Incidence of clinically evident isolated axillary nerve injury in 869 primary anatomic and reverse total shoulder arthroplasties without routine identification of the axillary nerve

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Background: It has been suggested that, during primary shoulder arthroplasty, surgeons should identify the axillary nerve through direct visualization, palpation, or the “tug test” to prevent iatrogenic nerve injury. Our goal was to document the rate of isolated axillary nerve injury (IANI) in patients who had undergone primary anatomic total shoulder arthroplasty (TSA) or reverse total shoulder arthroplasty (RTSA) without routine identification of the axillary nerve.

Methods: Data on 869 cases of primary shoulder arthroplasty (338 TSAs and 531 RTSAs) performed by 1 surgeon between 2003 and 2017 were reviewed. Neither the tug test nor identification of the axillary nerve through palpation or visualization was used in any case. The primary outcome was new IANI documented within 3 months after arthroplasty. The frequency of IANI was summarized using point estimates and 95% confidence intervals (CIs).

Results: Six cases met the criteria for IANI. The overall incidence of IANI was 0.7% (95% CI, 0.3%-1.4%). The incidence of IANI was 0.3% (95% CI, 0.0%-1.6%) after TSA and 0.9% (95% CI, 0.3%-2.1%) after RTSA. All IANIs were cases of neurapraxia, and all patients had experienced complete neurologic recovery at last follow-up.

Conclusion: Complete, permanent IANI resulting from direct surgical trauma during primary shoulder arthroplasty can be avoided without using the tug test or routine identification of the nerve. A low incidence of partial temporary IANI can be expected, which may be related to indirect traction injuries.

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subcoracoid space and over the axillary nerve. A gentle tug on the nerve with the subdeltoid index finger can be transmitted across the nerve and felt with the other index finger, allowing the surgeon to identify the nerve location and thereby protect it (Fig. 1). Although there is support for the use of the tug test, it has not been studied extensively or validated.\textsuperscript{26,27} Moreover, the risk of direct injury to the axillary nerve when performing primary shoulder arthroplasty without routine identification of the nerve is undetermined. A previous study of patients undergoing open capsular shifts for shoulder instability reported that neither isolating the nerve nor performing the tug test was necessary to protect the nerve from direct injury during the procedure.\textsuperscript{24}

The aim of this study was to document the rate and characteristics of clinically evident isolated axillary nerve injury (IANI) in patients who had undergone primary TSA or RTSA without axillary nerve identification through direct visualization, palpation of the nerve, or use of the tug test. We hypothesized that primary TSA and RTSA can be performed safely without isolating or palpating the axillary nerve or performing the tug test. The results of this study will help patients and surgeons understand the risks of clinically evident IANI after primary shoulder arthroplasty without routine identification of the axillary nerve.

**Materials and methods**

**Patient selection**

All patients aged 18 years or older who underwent elective shoulder arthroplasty at our institution performed by the senior author (E.G.M.) between January 1, 2003, and July 31, 2017, were identified using our shoulder arthroplasty database. We included patients who underwent primary shoulder arthroplasty (first elective TSA or RTSA). We excluded patients who underwent revision arthroplasty, which was defined as any procedure performed after failure of cup arthroplasty, hemiarthroplasty, TSA, or RTSA.

We identified 872 primary TSAs performed during the study period. Of these patients, 3 did not have at least 3 months of follow-up and were excluded, leaving 869 patients (99.7%) for analysis. Of these, 338 (39%) underwent primary TSA and 531 (61%) underwent primary RTSA. Our cohort consisted of 453 women and 416 men with a mean age (± standard deviation) of 67 ± 11 years.

**Surgical technique**

All patients underwent general anesthesia with or without a single-injection interscalene block or interscalene catheter. Patients underwent RTSA or TSA by a deltopectoral approach in a semi-sitting beach-chair position. The Solar Total Shoulder System (Stryker, Mahwah, NJ, USA) was used for all TSAs. RTSAs were performed using the Encore Reverse Shoulder Prosthesis (BJO Surgical, Austin, TX, USA) or the Reunion RTSA system (Stryker). When the subscapularis tendon was present, the interval between it and the conjoint tendon was developed. A knee retractor was placed between the 2 muscles to protect the brachial plexus. The subscapularis tendon and capsule were then released as a single unit from the lesser tuberosity and from the junction of the humeral head and the proximal humeral shaft. A blunt Hohmann retractor was placed along the axillary pouch to protect the axillary nerve while the capsule was released medially.

The rotator cuff interval was released, and the arm was then extended, adducted, and externally rotated to allow the humeral head to dislocate (Fig. 2). This arm position has been associated with nerve alerts related to brachial plexus traction.\textsuperscript{26,27} Although external rotation alone may be helpful during subscapularis and/or capsule peel off to draw the lesser tuberosity away from the axillary nerve,\textsuperscript{29} care was taken not to leave the arm in the extended, adducted, and externally rotated position for prolonged periods. Retractors were then placed circumferentially around the humeral head, with one placed inferiorly to protect the axillary nerve and one placed medially to protect the glenoid and brachial plexus. After the humeral head was removed, the glenoid was exposed. The labrum was resected circumferentially by releasing it from its capsular attachments. The capsule was released from the glenoid rim by use of electrocautery with care taken to stay directly on bone. We did not perform a 360° subscapularis release in any case.\textsuperscript{22,25} The described exposure allowed direct access to the glenoid without palpating or exposing the axillary nerve or using the tug test. Video 1 shows performance of this technique.

**Study outcome**

The primary outcome variable, IANI, was defined as the presence of an isolated new sensory or sensorimotor axillary nerve deficit documented in the medical record within 3 months of shoulder arthroplasty. The medical records of all included patients were reviewed in detail by 2 of the authors (C.L.L. and J.R.). Data on neurologic findings based on patient reports, physical examination findings, and electrodiagnostic studies were collected using a standard data collection form.

All physical examinations were performed by the senior author (E.G.M.) using the same examination protocol preoperatively and postoperatively. Preoperatively and postoperatively, the upper extremities were evaluated for sensation using light touch for the

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**Figure 1** Tug test (→). The index finger is directed posteriorly in the subdeltoid space while the other index finger is placed medially into the subcoracoid space and over the axillary nerve. A gentle tug on the nerve with the subdeltoid index finger can be transmitted across the nerve and felt with the other index finger, confirming that the nerve has been correctly identified. (Reproduced with permission from Chalmers PN, Van Thiel GS, Trenhaile SW. Surgical exposures of the shoulder. J Am Acad Orthop Surg 2016;24:250-8.)
axillary, ulnar, radial, and median nerves. Sensation for all nerves was compared meticulously side to side, and any differences preoperatively or postoperatively were noted. Motor testing of the axillary nerve consisted of resisted abduction of the deltoid muscle at 90° of abduction preoperatively and postoperatively; this was compared with the contralateral side. All patients were evaluated each day while hospitalized and at 8 to 10 days, 6 weeks, and 12 weeks postoperatively. Indications for electromyography (EMG) included signs of brachial plexopathy or individual nerve injury with neurologic deficits that did not improve during the first 6 weeks after arthroplasty.

All cases of IANI were followed until complete resolution of neurologic symptoms or the date of the last documented follow-up. Documentation of neurologic status was recorded within defined periods (<3, 3-6, 7-12, or >12 months) to assess the timing of neurologic recovery. The extent of neurologic recovery was defined clinically as complete (ie, neurologic status returned to baseline), partial (ie, symptoms improved but residual deficit remained), or none (ie, deficit was unchanged from initial description), consistent with criteria used in prior studies.

**Statistical analyses**

The incidence of IANI was estimated for the whole cohort and for the TSA and RTSA subgroups using point estimates with their corresponding 95% confidence intervals (CIs). CIs were calculated using the Jeffreys interval method considering the low rate of events. A two-tailed Z test was used to compare the incidence of IANI between primary TSA and RTSA. Continuous variables are reported as means and standard deviations, and categorical variables are reported as frequencies and percentages. P < .05 was considered statistically significant. All analyses were performed using Stata software (version 14; StataCorp, College Station, TX, USA).

**Results**

Six cases met our criteria for IANI, and all were neurapraxia lesions that resolved with no sequelae. Of the 6 IANIs, 5 occurred after RTSA and 1 occurred after TSA. The overall incidence of IANI for all patients was 0.7% (95% CI, 0.3%-1.4%). For TSA, the incidence of IANI was 0.3% (95% CI, 0%-1.6%). For RTSA, the incidence of IANI was 0.9% (95% CI, 0.3%-2.1%). Although RTSA was associated with a higher incidence of IANI than was TSA, this difference was not significant (P = .26).

The characteristics and clinical course of patients who experienced IANI are summarized in Table I. In the patient with an IANI after TSA, the deficit was both sensory and motor. Of the 5 patients with IANI after RTSA, 2 had both motor and sensory deficits whereas 3 had isolated sensory deficits. In the 3 patients with only sensory deficits, EMG was not performed and all deficits resolved within 6 months. In the 3 patients with sensory and motor changes (1 TSA and 2 RTSA patients), EMG confirmed partial IANI. All 3 patients experienced clinically complete recovery by 9 months. No patient had residual clinical sensory or motor changes at last follow-up.

**Discussion**

In our patients, the incidence of IANI after primary TSA or RTSA was less than 1% and all isolated injuries were cases of neurapraxia that resolved completely. These findings suggest that to prevent iatrogenic direct complete injury to the axillary nerve during primary shoulder arthroplasty, routine identification of the nerve by performing palpation, visualization, or the tug test is unnecessary if precautions are taken to protect the nerve. Despite not palpating the axillary nerve, directly observing the axillary nerve, or performing the tug test on the axillary nerve, neurovascular lesions were uncommon. This study supports the safety of not routinely identifying the axillary nerve during primary shoulder arthroplasty using the described surgical technique, but it does not show whether routine identification of the axillary nerve might reduce or prevent IANIs.

The incidence of IANI in our study (0.7%) is within the range of previously reported rates (0%-16%); Table II. Our reported incidence of IANI after TSA of 0.3% is also comparable to previously reported rates of 0.72% to 1.8%.

Although we found a higher incidence of IANI after RTSA (0.9%) than after TSA (0.3%), this difference was not significant. RTSA has been associated with a higher risk of nerve injury compared with TSA. Ladermann et al compared the incidence of nerve injury in TSA vs. RTSA using postoperative EMG and nerve conduction studies (NCS) and found that RTSA was associated with a 10.9-fold greater risk of postoperative nerve injury than TSA. They reported 10 isolated nerve injuries, 6 of which were in the axillary nerve and all of which were in the RTSA group. They speculated that the greater risk of any neurologic injury during RTSA compared with TSA was attributable to excessive arm lengthening.

We defined nerve injury as a clinically evident sensory and/or motor deficit consistent with an axillary nerve injury. The use of clinical detection to define the incidence of nerve injury may lead to underdetection of subclinical axillary nerve injury, but the use of clinical detection alone has precedence in the literature. Although electrodagnostic studies can often detect subclinical nerve injury, this was not a prospective study to determine subclinical axillary nerve dysfunction. Our results are consistent with what one would expect when performing an examination for nerve injury alone without routine EMG-NCS, and this practice
represents the usual standard of care for detecting nerve injury after primary shoulder arthroplasty. Intraoperative neuromonitoring and postoperative electrodiagnostic studies have suggested that there are more subclinical injuries to the brachial plexus and individual nerves than commonly appreciated.\textsuperscript{1,2,16,27} Nagda et al\textsuperscript{28} documented episodes of nerve dysfunction in up to 57% of patients undergoing intraoperative neuromonitoring during shoulder hemiarthroplasty and TSA. Of these episodes, 16.7% involved the axillary nerve alone. In most cases, nerve dysfunction alerts occurred during humeral and glenoid preparation and returned to baseline after the arm was returned to a neutral position. These authors found that the most common arm position creating stress on the nerves was extension and external rotation with abduction or adduction. In a study of 36 patients undergoing TSA or RTSA with intraoperative neuromonitoring, Parisien et al\textsuperscript{29} reported 203 nerve events in the 2 cohorts. Most nerve alerts in the brachial plexus occurred during humeral and glenoid preparation in both cohorts. The axillary nerve was the most frequently affected (27%) of all the peripheral nerves in both cohorts. Aleem et al\textsuperscript{1} used continuous intraoperative neuromonitoring in 282 patients who underwent TSA, RTSA, or hemiarthroplasty to determine whether nerve alerts were associated with postoperative peripheral nerve injury. The greatest frequency of nerve alerts was noted for the axillary nerve, but no axillary nerve injuries were detected clinically after surgery.

Two postoperative, clinically detectable peripheral nerve injuries occurred (0.7%), but both were radial nerve injuries. Lädermann et al\textsuperscript{16} documented subclinical electromyographic changes involving mainly the axillary nerve in 47% of patients who underwent routine EMG-NCS after RTSA. However, all were partial injuries that resolved completely in less than 6 months.

Other factors have been suggested to contribute to the incidence of nerve injury after TSA or RTSA. In an anatomic study, Lädermann et al\textsuperscript{17} suggested that the axillary nerve may also be at risk at the junction of the humeral head and humeral shaft in the posterior metaphyseal area. They recommended that care be taken when reaming the metaphysis to avoid posterior humeral cortical violation, particularly when having a low humeral cut and using a large reamer. It has been suggested that patients with decreased range of motion (<10\degree of passive external rotation with the arm at the side) or a history of open shoulder surgery may be at increased risk of nerve injuries. Similarly, patients with pre-existing peripheral neuropathy or cervical radiculopathy may be at risk of increased neurologic symptoms postoperatively.\textsuperscript{16} In our study, the small number of patients with IANI prohibited evaluation of these variables and their possible contributions to IANI.

To our knowledge, ours is the first study to report the incidence of clinically evident IANI after primary shoulder arthroplasty performed without routine identification of the axillary nerve. However, the study has limitations that should be considered when

### Table I
Characteristics and clinical course of axillary nerve injuries

| Case no. | Procedure | Sex | Age, yr | Diagnosis                  | Type of neurologic deficit | Time until neurologic deficit, wk | Subjective findings               | Objective findings                                      | EMG findings                      | Time until complete recovery, mo |
|----------|-----------|-----|---------|----------------------------|-----------------------------|-----------------------------------|-----------------------------------|---------------------------------------------------------|-----------------------------------|---------------------------------|
| 1        | TSA       | F   | 57      | Primary OA                | Sensorimotor                | 12                                | Weakness, numbness                | Weakness in elevation, drop-arm sign                    | Partial axillary neuropathy       | 9                               |
| 2        | RTSA      | M   | 63      | OA with bone loss         | Sensory                     | 11                                | Dysesthesia, numbness             | Decreased sensation, weakness in elevation, drop-arm sign | Not performed                    | 6                               |
| 3        | RTSA      | F   | 53      | IMRCT with pseudoparalysis| Sensorimotor                | 2                                 | Dysesthesia, numbness             | Dysesthesia, numbness                        | Partial axillary neuropathy        | 10                              |
| 4        | RTSA      | M   | 85      | CTA                       | Sensory                     | 2                                 | Numbness                          | Decreased sensation                                  | Not performed                    | 1                               |
| 5        | RTSA      | F   | 70      | RA                        | Sensorimotor                | 8                                 | Weakness                          | Decreased sensation                                  | Not performed                    | 6                               |
| 6        | RTSA      | M   | 59      | CTA                       | Sensory                     | 7                                 | None                              | Decreased sensation                                  | Not performed                    | 1                               |

EMG, electromyography; TSA, anatomic total shoulder arthroplasty; F, female; OA, osteoarthritis; RTSA, reverse total shoulder arthroplasty; M, male; IMRCT, irreparable massive rotator cuff tear; CTA, cuff tear arthroplasty; RA, rheumatoid arthritis.

\*Abnormal spontaneous activity of the deltoid with fast-firing voluntary motor units of increased duration and amplitude.

### Table II
Axillary nerve mononeuropathy reported in the literature after primary shoulder arthroplasty

| Author (year) | Procedure               | No. of shoulders | No. of axillary nerve injuries (%) | Diagnostic basis | Type of neurologic deficit | Recovery (time until recovery, mo) |
|---------------|-------------------------|------------------|-----------------------------------|------------------|---------------------------|-----------------------------------|
| Torchia et al\textsuperscript{1} (1997) | Primary TSA            | 113              | 1 (0.88)                          | NR               | NR                        | NR (NR)                           |
| Edwards et al\textsuperscript{10} (2002) | Primary HA and TSA     | 555              | 4 (0.72)                          | NR               | NR                        | Complete in 2 patients (NR), none in 2 patients (NA) |
| Goderich et al\textsuperscript{11} (2002) | Primary TSA            | 268              | 2 (0.75)                          | Clinical         | NR                        | None (NA)                         |
| Matsoukas et al\textsuperscript{13} (2003) | Primary HA and TSA     | 55               | 1 (1.8)                           | NR               | NR                        | Complete (NR)                     |
| Werner et al\textsuperscript{17} (2005) | Primary RTSA           | 58               | 1 (1.7)                           | NR               | NR                        | Complete (NR)                     |
| Boileau et al\textsuperscript{18} (2006) | Primary RTSA           | 45               | 1 (2.2)                           | NR               | NR                        | Partial (36)                      |
| Nagda et al\textsuperscript{1} (2007) | Primary HA and TSA, revision TSA | 30               | 3 (10)                            | EMG\textsuperscript{+} | NR                        | NR (NR)                           |
| Lädermann et al\textsuperscript{18} (2009) | Primary and revision RTSA | 199               | 1 (0.50)                          | NR               | NR                        | Complete (12)                     |
| Lädermann et al\textsuperscript{20} (2011) | Primary RTSA           | 23               | 0 (0)                             | NA               | NA                        | NA                                |
| Walch et al\textsuperscript{21} (2012) | Primary and revision RTSA | 19               | 3 (16)                            | EMG\textsuperscript{+} | Sensorimotor              | Complete (6)                      |

TSA, anatomic total shoulder arthroplasty; NR, not reported; HA, hemiarthroplasty; NA, not applicable; RTSA, reverse total shoulder arthroplasty; EMG, electromyography.

\* All patients underwent intraoperative neurologic monitoring. Patients with intraoperative nerve alerts underwent diagnostic EMG at least 4 weeks postoperatively.
\* All patients underwent EMG 3 weeks after surgery.
\* Patients did not undergo EMG routinely. EMG was performed after clinical diagnosis.
interpreting the results. First, the retrospective method of data collection introduces the possibility of missing transient yet clinically relevant events that may not have been reported by the patient or documented by the surgical team in the patient records. Second, without a control group in which the routine identification of the axillary nerve was performed, we cannot conclude that the presented technique is equivalent in, better at, or worse at preventing partial or complete IANI compared with routine performance of the tug test or palpation or visualization of the axial nerve. Third, our results are those of a high-volume, fellowship-trained shoulder and elbow surgeon who performs many primary and complex revision shoulder procedures.

The surgical technique described in this study may differ from the techniques used by other surgeons, who may resect the anterior capsule or release the subscapularis inferiorly to increase excursion of this muscle. Similarly, the presented surgical technique may differ from that required to expose the glenoid in patients with major scarring in the anterior-inferior glenoid. Consequently, our conclusions cannot be extrapolated to primary shoulder arthroplasty performed with a more extensive capsular release, such as that performed with the 360° subscapularis release, or to patients undergoing revision arthroplasty.

Other factors may have influenced our results. Prosthetic designs changed during the study period, which may have affected the axillary nerve position or tension. The retractors we used around the shoulder may not be the same as those used by other surgeons. During humeral and glenoid preparation, the arm was purposefully kept out of an abducted and externally rotated position, so it is possible that in the cohort studied, the final range of motion postoperatively, especially external rotation, might be less than that in other studies in which a 360° release was performed.

Conclusion

Our results suggest that when performing primary TSA or RTSA, the tug test or palpation or visualization of the axillary nerve may not be necessary to prevent complete, permanent IANI resulting from direct surgical trauma if proper precautions are taken to protect the nerve. When IANIs do occur, they are typically cases of neurapraxia that resolve within a year. This study is limited to 1 surgical technique for exposing the glenohumeral joint and should not be extrapolated to other techniques.

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Supplementary data

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