Randomized Controlled Trial of a Novel Silicone Device for the Packing of Cutaneous Abscesses in the Emergency Department: A Pilot Study

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Objective: Traditional treatment of abscesses in the emergency department includes packing with gauze; however, this can add pain and discomfort to the procedure and frequently involves a follow-up visit for packing removal. Alternatives to gauze packing have been proposed, but they may have disadvantages in the context of emergency care. The objective of this pilot study was to investigate the use of a novel silicone packing device — the Derma-Stent.

Methods: This was a randomized controlled pilot study of 50 patients from two urban emergency departments with uncomplicated superficial abscesses. The primary outcome was the likelihood of self-removal of packing gauze versus the silicone device. Secondary outcome measures included subjective clinician and patient metrics, such as pain and ease of use.

Results: Patients identified with simple cutaneous abscesses were randomized to intervention (packing with the novel silicone device, n=25) or standard care (gauze packing, n=25). Mean age was 36 years, 54% were female, and 96% identified as African American. Although it took longer to place, the silicone device (19.0 vs 15.3 minutes, \( p = 0.03 \)), pain scores were significantly lower (4.3/10 vs 7.1/10, \( p = 0.008 \)) and ease of use reported by physicians better in the silicone-device group (4.8/5 vs 4.0/5, \( p = 0.002 \)). A high unknown rate in the gauze-packing group limits discussion of the likelihood of self-removal; however, the silicone device was more likely to remain in place (60%) versus the gauze packing at 3 days (24%, \( p = 0.01 \)).

Conclusion: This pilot randomized controlled trial compared the treatment of packing cutaneous abscesses with gauze versus using the silicone device. Limitations in the data prevent discussion on likelihood of self-removal. However, the silicone device was more likely to remain in place at day 3 follow-up and was equally effective to gauze packing in abscess reduction while also improving patient-reported pain scores. It did take longer to place the silicone device; however, physicians reported better ease of use and removal. This pilot study is encouraging for additional larger-scale trials that are required to further assess the utility of this device in the emergency department.

Keywords: abscess, Derma-Stent, silicone device, gauze, packing

Introduction

Cutaneous abscesses are an infectious condition that affects millions of people each year, and their incidence is on the rise. Between 1993 and 2005, the annual number of emergency department (ED) visits in the US for skin and soft-tissue
infections increased from 1.2 to 3.4 million, largely attributed to a rise in the prevalence of methicillin-resistant *Staphylococcus aureus* in the community.\(^1\)\(^2\)\(^3\)\(^4\)

Although currently under debate,\(^3\)\(^4\) many providers believe that for uncomplicated cutaneous abscesses seen in the ED, the standard treatment is incision and drainage at the bedside.\(^5\) It is also not uncommon to insert a strip of gauze packing to maintain the open surgical incision, enhance drainage, and prevent reaccumulation of the abscess. If the abscess is packed, patients may be instructed to return to the ED in 2–3 days to have the packing removed. However, this gauze packing may not be effective, as it often falls out spontaneously and prematurely without providing any clinical benefit.\(^6\) Additionally, the standard packing procedure and its removal may be painful, due to the gauze hardening after absorbing purulent discharge and blood.

A more novel approach advocates for the use of a vessel loop, such as a silicone string loop, for purposes of packing and drainage.\(^7\)\(^8\)\(^9\)\(^10\) While this approach may reduce pain and may be superior in terms of healing time and treatment failure, particularly in children,\(^10\) it is not readily available in the ED or outpatient clinics. String-loop devices are typically sterile silicone devices used in vascular procedures. However, for abscesses, the smaller incision needed might preclude irrigation of the abscess cavity, and the vessel loops may be susceptible to malfunction, since they typically require a surgical knot through two incisions.

The objective of the present study was to investigate the use of a novel silicone packing device, the Derma-Stent, which has been developed for use in the ED as a commercial device similar to the vessel loop. The device has multiple benefits due to its silicone design: it does not absorb bodily fluids, it increases patient compliance, and it decreases device failure, while achieving the objective of wound healing by maintaining the opening of the surgical incision(s). The first use of this device was described in a case report, and led to a successful clinical and cosmetic outcome.\(^11\) The device is similar to other loop devices that have resulted in lower failure rates in children, but otherwise equivocal rates in adults.\(^10\) In this paper, we describe a pilot randomized control trial (RCT) of the silicone device. Compared to standard gauze packing, we posit that this novel commercial device will facilitate self-removal, be better tolerated by patients, and be more likely to remain in the abscess cavity until proper removal.

### Methods

We conducted a pilot RCT comparing the silicone device to traditional gauze packing in cutaneous abscesses of patients who presented to the ED. As a measurement of perceived agency over self-care, the primary outcome was the likelihood of self-removal of the silicone device versus gauze packing. Study participants were selected from consecutive patients presenting to either of two urban EDs in Detroit, MI, USA, with cutaneous abscesses large enough to require bedside incision, drainage, and packing, as determined by ED providers. Prospective participants were screened, consented, and enrolled by trained research associates 24 hours a day, 7 days a week between February 2017 and June 2018. Prior to randomization, abscesses had been measured with bedside ultrasound by trained personnel supervised by ultrasound fellowship–trained attending ED physicians. Inclusion criteria were age 18–90 years, total abscess dimensions \((x + y + z \text{ cm})\) on ultrasound \(\geq 5\) cm, assessment by an attending physician that the abscess would require incision, drainage, and packing, and informed consent. Exclusion criteria were patients requiring admission during initial ED visit, abscess drainage requiring procedural sedation or formal incision in the operating room, inability to comprehend consent or follow-up instructions, and prisoners.

Following consent and randomization, the treating ED physician sterilized the skin and anesthetized the abscess utilizing standard departmental protocols and supplies. The specific technique for localized anesthesia was at the discretion of the treating ED physician and not protocolized by the study; however, any patients receiving general anesthesia or systemic sedation were excluded from the study. All abscesses were irrigated, then packed using either an iodine-impregnated gauze strip (Curity Iodoform packing strip; Medtronic, Minneapolis, MN, USA) or the silicone device (Derma-Stent; Mar-Med, Grand Rapids, MI, USA). Figure 1 is a schematic illustration of the device and various configurations for its deployment, including single incision versus dual incision. In order to fit various sizes and shapes of abscess cavities, treating physicians can trim the body length or longitudinal arms of the device for insertion into one or two incisions. At physicians’ discretion, the single- or double-incision technique can be employed. Provision of antibiotics was also at the discretion of the providers, but institutional guidelines strongly recommend antibiotics with methicillin-resistant *Staphylococcus aureus* coverage.

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**Figure 1** A schematic illustration of the Derma-Stent device.
Enrolled patients were scheduled for follow-up on day 3, during which they answered a number of survey questions. All packing gauze or devices remaining in abscess cavities were removed by the patients themselves or by the treating physician, and abscess-cavity size was measured again. We performed a chart review at day 30 to assess for treatment failures, defined as a need for reincision or hospitalization for antibiotic therapy.

The ability of the patients to remove the device was the primary outcome. Other outcome measures included clinical failure, defined by the presence of fever, an increase in maximal diameter of erythema, or worsening of wound swelling and tenderness, cosmetic result at follow-up visits, as measured subjectively by participants utilizing a 1–5 Likert scale, pain during the procedure and packing removal measured using a 0–10 visual analogue scale, and acceptability and satisfaction surveys by both providers and participants regarding different aspects of the procedure, such as pain, duration, and ability to maintain a sterile field.

Descriptive statistics were used for comparison with data, and are presented as means ± SD. For exploratory analysis, 95% CIs were utilized to evaluate effect size. Categorical and continuous results were analyzed utilizing chi-squared test and t-test probabilities, respectively, with \( p<0.05 \) considered significant. Due to the pilot nature of this study, the effect size was unknown and formal power analysis could not be performed. As such, 25 patients per treatment group were selected. Study data were collected and managed using REDCap (Research Electronic Data Capture) electronic data-capture tools hosted at Wayne State University (Detroit, MI, USA). REDCap is a secure, web-based application designed to support data capture for research studies. Analysis was performed with SAS software (SAS Institute, Cary, NC, USA). The institutional review board of Wayne State University approved this human research study, which was performed in accordance with the Declaration of Helsinki. The study was registered on [www.clinicaltrials.gov](http://www.clinicaltrials.gov) (NCT03171714).

**Results**

Fifty subjects were recruited and randomized evenly between the investigational silicone device and gauze.
packing. Mean age of the subjects was 36 years, 54% were female, and 96% identified as African American. Of the 25 patients in each group, 15 in the gauze-packing group and 22 in the silicone-device group were seen at day 3 follow-up appointments (Figure 2). Postrandomization baseline data are presented in Table 1. Most variables were similar between groups; however, there were more unemployed patients with a lower level of education in the silicone-device group (Table 1). Average abscess size was similar between groups, though the procedure time for placement of the silicone device was almost 4 minutes longer (15.3 vs 19.0 minutes, \( p = 0.03 \); Table 2). Perceived pain was significantly lower (4.3/10 vs 7.1/10, \( p = 0.008 \)) and ease of use reported by the physician was better in the silicone-device group (\( p = 0.002 \)).

At the day 3 follow-up appointment, abscess sizes had improved in both groups compared to day 0, but no differences were detected between groups. Such measures as postinsertion pain and bleeding/discharge were not different, but ease of removal by the physician was improved in the silicone-device group (\( p = 0.011 \)). Although there were a high number of unknown removals (n=12) in the gauze-packing group, the silicone device was more likely to remain in place at the day 3 follow-up than the gauze packing (60% vs 24%, respectively; \( p = 0.010 \)).

There were no reported adverse outcomes during the initial 3 days of the study; however, 30-day chart review identified one patient with a history of HIV in the silicone-device group that did not return for the day 3 follow-up appointment but rather came back to the ED around 2 weeks later and was admitted for intravenous antibiotics. The patient was not febrile or septic, and did not require reincision of the abscess.

**Discussion**

Cutaneous abscesses remain a common cause of ED visits, and there a variety of management options available. Although there is some debate about the utility of wound packing,\(^3\)\(^4\)\(^6\) many providers continue to use packing materials to facilitate ongoing drainage after initial incision and drainage, especially for abscesses >5 cm in diameter or in patients who are at high risk of treatment failure. Because of the pain and discomfort associated with the use of traditional gauze packing, a more comfortable and secure alternative is desirable. The need for an effective packing material that can be well tolerated has led to improvised methods of packing from a variety of materials, as well as the development of novel silicone-packing solutions,\(^10\) including the silicone device.\(^11\)

The silicone device was derived from the concept of a price-tag fastener. Its small longitudinal bars can maintain its position, and can thus be modified based on the size and shape of the abscess. While similar devices have not resulted in reduced failure rates in adults, they have been shown to be superior to gauze packing in children.\(^10\) In this pilot RCT, the ability of the patients to remove the silicone device versus gauze packing was our primary outcome. However, limitations in the study design and patient follow-up did not allow for commentary as intended. Alternatively, presence of the packing device in the abscess cavity at day 3 heavily favored the silicone-packing device, validating the functionality of the device design. Furthermore, the silicone device was shown to be equally effective as gauze packing in abscess reduction while improving patient-reported pain scores, despite the procedure taking 3.7 minutes longer. The physicians reported better ease of use and removal. Despite an unequal number of missing data regarding removal of the packing gauze, the silicone device appeared to remain in the abscess more effectively than gauze. Overall, with the lone exception of procedure time (which may improve with experience), the silicone device was superior or...
equivalent in terms of patients, physicians, and clinical outcomes.

There was one potential adverse event identified on 30-day chart review. A patient in the silicone-device group with a history of HIV had a subsequent ED visit and admission for intravenous antibiotics. As the patient did not return at day 3, it could not be determined if the device played a role in treatment failure.

### Limitations

As a pilot, this study was limited primarily by its small sample and discrepancy in follow-up rates for the gauze-packing group, which directly impacted our primary outcome. With 12 patients (48%) categorized as “unknown removal”, this limited our conclusions regarding this pilot trial. Additionally, another shortcoming was the lack of standardized protocols for the provision of antibiotics and detailed, protocolized techniques for the incision and drainage procedures, such as method of anesthesia, amount of irrigation, length of incision, and amount of packing in the gauze packing. Although this strategy confers the benefits of a pragmatic trial design, it limits precise comparisons and isolation of the intervention as an independent variable. Subsequent RCTs with this device should also expand their follow-up times to further elucidate any possible relationship between the silicone device and adverse outcomes, such as seen in one patient.

### Conclusion

This pilot RCT compared the treatment of packing cutaneous abscesses with gauze versus using a silicone device. The silicone device was more likely to remain in place at day 3 follow-up, and was equally

| Table 1 Baseline variables for patients packed with gauze or silicone device |
|-------------------------------|-----------------------------|---------------------------|  
|                               | Gauze packing | Silicone device | p  
| n/mean (%/SD)                | n/mean (%/SD)   | p              |
| **Demographics**             |               |                |  
| Age (years)                  | 33 (10)       | 39 (14)       | 0.192 |
| Female                       | 16 (64)       | 11 (44)       | 0.156 |
| Employed or student          | 18 (72)       | 12 (48)       | 0.044 |
| Insured                      | 22 (88)       | 23 (92)       | 0.672 |
| **Education (highest level attained)** |               |                |  
| Elementary/high school       | 8 (33)        | 14 (61)       | 0.048 |
| Associate degree or technical school | 3 (12) | 2 (9) | 0.664 |
| College/university           | 13 (54)       | 7 (30)        | 0.081 |
| **Identified ethnicity**     |               |                |  
| African American             | 24 (96)       | 24 (96)       | —     |
| Hispanic or Latino           | 1 (4)         | 0             | —     |
| Other                        | 1 (4)         | 1 (4)         | —     |
| **Medical history**          |               |                |  
| Diabetes (type 2)            | 5 (20)        | 5 (20)        | —     |
| Hidradenitis suppurativa     | 5 (20)        | 3 (12.5)      | 0.855 |
| HIV                          | 1 (4)         | 2 (8)         | —     |
| Hypertension                 | 6 (24)        | 11 (44)       | 0.114 |
| Immunosuppressive therapy    | 0             | 1 (4)         | —     |
| Congestive heart failure     | 0             | 1 (4)         | —     |
| **Substances of abuse**      |               |                |  
| Alcohol                      | 3 (12)        | 0 (0)         | 0.235 |
| Tobacco                      | 18 (72)       | 22 (88)       | 0.157 |
| Heroin                       | 1 (4)         | 2 (8)         | —     |
| Cocaine                      | 1 (4)         | 2 (8)         | —     |

Note: “—” represents no significant difference or insufficient data for analysis.
effective as gauze packing in abscess reduction, while improving patient-reported pain scores. It did take longer to place the silicone device; however, physicians reported better ease of use and removal. These results are very promising as pilot data and provide justification for a larger, multicenter trial, but would likely need more standardized protocols for the incision and drainage procedures. Future models of the device could be modified and enlarged for use with abscesses drained in the operating room by surgeons.

### Data Sharing Statement
Deidentified study-participant data will be made available for academic purposes upon request to the corresponding author. Data include recruitment tools, consent forms, and participant baseline and outcome data. All reasonable requests will be honored.

### Author Contributions
All authors contributed to data analysis, drafting or revising the article, have agreed on the journal to which the article will be submitted, gave final approval to the version to be published, and agree to be accountable for all aspects of the work.

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### Table 2 Outcomes of patients randomized to abscesses packed with gauze or silicone device

|                      | Gauze packing | Silicone device | p      |
|----------------------|---------------|-----------------|--------|
|                      | Mean (SD)     | n               | Mean (SD) | n | 0.678 |
| **Day 0**            |               |                 |         |    |       |
| Abscess size (cm, x + y + z) | 6.5 (1.6) | 25 | 6.8 (1.6) | 25 | 0.030 |
| Procedure time (minutes) | 15.3 (11.3) | 25 | 19.0 (7.8) | 25 |        |
| **Patient-reported outcomes** |     |                 |         |    |       |
| Pain during procedure (0–10) | 7.1 (2.9) | 25 | 4.3 (3.6) | 25 | 0.008 |
| Experience as a whole (1–5) | 4.0 (1.4) | 25 | 4.5 (1.0) | 25 | 0.114 |
| Satisfaction with time (1–5) | 4.7 (0.6) | 25 | 4.7 (0.5) | 25 | 0.772 |
| **Physician-reported outcomes** |     |                 |         |    |       |
| Ease of use (1–5) | 4.0 (1.0) | 25 | 4.8 (0.5) | 25 | 0.002 |
| Ability to maintain sterile field (1–5) | 3.5 (1.2) | 25 | 4 (1.1) | 25 | 0.142 |
| **Day 3 follow-up** |               |                 |         |    |       |
| Abscess size (cm, x + y + z) | 3.6 (1.4) | 14 | 3.6 (1.5) | 21 | 0.884 |
| Device/packing already removed by patient |    | 7 | 6 | 0.732 |
| Device/packing removed by physician | 6 | 15 | 0.010 |
| Unknown removal | 12 | 4 | 0.031 |
| **Patient-reported outcomes** |     |                 |         |    |       |
| Pain since procedure (0–10) | 4.2 (2.7) | 14 | 3.4 (2.8) | 21 | 0.246 |
| Discharge and bleeding (1–5) | 3.8 (1.4) | 14 | 3.7 (1.2) | 21 | 0.711 |
| **Physician-reported outcomes** |     |                 |         |    |       |
| Ease of removal (1–5) | 4.8 (0.4) | 10 | 4.8 (0.5) | 19 | 0.570 |
| Satisfaction with removal (1–5) | 4.0 (1.0) | 11 | 4.8 (0.5) | 20 | 0.011 |
| Appearance of scar (1–5) | 3.9 (0.9) | 13 | 4.3 (0.9) | 21 | 0.211 |

Notes: Likert scale (1–5): 1, strongly disagree; 2, disagree; 3, neutral; 4, agree; 5 strongly agree. Pain characterized by visual analogue scale: 0, no pain; 5, distressing pain; 10, unbearable pain. Values in bold indicate p<0.05.
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