Surgical treatment of Neer type II and type V lateral clavicular fractures: comparison of hook plate versus superior plate with lateral extension: a retrospective cohort study

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Abstract
Purpose Different fixation methods are used for treatment of unstable lateral clavicle fractures (LCF). Definitive consensus and guidelines for the surgical fixation of LCF have not been established. The aim of this study was to compare patient-reported functional outcome after open reduction and internal fixation with the clavicle hook plate (CHP) and the superior clavicle plate with lateral extension (SCPLE).

Methods A dual-center retrospective cohort study was performed. All patients operatively treated for unstable Neer type II and type V LCF between 2011 and 2016, with the CHP (n = 23) or SCPLE (n = 53), were eligible for inclusion. The primary outcome was the QuickDASH score. Secondary outcomes were the numerical rating scale (NRS) pain score, complications, and implant removal.

Results A total of 67 patients (88%) were available for the final follow-up. There was a significant difference in bicortical lateral fragment size, 15 mm (± 4, range 6–21) in the CPH group compared to 20 mm (± 8, range 8–43) in the SCPLE group (p ≤ 0.001). There was no significant difference in median QuickDASH score (CHP; 0.00 [IQR 0.0–0.0], SCPLE; 0.00 [IQR 0.0–4.5]; p = 0.073) or other functional outcome scores (NRS at rest; p = 0.373, NRS during activity; p = 0.559). There was no significant difference in median QuickDASH score or other functional outcome scores between Neer type II and type V fractures. There was no significant difference in complication rate, CHP 11% and SCPLE 8% (relative risk 1.26; [95% CI 0.25–6.33; p = 0.777]). The implant removal rate was 100% in the CHP group compared to 42% in the SCPLE group (relative risk 2.40; [95% CI 1.72–3.35; p ≤ 0.001]).

Conclusion Both the CHP and SCPLE are effective fixation methods for the treatment of unstable LCF, resulting in excellent patient-reported functional outcome and similar complication rates. SCPLE fixation is an effective fixation method for the treatment of both Neer type II and type V LCF. The SCPLE has a lower implant removal rate. Therefore, if technically feasible, we recommend SCPLE fixation for the treatment of unstable LCF.

Keywords Lateral clavicle fracture · Unstable · Plate fixation · Clavicle hook plate · Superior clavicle plate with lateral extension
Introduction

The fracture of the clavicle is frequently encountered in the emergency department, accounting for 2.6–4% of fractures in the adult population. Furthermore, clavicle fractures represent 35–44% of fractures in the shoulder region. Although the majority involve the midshaft, lateral fractures account for 10–30% [1–6].

Lateral clavicle fractures (LCF) are classified according to Neer based on their relation to the coracoclavicular ligaments [6, 7]. Neer types I, III and IV are considered to be stable fractures and are generally treated conservatively. The unstable Neer type II and V fractures account for approximately 10–52% of LCF. Surgical management is recommended for these unstable LCF, as non-operative treatment results in a 22–50% non-union rate [1–6, 8, 9].

Neer type II fractures are unstable due to the detachment of the coracoclavicular ligaments from the medial fragment. Neer type V fractures have a comminuted character, with only an inferior fragment remaining attached to the coracoclavicular ligament [4, 6, 7].

Fixation of LCF proves to be a challenge as it can be difficult to get a firm hold on small lateral fragments. In addition, opposing forces contribute to considerable displacement of the fracture ends. Therefore, LCF can usually only be stabilized by rigid fixation methods [4, 9]. Different surgical fixation methods are available for the treatment of unstable LCF. However, at present, no consensus has been reached regarding the optimal fixation method.

The clavicle hook plate (CHP) is fixated with a small hook under the acromion posterior to the acromioclavicular joint. Complications related to the CHP such as acromial osteolysis, acromion fractures, rotator cuff tears and sub-acromial impingement have been reported [4, 5, 10, 11].

The superior clavicle plate with lateral extension (SCPLE) is a more recently developed locking compression plate. The SCPLE has multiple locking screws on the lateral end, divergently configured to maximize screw purchase on LCF fragments. The SCPLE does not interfere with the acromioclavicular joint and has a relatively low-profile [12–17]. Previous case series have shown the SCPLE to be an effective fixation method for the treatment of unstable Neer type II fractures [12–17]. However, the results after SCPLE fixation of Neer type V fractures have not yet been studied.

Currently, both the CHP and SCPLE are being used for the treatment of LCF. However, definitive consensus and guidelines for the surgical fixation of LCF have not yet been established. The aim of this study was to retrospectively evaluate patients treated with CHP and SCPLE fixation by comparing patient-reported functional outcome, complication-, and implant removal rates. Our hypothesis was that the SCPLE would result in better functional outcome and would lead to a reduction in complication- and implant removal rates.

Materials and methods

Study design

A retrospective cohort study was performed using data from two level II trauma centers. All patients with an unstable LCF who were treated operatively between January 2011 and June 2016 were eligible for inclusion. Inclusion criteria were: (1) acute LCF, (2) age 18 years or older, (3) Neer type II or type V fracture, (4) fixation with CHP or SCPLE, (5) fixation within 2 weeks of injury, and (6) minimum of one-year follow-up. Exclusion criteria were: (1) history of prior shoulder injuries or (2) neurovascular disorders of the affected shoulder. Data collection was performed by reviewing electronic medical records, operative reports, radiology reports, and telephone interviews by an independent research fellow. Electronic medical records were reviewed to collect baseline characteristics regarding affected shoulder, age, gender, trauma date, trauma mechanism, time from injury to surgery, fixation method, previous shoulder injuries, and lateral fragment size. Lateral fragment size was measured in millimeters (mm) on the anterior–posterior view radiograph. Overall lateral fragment size was defined as the total length of the largest lateral fragment. The largest intact bicortical fragment, which would allow for adequate screw fixation, was considered as the bicortical lateral fragment length. Informed consent was obtained from all subjects, and approval was granted by the institutional review board.

Surgical procedure

Patients were treated by means of open reduction and internal fixation (ORIF) using a CHP (3.5 mm LCP; Depuy Synthes GmbH, Oberdorf, Switzerland) or SCPLE (3.5/2.7 mm LCP; Depuy Synthes GmbH, Oberdorf, Switzerland). Implant selection was based on the surgeon’s preference. CHP and SCPLE fixation were performed by several surgeons in both trauma centers. Operations were performed under general anesthesia with the patient placed in a beach chair position. An incision was made using a standard superior approach. The fracture site was exposed preserving as much periosteum as possible. Reduction was performed under direct visualization, and fragments were temporarily fixated using K-wires or reduction forceps. Fracture reduction, implant position, and screw placement were checked under fluoroscopic guidance. Coracoclavicular ligament
repair was not routinely performed. Finally, the fascia and skin were closed in layers.

**Clavicle hook plate**

In cases of CHP fixation, a small incision was made in the posterior capsule of the acromioclavicular joint to allow subacromial hook placement. Trial plates were used to determine correct length and depth. Definitive CHP fixation was completed with the insertion of 3.5 mm angular stable or conventional screws (Fig. 1).

**Superior clavicle plate with lateral extension**

In cases of SCPLE fixation, there was no involvement of the acromioclavicular joint. A plate with an appropriate length was chosen to allow adequate fixation with 3.5 mm conventional or angular stable screws in the medial fragment and smaller 2.7 mm angular stable screws in the lateral end (Fig. 2).

**Postoperative management**

Both groups received the same postoperative management. Radiographs were taken 1 day postoperatively. Patients were temporarily immobilized in a sling until the pain subsided; early mobilization and active range of motion exercises were allowed when tolerated. Weight-bearing activities and resisted exercises were not permitted until approval from the treating surgeon. Follow-up visits were scheduled at 2, 4, and 12 weeks postoperatively. Additional outpatient visits were scheduled depending on fracture consolidation. Removal of the SCPLE was not routinely performed, as opposed to the CHP where removal was recommended to all patients.

**Primary outcome**

Functional outcome was assessed at least 12 months following ORIF, using the Dutch language version of the QuickDASH score. The QuickDASH is a validated and shortened version of the Disabilities of the Arm, Shoulder and Hand questionnaire (DASH). The QuickDASH is a patient-reported outcome instrument developed to measure upper extremity disability and symptoms, resulting in a score ranging from 0 (no disability) to 100 (most severe disability) [18, 19].

**Secondary outcome**

Secondary outcomes were the numerical rating scale (NRS) pain score at rest and during activity, complications, revision surgery and implant removal. The NRS is a reliable and commonly used 11-point scale to measure pain intensity, ranging from 0 (no pain) to 10 (worst imaginable pain) [20]. Complications included infection, non-union, malunion, implant failure, and implant removal-related complications. Infections were subdivided in superficial-skin or deep-wound infection. Superficial infection was defined as redness, swelling, or purulent discharge from the wound that was treated with antibiotics alone. If surgical irrigation and
debridement was required, it was considered a deep infection. Non-union was defined as the absence of fracture consolidation 6 months after surgery. Mal-union was defined as a symptomatic deformity of the clavicle. Implant failure was defined as implant displacement, implant breakage, or breakage of screws. Revision surgery was defined as the need for subsequent surgery other than implant removal. Infection and re-fracture following implant removal were considered implant removal-related complications. Implant-related irritation and indication for implant removal were analyzed using a series of questions developed by Hulsmans et al. [21]. Responses to these questions allowed categorization of implant removal into (1) routinely or on patient’s request without irritation or (2) patient’s request due to irritation. Patients with the implant still in situ received a different series of questions, leading to categorization of why implant was not removed; (1) not experiencing irritation, (2) experiencing irritation but removal not necessary, (3) experiencing irritation but no request for removal due to fear of re-operation, or (4) experiencing irritation, considering removal.

**Results**

**Study population**

A flowchart of the patient cohort is shown in Fig. 3. In total, 76 patients met the inclusion criteria. However, eight patients could not be contacted, and one patient refused participation. This resulted in the inclusion of 67 patients (88%) for analysis. The baseline characteristics are shown in Table 1. The CHP group included 19 patients (28%) compared to 48 patients (72%) in the SCPLE group. The most frequent fracture pattern was Neer type II found in 43 patients (64%). The overall lateral fragment size was 39 mm (± 12, range 14–83). There was a significant difference in bicortical lateral fragment size, 15 mm (± 4, range 6–21) in the CPH group compared to 20 mm (± 8, range 8–43) in the SCPLE group (p ≤ 0.001). The mean time from injury to surgery was 6.9 days (± 3.6, range 0–14). The mean follow-up was 37.5 months (± 17.9, range 12–76).

**Functional outcome**

There was no significant difference in functional outcome, as shown in Table 2. The median QuickDASH score in the CHP group was 0.00 (IQR; 0.0–0.0), as opposed to 0.00 (IQR; 0.0–4.5) in the SCPLE group (p = 0.073). There were 15 patients (79%) with a QuickDASH score of 0 in
### Table 1  Baseline characteristics

|                      | Overall n (%) | CHP n (%) | SCPLE n (%) | 95% CI of the difference | p value |
|----------------------|---------------|-----------|-------------|--------------------------|---------|
| Patients             | 67            | 19        | 48          |                          |         |
| Age [mean ± SD]      | 43 (14)       | 42 (17)   | 43 (12)     | −8.29 to 6.54            | 0.814   |
| Gender               |               |           |             |                          |         |
| Male                 | 54 (81)       | 13 (68)   | 41 (85)     |                          | 0.169   |
| Female               | 13 (19)       | 6 (32)    | 7 (15)      |                          |         |
| Side injury          |               |           |             |                          |         |
| Left                 | 39 (58)       | 8 (42)    | 31 (65)     |                          | 0.108   |
| Right                | 28 (42)       | 11 (58)   | 17 (35)     |                          |         |
| Affected side dominant side |         |           |             |                          |         |
| Yes                  | 27 (40)       | 10 (53)   | 17 (35)     |                          | 0.270   |
| No                   | 40 (60)       | 9 (47)    | 31 (65)     |                          |         |
| Neer classification  |               |           |             |                          |         |
| Type II              | 43 (64)       | 13 (68)   | 30 (63)     |                          | 0.780   |
| Type V               | 24 (36)       | 6 (32)    | 18 (38)     |                          |         |
| Overall lateral fragment (mm) [mean ± SD] | 39 (12) | 37 (12) | 40 (12) | −9.39 to 3.55 | 0.371 |
| Bicortical lateral fragment (mm) [mean ± SD] | 19 (7) | 15 (4) | 20 (8) | −8.40 to 2.64 | <0.001 |
| Time injury to surgery (days) [mean ± SD] | 6.9 (3.6) | 7.5 (3.5) | 6.7 (3.6) | −1.15 to 2.72 | 0.419 |
| Follow-up (months) [mean ± SD] | 37.5 (17.9) | 31.3 (16.3) | 40.0 (18.0) | −18.25 to 0.77 | 0.071 |

*Percentages may not add up to 100 due to rounding. Bold values indicate statistically significant results (e.g., p < 0.05)*

### Table 2  Functional outcome and implant-related complications

|                      | CHP (n = 19) n (%) | SCPLE (n = 48) n (%) | Relative risk (95% CI) | p value |
|----------------------|--------------------|----------------------|-------------------------|---------|
| QuickDASH median [IQR] | 0.00 (0.0–0.0) | 0.00 (0.0–4.5) | 0.073 |
| QuickDASH distribution [range] | 0–21 | 0–23 | | |
| 0                    | 15 (79)            | 25 (52)              |                         |         |
| 0–10                 | 3 (16)             | 19 (40)              |                         |         |
| 10–20                | 0                  | 3 (6)                |                         |         |
| 20–25                | 1 (5)              | 1 (2)                |                         |         |
| NRS pain at rest [median, IQR] | 0.00 (0.0–0.0) | 0.00 (0.0–0.0) | 0.373 |
| NRS pain at rest distribution [range] | 0–6 | 0–3 | | |
| 0                    | 16 (84)            | 44 (92)              |                         |         |
| 0–3                  | 2 (11)             | 3 (6)                |                         |         |
| 3–6                  | 1 (5)              | 1 (2)                |                         |         |
| NRS pain during activity [median, IQR] | 0.00 (0.0–1.0) | 0.00 (0.0–2.0) | 0.559 |
| NRS pain during activity distribution [range] | 0–8 | 0–7 | | |
| 0                    | 14 (74)            | 30 (63)              |                         |         |
| 0–3                  | 1 (5)              | 7 (15)               |                         |         |
| 3–6                  | 2 (11)             | 8 (17)               |                         |         |
| 6–8                  | 2 (11)             | 3 (6)                |                         |         |
| Complications        | 2 (11)             | 4 (8)                | 1.26 (0.25–6.33)        | 0.777   |
| Complication classification |     | | | 0.929 |
| Implant failure      | 1 (5)              | 3 (6)                |                         |         |
| Non-union            | 1 (5)              | 1 (2)                |                         |         |
| Revision surgery     | 1 (5)              | 2 (5)                | 1.26 (0.12–13.13)       | 1.000   |

*Percentages may not add up to 100 due to rounding. QuickDASH score: 0 = no disability to 100 = most severe disability. NRS pain score: 0 = no pain to 10 = worst imaginable pain. Bold values indicate statistically significant results (e.g., p < 0.05)*
the CHP group (range 0–21) compared to 25 patients (52%) in the SCPLE group (range 0–23). The median NRS pain score at rest was 0.00 (IQR; 0.0–0.0) in the CHP group and 0.00 (IQR; 0.0–0.0) in the SCPLE group (\( p = 0.373 \)). There were 16 patients (84%) with a NRS pain score at rest of 0 in the CHP group (range 0–6) compared to 44 patients (94%) in the SCPLE group (range 0–3). In the CHP group, the median NRS pain score during activity was 0.00 (IQR; 0.0–1.0) compared to 0.00 (IQR; 0.0–2.0) in the SCPLE group (\( p = 0.559 \)). There were 14 patients (74%) with a NRS pain score during activity of 0 in the CHP group (range 0–8) compared to 30 patients (63%) in the SCPLE group (range 0–7).

### Functional outcome according to Neer type

In both treatment groups, there was no significant difference in median QuickDASH score or other functional outcome scores between the Neer type II and type V fractures (Table 3). The median QuickDASH score in the Neer type II group following CHP fixation was 0.00 (IQR; 0.0–2.3), as opposed to 0.00 (IQR; 0.0–0.6) in the Neer type V group (\( p = 0.623 \)). In the SCPLE group, the median QuickDASH score in the Neer type II group was 0.00 (IQR; 0.0–5.1), as opposed to 2.30 (IQR; 0.0–4.5) in the Neer type V group (\( p = 0.764 \)).

### Implant removal

Implant removal rates and indications are presented in Table 4. CHP fixation was associated with a significant higher removal rate. CHP removal was, according to protocol, performed in all 19 patients (100%) compared to 20 patients (42%) in the SCPLE group (relative risk 2.40; 95% CI 1.72–3.35; \( p \leq 0.001 \)). The mean time to removal was 4.3 months (± 2.2, range 2–10) and 13.6 months (± 11.5,

### Table 3 Functional outcome according to Neer classification

| Neer            | Type II | Type V | \( p \) value |
|-----------------|---------|--------|---------------|
| CHP n (%)       | 13 (68) | 6 (32) | 0.623         |
| QuickDASH median [IQR] | 0.00 (0.0–2.3) | 0.00 (0.0–0.6) | 1.000 |
| NRS pain score at rest [median, IQR] | 0.00 (0.0–0.0) | 0.00 (0.0–0.0) | 0.734 |
| NRS pain score during activity [median, IQR] | 0.00 (0.0–2.0) | 0.00 (0.0–2.0) | 0.734 |
| SCPLE n (%)     | 30 (63) | 18 (38) | 0.609         |
| QuickDASH median [IQR] | 0.00 (0.0–5.1) | 2.30 (0.0–4.5) | 0.764 |
| NRS pain score at rest [median, IQR] | 0.00 (0.0–0.0) | 0.00 (0.0–0.0) | 0.609 |
| NRS pain score during activity [median, IQR] | 0.00 (0.0–3.3) | 0.00 (0.0–1.0) | 0.100 |

Table 3 Functional outcome according to Neer classification

### Table 4 Implant removal rate and indication

|                      | CHP (n = 19) n (%) | SCPLE (n = 48) n (%) | Mean difference (95% CI) | Relative risk (95% CI) | \( p \) value |
|----------------------|--------------------|----------------------|--------------------------|------------------------|--------------|
| Implant removal      | 19 (100)           | 20 (42)              | 2.40 (1.72–3.35)         | 2.40 (1.72–3.35)       | \( < 0.001 \) |
| Reason implant removed |                   |                      |                          |                        | 0.695        |
| Routinely or patient’s request, without irritation | 3 (16) | 5 (25) | 0.63 (0.17–2.29) |                       |              |
| Due to irritation    | 16 (84)            | 15 (75)              | 1.23 (0.81–1.55)         |                        |              |
| Time to implant removal (months) [mean±SD] | 4.3 (2.2) | 13.6 (11.5) | -9.287 (-14.757 to 3.817) |                        | \( 0.002 \) |
| Status implant not removed    |                    |                      |                          |                        | NP           |
| Not experiencing irritation | 0  | 12 (43)  |                      |                        |              |
| Irritation, but implant removal not necessary | 0 | 6 (21)  |                      |                        |              |
| Irritation, no request removal due to fear re-operation | 0 | 5 (18)  |                      |                        |              |
| Irritation, considering removal | 0 | 5 (18)  |                      |                        |              |

Table 4 Implant removal rate and indication

Bold values indicate statistically significant results (e.g., \( p < 0.05 \)). NP statistical analyses are not possible because all CHP implants were removed.
range 5–50) in the CHP and SCPLE groups, respectively (mean difference −9.287; 95% CI −14.757 to 3.817; \( p=0.002 \)). In the CHP group, three patients (16%) reported removal without irritation and 16 patients (84%) reported removal due to irritation. There were no cases of implant removal-related complications. In the SCPLE group, 28 patients (58%) did not have the implant removed and 12 patients (43%) reported not to experience irritation.

**Complications**

Complications were reported in two patients (11%) in the CHP group compared to four patients (8%) in the SCPLE group (relative risk 1.26; 95% CI 0.25–6.33; \( p=0.777 \)) (Table 2). Complications in the CHP group consisted of one case of implant failure due to implant displacement and one case of non-union. Complications in the SCPLE group included three cases of implant failure and one case of non-union. The implant failures in SCPLE group consisted of two implant displacements and one case of screw breakage. No cases of infection or mal-union were observed. In total, there were three patients that needed revision surgery. In the CHP group, one patient received a lateral clavicle resection due to non-union. Two revision surgeries were performed in the SCPLE group, one due to severe implant displacement and one case of non-union. The SCPLE implant displacement was treated by repeat SCPLE fixation. The non-union was treated with temporary K-wires fixation for 9.5 months.

**Discussion**

There was no significant difference in patient-reported functional outcome or complication rate between CHP and SCPLE fixation. However, the CHP was used more often on fractures with a small lateral bicortical fragment. There was no significant difference in patient-reported functional outcome between Neer type II and type V LCF fractures. Furthermore, there was a significant higher implant removal rate in the CHP group. In the SCPLE group, 57% of patients with the implant still in situ reported varying degrees of implant-related irritation.

Both the SCPLE and CHP result in excellent functional outcome. These findings are in accordance with previous comparative studies. Zhang et al. [22] compared functional outcome of 36 patients with the SCPLE implant to 30 patients with the CHP using the Constant–Murley score and demonstrated no significant difference between groups. Erdle et al. [23] compared the results of 19 patients with CHP and 13 patients with SCPLE fixation, and they reported no significant difference between the groups when using the Constant score, the Oxford shoulder score, and the subjective shoulder value.

In the current study, the bicortical lateral fragment size was significantly smaller in the CHP group. Erdle et al. [23] reported no significant difference in lateral fragment size; however, they did not report whether the intact lateral fragment was bicortical. In the current study, the largest intact bicortical lateral fragment size which would allow for adequate screw fixation was measured. Our results indicate implant selection was influenced by the bicortical lateral fragment size. We recommend further research to focus on lateral fragment size to determine whether lateral fragment size negatively affects functional outcome and complication rates with the use of different implants.

Previous case series have shown the SCPLE to be an effective fixation method for the treatment of unstable Neer type II fractures [12–17]. Zhang et al. [22] treated fractures with a lateral fragment size larger than 2 cm with the SCPE, and comminuted fractures close to the acromioclavicular joint were treated with the CHP with additional ligament repair. The comparative study by Erdle et al. [23] only included Neer type IIb fractures. To our knowledge this is the first study to evaluate the use of SCPE fixation for the treatment of Neer type V fractures. In the current study, treatment with SCPE fixation resulted in good functional outcome in both 30 patients (63%) with Neer type II and 18 patients (38%) with Neer type V fractures. These findings indicate SCPE fixation is also an acceptable treatment option for acute Neer type V fractures, despite their comminuted character.

There was no a significant difference in complication rate between CHP and SCPE fixation, which is in contrast to previous comparative studies. Zhang et al. [22] found a significantly higher complication rate, 23.3% in the CHP group compared to 5.6% in the SCPE group (\( p=0.04 \)). However, Zhang et al. [22] included symptomatic hardware as a complication, and they reported three cases (10%) of symptomatic hardware in the CHP group and none in the SCPLE group. Erdle et al. [23] also reported a significantly higher overall prevalence of complications in the CHP cohort (89%) compared to the SCPE cohort (38%) (\( p=0.014 \)). Erdle et al. [23] included radiographical proof of persistent acromial osteolysis and posttraumatic acromioclavicular joint arthrosis as complications.

The previous comparative studies included complications such as acromial osteolysis, posttraumatic acromioclavicular joint arthrosis, and sub-acromial impingement syndrome. These complications could be regarded as CHP implant specific. The CHP is fixated with a small hook under the acromion, posterior to the acromioclavicular joint which acts as a lever and maintains fracture reduction. However, this mechanism not only limits abduction of the arm, it may also affect the acromion and induce discomfort. The SCPE does not interfere with the acromioclavicular joint, which results in the absence of
acromial and impingement complications. Furthermore, there are several reports that indicated that these CHP implant-specific complications can resolve after removal [11, 24]. Renger et al. [11] evaluated the use of the CHP in 44 patients, and 30 patients (68%) reported implant-related discomfort. Renger et al. [11] found all implant-related complaints and osteolytic defects to disappear after implant removal.

Implant-related irritation and implant removal were analyzed using the series of questions developed by Hulsmans et al. [21]. In the current study, all CHP implants were removed after a mean of 4.3 months, in line with previous studies recommending CHP removal after fracture consolidation [11]. The comparative study by Zhang et al. [22] reported all CHPs were removed compared to 12 SCPLEs (33%). Erdle et al. [23] reported CHP removal was recommended and all CHP implants were removed after a mean period of 4.7 months. In the Erdle et al. [23] study, 77% of SCPLEs implants were removed after a mean period of 12.5 months due to local irritation or on patient’s explicit request. In the current study, after a minimum of 12 months following ORIF, 42% of SCPLEs implants were removed. Moreover, 43% of the patients with the SCPE still in situ reported not to experience any irritation.

This study has some limitations. First, the study is limited by the retrospective nature. This study did not include prospective collection of functional and radiological measures during different follow-up times, which would increase the understanding of the impact implants have prior to implant removal. Second, fixation method was based on surgeon’s preference, which could cause bias through selection-by-indication. Therefore, different measurements were performed to determine whether lateral fragment size influenced implant selection. Finally, our study is limited by the small number of included patients in the treatment groups. However, this number is in accordance with previous comparative studies. Unfortunately, results after the use of CHP and SCPE fixation have not yet been widely studied.

To our knowledge, this is the first study to evaluate the use of the CHP and SCPE, focusing solely on implant selection without major differences in surgical technique or ligament repair. Furthermore, this is the first study to present the results of SCPE fixation for the treatment of both Neer type II and type V fractures. Unfortunately, comparison of literature remains difficult due to small sample sizes, wide variety of functional outcome scores, definitions and surgical techniques. Therefore, a large multicenter study might provide insight into long-term results following different treatment modalities, influence of different LCF fractures types, and different patient populations.

Conclusion

Both the CHP and SCPE are effective fixation methods for the treatment of unstable LCF resulting in excellent patient-reported functional outcome and similar complication rates. SCPE fixation is an effective surgical fixation method for the treatment of both Neer type II and type V LCF. The SCPE has a lower implant removal rate compared to the CHP. Therefore, if technically feasible, we recommend SCPE fixation for the treatment of unstable LCF.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval This study was approved by the local institutional review board.

Informed consent Informed consent was obtained from all individual participants included in this study.

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References

1. Nordqvist A, Petersson C (1994) The incidence of fractures of the clavicle. Clin Orthop Relat Res 300:127–132
2. Robinson CM (1998) Fractures of the clavicle in the adult. Epidemiology and classification. J Bone Joint Surg Br 80:476–484
3. Postacchini F, Gumina S, De Santis P, Albo F (2002) Epidemiology of clavicle fractures. J Shoulder Elbow Surg 11:452–456
4. Sambandam B, Gupta R, Kumar S, Maini L (2014) Fracture of distal end clavicle: a review. J Clin Orthop Trauma 5:65–73. https://doi.org/10.1016/j.jcot.2014.05.007
5. Oh JH, Kim SH, Lee JH et al (2011) Treatment of distal clavicle fracture: a systematic review of treatment modalities in 425 fractures. Arch Orthop Trauma Surg 131:525–533. https://doi.org/10.1007/s00402-010-1196-y
6. Neer CS 2nd (1968) Fractures of the distal third of the clavicle. Clin Orthop Relat Res 58:43–50
7. Neer CS 2nd (1963) Fracture of the distal clavicle with detachment of the coracoclavicular ligaments in adults. J Trauma 3:99–110
8. Rokito AS, Zuckerman JD, Shaari JM et al (2002) A comparison of nonoperative and operative treatment of type II distal clavicle fractures. Bull Hosp Jt Dis 61:32–39
9. Ropars M, Thomazeau H, Huten D (2017) Clavicle fractures. Orthop Traumatol Surg Res 103:S53–S59. https://doi.org/10.1016/j.otsr.2016.11.007
10. Muramatsu K, Shigetomi M, Matsunaga T et al (2007) Use of the AO hook-plate for treatment of unstable fractures of the distal clavicle. Arch Orthop Trauma Surg 127:191–194. https://doi.org/10.1007/s00402-006-0284-5

11. Renger RJ, Roukema GR, Reurings JC et al (2009) The clavicle hook plate for Neer type II lateral clavicle fractures. J Orthop Trauma 23:570–574. https://doi.org/10.1097/BOT.0b013e318193d878

12. Lee SK, Lee JW, Song DG, Choy WS (2013) Precontoured locking plate fixation for displaced lateral clavicle fractures. Orthopedics 36:801–807. https://doi.org/10.3928/01477447-20130523-28

13. Andersen JR, Willis MP, Nelson R, Mighell MA (2011) Pre-contoured superior locked plating of distal clavicle fractures: a new strategy. Clin Orthop Relat Res 469:3344–3350. https://doi.org/10.1007/s11999-011-2009-5

14. Tiren D, Vroemen JP (2013) Superior clavicle plate with lateral extension for displaced lateral clavicle fractures: a prospective study. J Orthop Traumatol 14:115–120. https://doi.org/10.1007/s10195-013-0228-0

15. Beirer M, Siebenlist S, Cronlein M et al (2014) Clinical and radiological outcome following treatment of displaced lateral clavicle fractures using a locking compression plate with lateral extension: a prospective study. BMC Musculoskelet Disord 15:380. https://doi.org/10.1186/1471-2474-15-380

16. Fleming MA, Dachs R, Maqungo S et al (2015) Angular stable fixation of displaced distal-third clavicle fractures with superior precontoured locking plates. J Shoulder Elb Surg 24:700–704. https://doi.org/10.1016/j.jse.2014.09.024

17. Vaishya R, Vijay V, Khanna V (2017) Outcome of distal end clavicle fractures treated with locking plates. Chin J Traumatol 20:45–48. https://doi.org/10.1016/j.chj.2016.05.003

18. Gummesson C, Ward MM, Atroshi I (2006) The shortened disabilities of the arm, shoulder and hand questionnaire (QuickDASH): validity and reliability based on responses within the full-length DASH. BMC Musculoskelet Disord 7:44. https://doi.org/10.1186/1471-2474-7-44

19. Beaton DE, Wright JG, Katz JN (2005) Development of the QuickDASH: comparison of three item-reduction approaches. J Bone Joint Surg Am 87:1038–1046. https://doi.org/10.2106/JBJS.D.02060

20. Ferreira-Valente MA, Pais-Ribeiro JL, Jensen MP (2011) Validity of four pain intensity rating scales. Pain 152:2399–2404. https://doi.org/10.1016/j.pain.2011.07.005

21. Hulsmans MHI, van Heijl M, Houwert RM et al (2017) High irritation and removal rates after plate or nail fixation in patients with displaced midshaft clavicle fractures. Clin Orthop Relat Res 475:532–539. https://doi.org/10.1007/s11999-016-5113-8

22. Zhang C, Huang J, Luo Y, Sun H (2014) Comparison of the efficacy of a distal clavicular locking plate versus a clavicular hook plate in the treatment of unstable distal clavicle fractures and a systematic literature review. Int Orthop 38:1461–1468. https://doi.org/10.1007/s00264-014-2340-z

23. Erdle B, Izadpanah K, Jaeger M et al (2017) Comparative analysis of locking plate versus hook plate osteosynthesis of Neer type IIB lateral clavicle fractures. Arch Orthop Trauma Surg 137:651–662. https://doi.org/10.1007/s00402-017-2645-7

24. Tiren D, van Bemmel AJ, Swank DJ, van der Linden FM (2012) Hook plate fixation of acute displaced lateral clavicle fractures: mid-term results and a brief literature overview. J Orthop Surg Res 7:2. https://doi.org/10.1186/1749-799x-7-2

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