Effects of Electrical Muscle Stimulation for Preventing Deltoid Muscle Atrophy after Rotator Cuff Repair: Preliminary Results of a Prospective, Randomized, Single-blind Trial

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Background: This study investigates the effects of neuromuscular electrical stimulation (NMES) in preventing deltoid atrophy during the first 12 weeks after arthroscopic rotator cuff repair.

Methods: Eighteen patients undergoing arthroscopic repair of a medium-sized rotator cuff tear by a single surgeon, were randomized into two groups: NMES and transcutaneous electrical nerve stimulation (TENS). Each group used the respective device for 6 weeks after surgery. Pain was measured at baseline, 6, and 12 weeks postoperatively, using the visual analogue scale (VAS); range of motion (ROM), abduction strength and functional scores were measured at baseline and 12 weeks postoperatively. Deltoid thickness and cross-sectional areas were measured using magnetic resonance imaging at 12 weeks postoperatively.

Results: At 12 weeks post-surgery, no statistically significant difference was observed between the NMES and TENS groups in the pain VAS, the Disabilities of the Arm, Shoulder and Hand score, ROM, and abduction strength. Postoperative decrease in the thickness of the anterior, middle, and posterior deltoid, at the level just below the coracoid, was -2.5%, -0.7%, and -6.8%, respectively, in the NMES group, and -14.0%, -2.6%, and -8.2%, respectively, in the TENS group (\(p=0.016\), \(p=0.677\), and \(p=0.791\), respectively). At the level of the inferior glenoid tubercle, postoperative decrease in area of the deltoid was -5.4% in the NMES group and -14.0% in the TENS group, which was significantly different (\(p=0.045\)).

Conclusions: NMES has the potential for reducing deltoid atrophy after arthroscopic rotator cuff repair, suggesting that NMES might help minimize postoperative atrophy after various shoulder surgeries.

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Key Words: Shoulder; Rotator cuff; Deltoid muscle; Muscular atrophy; Electric stimulation

Introduction

An aging population and advancements in arthroscopic techniques have led to a steady increase in the number of repairs performed annually. There are approximately 300,000 cases of rotator cuff tear per year worldwide, making rotator cuff repair one of the most common orthopedic surgeries.\(^1\) Although rotator cuff repair generally improves pain and function, the complication rate is high.\(^2\) The most common complications are retear and stiffness, and several studies have also focused on atrophy of the shoulder muscles.\(^3-8\) Atrophy of the deltoid muscle, especially that of its anterior fibers, makes shoulder flexion difficult since no other muscle can effectively compensate for its functional loss.\(^9\) A previous study reported postoperative decrease in the thickness of the deltoid, with no recovery 1 year later.\(^7,8\) However, there is no report on the prevention of deltoid atrophy. Neuromuscular electrical stimulation (NMES) elicits skeletal muscle contractions through percutaneous electrodes that de-
polarize underlying motor nerves. NMES using percutaneous electrodes is noninvasive and easy-to-use, and has reportedly shown good results in preventing muscle weakness and decreasing the thickness of the quadriceps muscle after anterior cruciate ligament surgery.\(^{10-15}\) We therefore hypothesized that NMES could prevent deltoid muscle atrophy after arthroscopic rotator cuff repair, and performed a single-blind randomized study to compare the decrease in deltoid thickness in the NMES group and the transcutaneous electrical nerve stimulation (TENS) group (control).

**Methods**

**Patient Selection**

This investigation conformed to the principles outlined in the Declaration of Helsinki. All patients provided informed consent prior to experimentation. The study protocol was approved by the Medical Ethics Committee of our hospital. This study was a prospective, randomized, controlled trial. The inclusion criteria were medium-sized rotator cuff tears (tear size, 1–3 cm), planned arthroscopic repair surgery, and patient age between 40 and 75 years. Tear size was initially measured in preoperative magnetic resonance imaging (MRI), and confirmed arthroscopically using a calibrated probe at the time of surgery.

The exclusion criteria were incomplete repair of the footprint, preoperative fatty degeneration of the deltoid muscle, poor understanding of the stimulation devices (NMES or TENS), and refusal to participate in the study.

Sample size was determined by G power analysis (alpha error: 0.05, power: 0.80, calculated actual power: 0.82); a total of 22 patients were initially recruited as candidates for the study, 4 of whom were excluded due to preoperative fatty degeneration of the deltoid muscle (n=1) and poor understanding of the stimulation devices (n=3). Finally, 18 patients were enrolled between May 2018 and December 2018. Patients were assigned to either the NMES or TENS group, with a 1:1 allocation ratio based on the allocation sequence yielded by a computer-based block randomization process (block size, 4). Clinical and radiological evaluations were included in the follow-up at 12 weeks after surgery. The 3-month follow-up MRI was refused by 2 patients (NMES 1, TENS 1); thus, 8 patients in the NMES group and 8 in the TENS group were available for the final review (Fig. 1). Preoperative clinical data did not differ between the two groups (Table 1).

**Surgical Procedures**

Surgical procedures were performed by a single surgeon.

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### Table 1. Preoperative Data on Patients

| Variable                        | NMES group | TENS group | p-value |
|---------------------------------|------------|------------|---------|
| No. of patients                 | 8          | 8          |         |
| Male/female                     | 5/3        | 2/6        | 0.131   |
| Mean age (yr)                   | 62.0 (49–75) | 62.6 (52–73) | 0.878   |
| Dominant/nondominant shoulder   | 6/2        | 7/1        | 0.522   |
| Mean tear size (mm)             | 19.1 (11–29) | 21.4 (13–29) | 0.887   |
| ROM (degrees)                   |            |            |         |
| Forward Flexion                 | 154.4 ± 32.7 | 173.8 ± 9.2 | 0.145   |
| External Rotation               | 73.1 ± 25.5 | 86.9 ± 8.8  | 0.185   |
| Abduction                       | 143.8 ± 39.6 | 169.4 ± 19.7 | 0.171   |
| Pain VAS                        | 6.2 ± 0.4  | 6.1 ± 0.4  | 0.915   |
| DASH score                      | 40.7 ± 21.9 | 27.9 ± 14.5 | 0.274   |
| Peak torque, abduction (Nm)     | 9.2 ± 10.7 | 12.5 ± 14.0 | 0.615   |

Values are presented as number only, mean (range), and mean ± standard deviation.

NMES: neuromuscular electrical stimulation, TENS: transcutaneous electrical nerve stimulation, ROM: range of motion, VAS: visual analogue scale, DASH: Disabilities of the Arm, Shoulder and Hand.

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Fig. 1. Flow diagram: consolidated standards of reporting trials (CONSORT).

NMES: neuromuscular electrical stimulation, TENS: transcutaneous electrical nerve stimulation, MRI: magnetic resonance imaging.
Under general anesthesia, patients were placed in the beach-chair position. A posterior portal was first established for the evaluation of the glenohumeral joint. The associated biceps tendon lesion was treated by tenotomy (2 patients) in cases with subluxation or tear involving >50% of the long head biceps. The arthroscope was subsequently removed from the gleno-humeral joint and redirected into the subacromial space. The bursal tissue was removed for space clearance, and arthroscopic subacromial decompression was routinely performed to create a flat acromial undersurface in all patients. The footprint was prepared by removing the soft tissue and multiple channeling. The tear size was measured intraoperatively using a calibrated probe, after debridement of the degenerated tendon edges. All torn tendons were repaired using the suture bridge technique.

Postoperative Rehabilitation and Intervention
All patients underwent the same rehabilitation regimen. Briefly, shoulders were immobilized for 6 weeks with an abduction brace. Range of motion (ROM) exercises for the elbow, wrist, and fingers were started immediately after surgery. Pendulum exercise, passive forward elevation exercise, and shrugging of both shoulders were encouraged immediately after surgery. Active assisted ROM exercises were allowed after the patients were weaned off the brace. Muscle strengthening commenced at 12 weeks after surgery.

A portable EMS (EMS-1000; Cybermedic, Iksan, Korea) was used for the NMES intervention. This stimulator is practical for home use. During stimulation, the upper limb is secured by an abduction brace. Self-adherent, flexible rectangular electrodes are placed on the lower anterior and posterior portions of the deltoid muscle, and are marked to ensure consistent reapplication by the participant (Fig. 2).

NMES from the portable electrical stimulator was applied to the resting muscle, and the participant was instructed to relax during the induced muscle contraction. The intensity was set to the maximal intensity tolerated during each session, and participants were repeatedly encouraged to increase the intensity as tolerated. The stimulator was set to deliver a biphasic current, using a symmetrical waveform, at 35 Hz for 5 seconds (including a 2.5 seconds ramp-up time and a 2 seconds ramp-down time) and a 5 seconds off time with pulse duration of 250 µseconds.

A portable TENS (KD-PRO5000; Kukdong Electronics, Ansan, Korea) was used for the TENS group. Active TENS units delivered a balanced, asymmetric, biphasic waveform with alternating pulses between channels. Similar to prior studies, the TENS units used a continuous frequency of 5 to 56 Hz for 60 seconds with pulse duration of 150 µseconds. Patients were advised to adjust the intensity of the TENS unit to the highest tolerable setting that did not exacerbate their pain.

These interventions were initiated 24 hours after surgery, in patients assigned to the respective NMES and TENS groups. NMES or TENS were performed during each session, once daily, for 6 weeks after rotator cuff repair. An emphasis was placed on the importance of using the stimulator at a tolerable but slightly uncomfortable intensity, although there was no minimum intensity required for the study protocol. Moreover, the participants were repeatedly instructed to continue increasing to a tolerable intensity within and between sessions. Most participants demonstrated safe and proper use of the stimulator in the hospital. All patients visited the hospital 2 weeks after surgery, and we verified to confirm appropriate use as instructed.

Clinical Evaluation
The clinical assessment included ROM (forward flexion, external rotation, and abduction), pain visual analogue scale (VAS), deltoid muscle power (peak torque at abduction) and functional score (the Disabilities of the Arm, Shoulder and Hand [DASH] score, range: 0–100). These outcome measures were evaluated preoperatively, postoperatively at 6 weeks (pain VAS, DASH), and 12 weeks (pain VAS, ROM, DASH, and peak torque at abduction). The strength of shoulder abduction muscle was measured thrice by peak torque using Biodex (Biodex Corp., Shirley, NY).
NY, USA), for 5 seconds at 60 degrees of shoulder abduction.

**Radiological Evaluation**

Anatomic evaluation of the rotator cuff and the deltoid muscle was achieved by MRI, because this method provides the benefit of multiplanar imaging of the shoulder.\(^1\) All MRI studies were done with a 3.0-T unit (Achieva TX; Philips Medical System, Eindhoven, the Netherlands) and a dedicated shoulder array. Patients underwent imaging in the supine position, with the examined arm alongside the body in slight external shoulder rotation with the palm facing upward. Patients underwent MRI scans preoperatively and at 12 weeks postoperatively. Two orthopedic surgeons independently performed the MRI analysis in a randomized blind fashion. Measurements were performed on MRI images using the Picture Archiving and Communication System (PACS; Maroview ver. 5.4; MAROTECH Inc., Seoul, Korea).

The area of the deltoid muscle and the thickness of the anterior, lateral, and posterior deltoids were measured to compare the preoperative and postoperative deltoid volume, using the axial T2-weighted MRI images at the level just below the coracoid process (Fig. 3). Line 1 was drawn from the anterior edge to the posterior edge of the glenoid. Line 2 was drawn parallel to line 1 and passing the center of the humeral head, and the anterior and posterior deltoid thickness was measured along line 2 at the level just below the coracoid process. Line 3 was drawn perpendicular to line 1, and also passing through the center of the humeral head. Lateral thickness was measured along line 3. The cross-sectional area was measured on the same axial image (Fig. 3). The cross-sectional area of the deltoid muscle was measured at the level of the inferior glenoid tubercle using PACS (Fig. 4).

**Statistical Analysis**

A paired t-test was performed to assess differences in the preoperative and postoperative parameters of the deltoid muscle and the clinical outcomes of each group. An independent t-test

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**Fig. 3.** Measurement of the deltoid thickness and the area at the level just below the coracoid process. (A) The thickness of the anterior, lateral, and posterior deltoids was measured using the axial T2-weighted magnetic resonance images (MRI) at the level just below the coracoid process. Line 1 was drawn from the anterior edge to the posterior edge of the glenoid. Line 2 was drawn parallel to line 1 and passing the center of the humeral head, and the anterior and posterior deltoid thickness was measured along line 2 at the level just below the coracoid process. Line 3 was drawn perpendicular to line 1 and also passing through the center of the humeral head. Lateral thickness was measured along line 3. (B) Cross-sectional area was measured on the same axial image.

**Fig. 4.** Measurement of cross-sectional area of the deltoid thickness (at the level of inferior glenoid tubercle). (A) Cross-sectional area of deltoid was measured on axial T2-weighted magnetic resonance imaging (MRI) at the level of inferior glenoid tubercle. (B) Cross-sectional line a showed the level just below the coracoid, and line b showed the level of inferior glenoid tubercle.
was applied to compare postoperative alterations of the deltoid muscle and the clinical outcomes between the two groups. Statistical significance was set at \( p < 0.05 \) with 95% confidence intervals. All statistical tests were performed using IBM SPSS for Windows ver. 25.0 (IBM Corp., Armonk, NY, USA).

**Results**

No statistically significant differences were observed between the NMES and TENS group when comparing pain VAS, DASH score, ROM, and abduction strength (peak torque) at 12 weeks after surgery. MRI revealed three retears in the repaired rotator cuff at 12 weeks after surgery (Sugaya type IV & V; one retear in NMES group, two retears in TENS group) (Table 2).

Good interobserver reliability was obtained for measurement of thickness and area of the deltoid muscle on MRI (correlation coefficients 0.84 and 0.75, respectively).

The mean postoperative thickness of the anterior and posterior deltoid was significantly thinner than the mean preoperative thickness (\( p = 0.005 \) and \( p = 0.018 \), respectively). The mean postoperative thickness of the lateral deltoid was not significantly different from the preoperative thickness (\( p = 0.324 \)). The mean postoperative area of the deltoid was significantly smaller than the mean preoperative area at the level just below the coracoid process (\( p = 0.001 \)) and at the level of the inferior glenoid tubercle (\( p = 0.008 \); Table 3).

Postoperative decrease in the thickness of the anterior, middle, and posterior deltoid at the level just below the coracoid was -2.5%, -0.7%, and -6.8%, respectively, in the NMES group and -14.0%, -2.6%, and -8.2%, respectively, in the TENS group. Significant difference was obtained in the thickness of the anterior deltoid between the two groups (\( p = 0.016 \); Fig. 5). However, there was no significant difference in the thickness of the lateral and posterior deltoids between the groups (\( p = 0.677 \) and \( p = 0.791 \), respectively). The postoperative decrease of the deltoid area was -7.2% in the NMES group and -10.1% in the TENS group, which was not significantly different (\( p = 0.657 \); Fig. 6). However, at the level of the inferior glenoid tubercle, the postoperative decrease in area of the deltoid was -5.4% in the NMES group and -14.0% in the TENS group, which was significantly different (\( p = 0.045 \)).

Table 2. Comparison of Postoperative Results between NMES and TENS Groups

| Variable                  | NMES group | TENS group | \( p \)-value |
|---------------------------|------------|------------|---------------|
| At 6 weeks after surgery  |            |            |               |
| Pain VAS                  | 3.5 ± 2.7  | 4.1 ± 1.5  | 0.523         |
| DASH score                | 45.6 ± 22.0| 46.7 ± 22.0| 0.922         |
| At 12 weeks after surgery |            |            |               |
| Pain VAS                  | 2.6 ± 2.5  | 3.7 ± 2.1  | 0.379         |
| DASH score                | 29.3 ± 18.7| 30.3 ± 9.2 | 0.897         |
| ROM (degrees)             |            |            |               |
| Forward Flexion           | 149.4 ± 25.1| 156.4±8.5 | 0.493         |
| External Rotation         | 57.5 ± 18.3| 58.6±10.3 | 0.482         |
| Abduction                 | 150.0 ± 32.2| 159.3±11.0| 0.890         |
| Peak torque, abduction (Nm)| 13.3 ± 16.4| 9.9 ± 11.4 | 0.671         |
| Retear*                   | 1          | 2          | 0.522         |

Values are presented as mean ± standard deviation or number only.
NMES: neuromuscular electrical stimulation, TENS: transcutaneous electrical nerve stimulation, VAS: visual analogue scale, DASH: Disabilities of the Arm, Shoulder and Hand, ROM: range of motion.

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Fig. 5. At the level just below the coracoid, the postoperative decrease in the thickness of the anterior, middle, and posterior deltoid and -0.7%, and -6.8%, respectively, in the NMES group, and -14.0%, -2.6%, and -8.2%, respectively, in the TENS group. Significant difference was obtained in the thickness of the anterior deltoid between the two groups (\( p = 0.016 \); Fig. 5). However, there was no significant difference in the thickness of the lateral and posterior deltoids between the groups (\( p = 0.677 \) and \( p = 0.791 \), respectively). The postoperative decrease of the deltoid area was -7.2% in the NMES group and -10.1% in the TENS group, which was not significantly different (\( p = 0.657 \); Fig. 6). However, at the level of the inferior glenoid tubercle, the postoperative decrease in area of the deltoid was -5.4% in the NMES group and -14.0% in the TENS group, which was significantly different (\( p = 0.045 \)).

Table 3. Decrease of Deltoid Volume at 12 Weeks after Surgery in all Patients

| Variable        | Before surgery | 12 weeks after surgery | \( p \)-value |
|-----------------|----------------|------------------------|---------------|
| Anterior thickness (mm) | 16.0 ± 4.6 | 14.7 ± 4.6 | 0.005         |
| Lateral thickness (mm) | 9.6 ± 2.5 | 9.3 ± 2.0 | 0.324         |
| Posterior thickness (mm) | 20.8 ± 4.0 | 19.2 ± 4.3 | 0.018         |
| Deltoid area (mm\(^2\)) | 3,152.37 ± 715.67 | 2,904.89 ± 707.82 | 0.008         |

Values are presented as mean ± standard deviation.

*At the level below the coracoid. †At the level of inferior glenoid tubercle.
Results of this study reveal a significant decrease in anterior deltoid muscle thickness and deltoid area in the NMES group after rotator cuff repair as compared to the TENS group, suggesting that early postoperative application might be useful for reducing atrophy of the deltoid muscle. The TENS group was employed as a control group, intended to allow blinding of patients by providing sham stimulation (placebo stimulation).

Subsequent to rotator cuff repair, it is common to immobilize the shoulder for 6 weeks with an abduction brace, and commence muscle strengthening at 12 weeks. We deliberated that this rehabilitation protocol is likely to cause atrophy of the muscles around the shoulders, as has been indicated in few studies on deltoid atrophy after rotator cuff repair: Cho et al.\(^\text{7}\) reported a 7.3% decrease in the thickness of the anterior deltoid after open rotator cuff repair; Hata et al.\(^\text{8}\) reported that mean thickness of the anterior deltoid decreases by 11.4% (18.4 to 16.3 mm) at 6 months and 15.2% (18.4 to 15.6 mm) at 12 months postoperatively, and is significantly smaller than it was prior to surgery \(p<0.05\). In the present study, the mean thickness of the anterior deltoid decreased by 2.5% in the NMES group and by 14.0% in the TENS group at the level just below the coracoid, which was significantly different. At the level of the inferior glenoid tubercle, which occurred early in the study. Isometric evaluation of the deltoid muscle was performed at 6 and 12 weeks after surgery, and we believe that this early muscle strength evaluation had a negative effect. Hence, muscle strength evaluation was subsequently excluded at 6 weeks postoperatively, and there were no further cases of retear. The mean peak torque value of abduction strength was significantly not different between the groups in our study. This is probably due to the small sample size and short follow-up period.

In the current study, there were two retears in the TENS group and one in the NMES group after rotator cuff repair, all of which occurred early in the study. Isometric evaluation of the deltoid muscle was performed at 6 and 12 weeks after surgery, and we believe that this early muscle strength evaluation had a negative effect. Hence, muscle strength evaluation was subsequently excluded at 6 weeks postoperatively, and there were no further cases of retear. The mean peak torque value of abduction strength was significantly not different between the groups, but the mean value trended to increase (9.2 Nm to 13.3 Nm) in the NMES group, whereas values in the TENS group showed a tendency to decrease (12.5 Nm to 9.9 Nm). In a subsequent study, we intend to recruit additional patients and measure the isokinetic strength at 12, 24, 48 weeks after surgery. There was no significant difference in clinical outcomes at 12 weeks after surgery; however, Mahure et al.\(^\text{17}\) reported that the use of TENS

*Fig. 6. Postoperative decrease in cross-sectional area of the deltoid muscle in the neuromuscular electrical stimulation (NMES) group and transcutaneous electrical nerve stimulation (TENS) group. (A) At the level just below the coracoid, the postoperative decrease in area of the deltoid was -7.2% in the NMES group and -10.1% in the TENS group, which was significantly not different \(p=0.657\). (B) At the level of inferior glenoid tubercle, the postoperative decrease in cross-sectional area of the deltoid was -5.4% in the NMES group and -14.0% in the TENS group, which was significantly different \(p=0.045\).

Preop-area: preoperative area of deltoid, Postop-area: postoperative area of deltoid.

*Significantly different between two groups \(p<0.05\).*

Discussion

Results of this study reveal a significant decrease in anterior deltoid muscle thickness and deltoid area in the NMES group after rotator cuff repair as compared to the TENS group, suggesting that early postoperative application might be useful for reducing atrophy of the deltoid muscle. The TENS group was employed as a control group, intended to allow blinding of patients by providing sham stimulation (placebo stimulation).

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Preop-area: preoperative area of deltoid, Postop-area: postoperative area of deltoid.

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significantly reduces opioid use by >25% at both 48 hours and 1 week after arthroscopic rotator cuff repair, with statistically significant reductions in the pain scores during the same period.

The results of our study suggest that NMES could be useful in preventing anterior and inferior deltoid atrophy after various shoulder surgeries, such as replacement arthroplasty and fractures, and will have a positive effect on rehabilitation after shoulder surgery.

Our study has several strengths. First, all surgeries were performed by a single surgeon using the suture bridge technique, and all patients were subjected to the same rehabilitation protocol, performed by a single rehabilitation physician. We also performed radiological evaluations using the same MRI scan in the same position. Second, this study was a prospective and randomized controlled trial, and we employed placebo stimulation by TENS for single-blind trial. Third, this study was the first intervention study that prevented muscle atrophy after rotator cuff repair.

The present study also has a few limitations. First, this study had a 3-month follow-up period. We thought that deltoid atrophy was most likely to be prominent at 12 weeks after surgery, before the start of muscle strengthening; hence, postoperative MRI was performed at 12 weeks. However, comparison of clinical outcome seems to be insufficient after 12 weeks. In a follow-up study, we intend to compare clinical outcomes at 6 and 12 months after surgery, and the final muscle thickness will also be compared at 12 months through MRI or computed tomography. Second, this study had a limited sample size for comparing the deltoid area. The sample size obtained from G power analysis was 11 per group (alpha error: 0.05, power: 0.80, calculated actual power: 0.82, total sample size: 22). However, only 16 patients were finally included in the analysis, after 4 patients were excluded as per the selection criteria and 2 were lost to follow-up. Third, the frequency and duration of machine use were evaluated as per the report submitted by patients. Most patients believed that electrical stimulation was helpful for the treatment, and the stimulation devices were portable and easily applicable. Since none of the patients reported that the stimulation device was utilized less than once a day, we, therefore, assessed that stimulation was done properly.

Conclusion

Results from this prospective single-blind randomized trial demonstrate that compared with TENS, NMES is more efficient in reducing deltoid atrophy after arthroscopic rotator cuff repair, thereby suggesting that NMES has the potential to minimize postoperative atrophy after diverse shoulder surgeries.

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