |
| General                              | 3 (30.0)       |

NRS, numeric rating scale.
Surgical treatment of painful varicocele is controversial, because a few literature supports its use and surgical ligation is only recommended in men who have specific pain complaints and in whom conservative treatments has failed.4-6 The studies by Peterson, et al.11 and by Yaman, et al.3 reported complete resolution of pain in 86% and 88% of patients, respectively. Yaman, et al.3 suggested that the failure rate was associated with preoperative varicocele grade. Also, Karademir, et al.1 reported similar results using inguinal and subinguinal ligation and suggested that surgical approaches may influence outcomes with best outcomes achieved with inguinal or sub-inguinal ligation. However, Al-Buheissi, et al.12 showed that although surgical ligation is an effective

DISCUSSION

The most common grievance in men with varicocele is a dull and throbbing scrotal pain that aggravated during straining and exercise.1 Urologists often meet patients with painful varicocele, whose jobs need working mostly in standing positions or vigorous physical effort. However, most urologists cannot form an objective opinion based on their own experience.

Conventional treatment of painful varicocele is conservative treatment followed by surgery. Generally used conservative methods are scrotal elevation, nonsteroidal anti-inflammatory medication and decreased physical activity, that often lead to unacceptable lifestyle restrictions.10 Also, percutaneous embolization or various surgical options, such as high inguinal, inguinal, subinguinal, laparoscopic, and microsurgical ligation, might be implemented to relieve pain of varicoceles.2

Surgical treatment of painful varicocele is controversial, because a few literature supports its use and surgical ligation is only recommended in men who have specific pain complaints and in whom conservative treatments has failed.4-6 The studies by Peterson, et al.11 and by Yaman, et al.3 reported complete resolution of pain in 86% and 88% of patients, respectively. Yaman, et al.3 suggested that the failure rate was associated with preoperative varicocele grade. Also, Karademir, et al.1 reported similar results using inguinal and subinguinal ligation and suggested that surgical approaches may influence outcomes with best outcomes achieved with inguinal or sub-inguinal ligation. However, Al-Buheissi, et al.12 showed that although surgical ligation is an effective

## Table 3. Univariate Analysis of Preoperative and Intraoperative Parameters of Determinants for Surgical Success Rates at 1 Year Follow Up State

|                                | Odds ratio (95% CI) | p value |
|--------------------------------|---------------------|---------|
| Age                            | 1.25 (0.64-2.12)    | 0.472   |
| BMI                            | 1.06 (0.52-2.02)    | 0.934   |
| Left location                  | 1.00 (reference)    |         |
| Both location                  | 2.20 (1.17-4.86)    | 0.712   |
| Pain duration before surgery   | 0.87 (0.29-2.63)    | 0.806   |
| Varicocele                     |                     |         |
| Grade I                        | 1.00 (reference)    |         |
| Grade II-III                   | 2.42 (1.19-4.84)    | 0.681   |
| Preoperative NRS scale, mild (1-3) or moderate (4-6) group | 1.00 (reference)    |         |
| Preoperative NRS scale, severe (7-10) group | 2.24 (1.24-4.07)    | 0.002   |
| Preoperative pain quality, dull or dragging nature | 1.00 (reference)    |         |
| Preoperative pain quality, aching nature | 1.91 (1.13-3.24)    | 0.003   |
| Operation time                 | 0.76 (0.43-1.71)    | 0.542   |
| Hospitalization duration       | 0.65 (0.38-1.43)    | 0.514   |
| Anesthesia                     |                     |         |
| Spinal                         | 1.00 (reference)    |         |
| General                        | 1.06 (0.45-2.98)    | 0.932   |

95%CI, 95% confidence interval; BMI, body mass index; NRS, numeric rating scale.

**Fig 1.** Success rates by preoperative NRS scale and pain quality. The multivariate analysis of preoperative factors demonstrated that NRS scale and pain quality might predict the success rates of microsurgical ligation in painful varicocele patients. NRS, numeric rating scale.
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treatment of painful varicocele, there was no correlation be-
tween the surgical approach used and failure of pain resolu-
tion. They also reported that the quality of pain was a deter-
imental factor when selecting patients for varicocele ligation.

In our study, of the 114 patients, 104 (91.2%) reported
complete or marked resolution of pain at follow-up 1 year
after surgery. We also found that the quality and quantity
of preoperative pain were independent factors related to suc-
cess rates (p=0.012, OR 1.76 and p=0.004, OR 1.62, respec-
tively) in the present multivariate analysis. Accordingly, the
degree of postoperative pain resolution was related with low
quantity ≤6 points in visual analogue scale (VAS) scores] and
dragging or dull natured preoperative pain. However,
when the relationships between pain resolution and patient
age, BMI, duration of pain before surgery, grade and loca-
tion of varicocele were investigated, none of these factors
significantly predicted the resolution of pain (p>0.05).

With regard to varicocele recurrence as postoperative
complications, microsurgical varicocelectomy is better than
non-microsurgical methods.13,14 It has been reported that
varicocele recurrence generally results from incomplete li-
gation of collateral veins,2 and that magnification of the
spermatic vessels by operating microscope reduces the pos-
sibility of such complications.15 Levine, et al.16 also report-
ed that complete dissection of the spermatic cord using op-
erative microscope might be effective for chronic scrotal
pain because that procedure might result in partial denerva-
tion of the testis.

In this study, we used microsurgical inguinal varicocele-
tomy with complete dissection of the spermatic cord, such
that it only included the spermatic artery, lymphatics, and
the vas deferens and its accompanying artery. As a result,
10 patients (8.8%) had recurrent or persistent scrotal pain
and only one (0.8%) had varicocele recurrence at 1 year of
follow-up; however, no patients had hydrocele formation,
evidence of testicular loss or progressive hypotrophy.

In our study, the 11-point NRS was used to evaluate the
quantity of pain. Although the VAS and the NRS have been
reported that they provide almost identical results in the same
patient at several times after surgery, the NRS, with numbers
from 0 to 10 (‘no pain’ to ‘worst pain imaginable’), is more
practical and easier to understand than the VAS.17 Therefore,
in our study, response criteria is based on subjective patients
against other most studies.

In conclusion, microsurgical ligation has a high success
rate (91.2%) with low recurrence rate (0.8%) for the treat-
ment of painful varicocele. Also, we show that the quantity
and quality of preoperative pain are independent predictive
factors of pain resolution after surgery. A prospective ran-
domized study with long term follow-up comparing surgi-
tical treatments of painful varicocele using different surgical
approaches is needed to support the present data.

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