A novel approach wound closure for total joint arthroplasty procedures

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Introduction

The importance of a successful wound closure is well known and well described in the orthopedic literature. While wound appearance is important to the patient, it is the successful healing of that surgical wound that is of utmost importance to the surgeon. An evolution has taken place over the last decade with the goal of improvement in not only the appearance of the post-operative wound, but also the elimination of wound complications.

This trend has occurred in all aspects of wound closure. For example, barbed sutures were introduced as an attempt to control the post-operative hemorrhage often seen in the post-operative period and have demonstrated effective, water-tight closure. The importance of a water-tight closure is recognized in the literature, as are the potential complications if it is not obtained and extended wound drainage occurs [1-4]. Wound drainage has been associated with periprosthetic joint infection, morbidity, mortality, length of stay, and wound complications [5,6]. These complications are usually treated with absorbent dressings, temporary joint immobilization, or surgical treatment such as debridement, irrigation, and antimicrobial agent application, placing a costly burden on patients and the healthcare system [4].

The barbed suture offers the theoretical advantage of creating a more even distribution of tension across the wound. It has become popular in orthopedic surgery due to the perception that it can shorten total operative time while maintaining similar rates of infection [7]. Nett et al reported superior water-tightness using barbed sutures compared to the standard interrupted technique (with fluid leakage of 174 ml vs. 601 ml on average, respectively) [8]. Several studies report on the shorter closure time with the use of barbed sutures [9,10], but a 2014 meta-analysis found a lower superficial infection rate with standard sutures [11]. It has been suggested that the increased surface area of barbed sutures may provide larger surfaces for bacterial proliferation and nidi for infection [12,13].

While surgical staples have long been the gold standard for wound closure, increased superficial infections have been described with their use. Some evidence shows that there may be no difference or superior outcomes when using staples in a variety of specialties [14-16], but others report increased rates of complications. Ando et al showed increased rates of wound infections with patients receiving staples (8.0 %) compared to those receiving sutures (0 %) in spinal surgery, and the majority of the infections in the staples group were deep infections [17]. Smith et al. [18] report a meta-analysis on closure technique used on orthopedic patients; data from six studies, consisting of 683 orthopedic procedures, showed that the risk of infection was more than three times higher when staples were used (relative risk 3.83, 95 % confidence interval 1.38 to 10.68). On subgroup analysis of hip surgery alone, the authors found the risk of infection was over four times greater when staples were used (relative risk 4.79, 95 % confidence interval 1.24 to 18.47). However, no difference was seen in rates of inflammation, discharge, dehiscence, necrosis, or allergic reaction in the same study. There is considerable heterogeneity between these studies, and a clear answer to the optimal approach to wound closure has not been established. Due to these potential complications, the search continues for improved wound closure techniques that are efficacious as well as cosmetically pleasing to the patient.

ClozeX® is an FDA approved medical device with well-established effectiveness in over 10,000 surgical procedures [19]. This new technology utilizes an interlaced design to rapidly apply tension along incision lines and secure wound closure. In practice, this technique provides shorter closure times and improved or similar outcomes to closure by conventional means.

The ClozeX® system is recommended for low-to medium-tension surgical wounds, or as an adjunct to high tension wounds closed with conventional methods [19]. It is not recommended for areas of skin with excess fluids, oils, or hair, which render poor surfaces for adhesion of the ClozeX® system. It is also not recommended for use on infected areas due to the potential risk for device failure.

The technique

At the conclusion of surgery, deep tissue layers are closed in the conventional manner. The subcutaneous tissue is also closed in either a running or interrupted fashion based on surgeon preference. The author prefers a running stitch to further enhance the watertight properties of the wound. Prior to skin closure, the ClozeX® device is selected by first choosing the appropriate size to match the length of the wound. There are 11 different sizes, ranging from the long and narrow 3/8" (10mm) to the squarer shaped 4" (100 mm). If a large wound must be closed, as with long surgical incisions, multiple ClozeX® devices may be combined at the time of application. This may also be done for incisions on irregular or contoured surfaces, such as those with multiple skin folds. For example, for a typical Total Knee Arthroplasty (TKA) wound, 3 or 4 of the 40mm ClozeX® devices would be utilized.

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The skin is cleaned prior to application of the film. All debris, exudates, and oils are effectively removed from an area of 5-7 cm around the incision. The area is then dried to ensure proper adhesion of the film.

The film’s adhesive portions are covered with numbered protective liners (1,2,3), which help facilitate liner removal and device application. “Liner 1” is first taken off to expose the adhesive pads. While holding the two “Liner 2s” on either side, the film is applied 1-2 mm from each side of the wound edge with the adhesive surface applied on the skin. Each Liner 2 is removed gently in succession and each pad is pressed to secure it firmly on both sides of the wound. The pull tabs are then lifted in either hand, and the two Liner 3s are removed to secure the adhesive straps. The tabs are pulled in opposite directions to close the wound gently and uniformly. This provides excellent 3D control of the skin edges. When the skin edges are properly aligned under tension, the straps are lowered onto the adhesive pads below. Finally, the pulled tabs are easily removed along a perforated line (Figures 1 and 2).

The film is never lifted or reapplied once it touches the skin. If erroneously placed, a new film should be used.

Discussion

Over the last decade, new techniques have been described to aid in the final layer of wound closure. In an attempt to produce superior outcomes, devices such as Dermabond®, Dermabond® Prineo®, the Zip® Skin Closure System, and ClozeX® have been developed.

Dermabond® is a topical skin adhesive used to augment tensile strength to deep dermal sutures and provide a barrier to surgical wounds. It is composed of a 2-octyl cyanoacrylate (2-OCA) liquid that forms a protective glue on incision sites. Daykan et al. [20] recently conducted a randomized trial and reported similar outcomes between Dermabond® and monofilament epidermal sutures in cesarean procedures, including length of stay, closure time, closure appearance, blood loss, infections, and wound disruption. Martin et al. [20] found no difference between cosmetic outcomes after closure with Dermabond® and subcuticular sutures, although mean operative time was shorter in the Dermabond® group (1.4 minutes vs. 8.6 minutes) [21].

The Dermabond® Prineo® skin closure system utilizes the same 2-OCA technology along with a self-adhering mesh. This system has demonstrated improved cosmesis, reduced healing time, and shorter operative time when compared to traditional alternatives [22,23]. However, cutaneous complications, such as severe allergic dermatitis, have been linked to its use [24,25]. Huemer et al. [22] reported 4 cases of contact dermatitis, and postulate this may be due to prior exposure as each patient had previously used this device. Parvizi et al. [23] suggest that the Prineo® system should not be used in patients with known or suspected sensitivity to cyanoacrylate, formaldehyde, or adhesives. It is also recommended that tissue adhesives not be used to close contaminated or infected wounds, and should be avoided in areas prone to frequent moisture or friction, such as the hands and feet [26].

In addition, the cost of adhesives may be up to 4 times as expensive as sutures, and they require a meticulous technique to ensure no gaps or bleeding exist between wound edges [27].

Another alternative to conventional sutures is the Zip® 16 Surgical Skin Closure System, an adjustable and reversible skin closure device. Patients have reported less pain during removal when compared to staples [28] or sutures, [29] and fewer wound complications have been reported [30]. The existing literature regarding this technology has been positive, with adverse events being, at worst, similar to those of conventional methods [30-32]. It is unknown whether the incidence of surgical site infection is lower with the closure device compared to alternative methods, [33] however, and whether holding strength is different from that of staples [34].

The ClozeX® system has been shown to be an improvement on conventional wound closure methods (Figures 3–5). In a study comparing coaptive film and monocryl sutures in pediatric spine surgery, Gottklau et al. [35] found that using ClozeX® resulted in a mean of 384.13 seconds shorter closure time ($p < 0.01$), and no difference in cosmetic score. Shorter closure time, improved aesthetics, less inflammation, and similar rates of dehiscence, pain, and infection have also been reported [36-40].
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Figure 3. Healing Surgical Wound after ClozeX® Application in Hip Surgery

Figure 4. Healing Surgical Wound after ClozeX® Application in Spine Surgery

Figure 5. ClozeX Closure Application
ClozeX® also appears to be a more cost-effective closure method. Rebello et al. [19] found that estimated operating room cost is about $2,250 for 30 minutes, which translates to savings of greater than $550 if the ClozeX® system is used due to its fast and efficient application. Overall cost savings are estimated to be $964, based on these data [19]. In a study by Shipsett [41], the average per-minute operating room fee in the United States was $66 in 2005. However, fees charged to patients usually do not reflect the actual costs. According to an editorial by Macario [42], some operating room administrators consider a ballpark “true cost” of operating room time to be approximately $15 to $20 per minute for a basic surgical procedure. With this novel wound closure device, shorter operating times, low complication rates, and no need for follow up visits to remove sutures translate to considerable cost savings.

Conclusion

The ClozeX™ technique shows promise to not only improve the post-operative appearance of a wound, but also a cost-effective alternative to enhance wound closure. It reduces operative time, lowers the rate of surgical site infections, and produces similar or superior outcomes to those of staples and skin adhesives. This ClozeX™ system provides a reasonable alternative to conventional and other novel wound closure devices. This device may be of special use in high risk wounds where a post-operative wound vacuum dressing is being considered. The ClozeX® device would allow not only a cosmetic and efficacious closure, but would not hamper the potential positive effects of wound vac therapy.

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