Closed-wound negative pressure therapy dressing after loop ostomy closure: a retrospective comparative study

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Closed-wound negative pressure wound therapy (NPWT) dressings were recently introduced with the purpose to reduce incisional surgical site infections (iSSI) in high-risk wounds. The aim of this study was to compare iSSI rates in patients after ostomy closure with and without additional application of a closed-wound NPWT dressing. Single-center retrospective analysis of consecutive patients undergoing ileo- or colostomy closure over an 8-year period (January 2013—January 2021). Intradermal non-purse string technique with absorbable sutures were used in all patients. Since November 2018, all patients (study group) received a NPWT device for a maximum of 5 days postoperatively (PICO, SMITH AND NEPHEW). Primary outcome was iSSI rate within 30 days of surgery. SSI was defined in accordance with the Center of Disease Control (CDC) classification and included superficial and deep incisional SSI. Data was retrieved from the institutional enhanced recovery after surgery (ERAS) database, with standardized complication assessment by trained abstractors. In total, 85 patients (25%) in the study group were comparable with 252 (75%) patients in the control group regarding demographics (age, gender, body mass index, ASA score), ostomy type and anastomotic technique (all \( p > 0.05 \)), but not wound contamination class (class III: 5% vs 0%, \( p < 0.001 \)). Median time to NPWT removal was 4 (IQR 3–5) days. Incisional SSI were observed in 4 patients (4.7%) in the study group and in 27 patients (10.7%) in the control group (\( p = 0.097 \)). These preliminary results suggest a potential benefit of systematic application of the NPWT device after loop ostomy closure. A randomized controlled study is needed.

Abbreviations

ASA  American society of anesthesiologists
BMI  Body mass index
CDC  Center of disease control
CHUV  Centre hospitalier universitaire vaudois
ERAS  Enhanced recovery after surgery
IQR  Interquartile range
iSSI  Incisional surgical site infection
LOS  Length of stay
OR  Operating room
NPWT  Negative pressure wound therapy
POD  Post-operative day
SD  Standard deviation
SSI  Surgical site infection
VAC  Vacuum-assisted closure

Incisional surgical site infection (iSSI) is the most common complication after stoma closure, with reported rates varying from 2 to 41%⁶.⁷

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Among various skin closure techniques for ostomy wounds, purse-string closure has shown superiority in reducing SSI compared to primary skin closure. Nevertheless, purse-string wounds take longer to heal than primary closure techniques and require daily wound care.

Closed-wound negative pressure wound therapy (NPWT) has been suggested as an alternative in high-risk wounds. However, data on closed-wound NPWT application after loop ostomy closure are scarce, despite encouraging preliminary results regarding clinical benefits and ease of use.

The aim of this study was to compare iSSI rates in patients after loop ostomy closure with primary skin closure with or without application of a closed-wound NPWT dressing.

Methods
Retrospective comparative study of consecutive patients undergoing loop ileostomy or colostomy closure between January 2013 and January 2021, conducted at Lausanne University Hospital (CHUV). Since January 2013, all patients were treated within an Enhanced Recovery After Surgery (ERAS) pathway and data were prospectively collected in a dedicated ERAS database. A new dressing protocol including the NPWT device was implemented systematically in November 2018 for all patients undergoing ostomy closure. These patients (study group) were compared with the cohort operated before this date (control group). The present study was approved by the ethics committee of Canton de Vaud (Commission cantonale d'éthique de la recherche sur l'être humain—CER-VD). Due to retrospective nature of the study informed consent was waived by the ethics committee (decision # 2020–238). All methods were performed in accordance with the relevant guidelines and regulations.

Baseline demographics included age, gender, body mass index (BMI), American Society of Anesthesiologists (ASA) score, current smoking status, pre-existing diabetes requiring medication, ongoing systemic chemotherapy (within 3 months before surgery) and immunosuppressive medication (within two weeks prior to surgery). Immunosuppressive medications included systemic steroid therapy (> 10 mg/day) and specific immunosuppressive drugs. Intra-operative details such as wound contamination class (defined according to the CDC classification), type of anastomosis and duration of surgery were collected.

All procedures were elective loop ileostomy or colostomy closures through direct approach. All procedures were carried out by board-certified general or colorectal surgeons. Patients not directly operated through the existing stoma site or undergoing additional abdominal procedures were excluded.

Wound management. Ostomy wounds were closed plane-by-plane in a standardized fashion. The fascia was closed with separate stitches of monofilament synthetic absorbable sutures (PDS-1, ETHICON INC, Raritan, USA) and subcutaneous tissue was adapted through separate stitches of synthetic absorbable braided sutures (Vicryl, ETHICON INC, Raritan, USA). The cutaneous layer was closed with intradermal absorbable sutures (Monocryl 4.0, ETHICON INC, Raritan, USA) in a linear fashion, according to institutional protocol (Fig. 1A). Wound dressings included surgical glue (Histoacryl, B BRAUN INC, Sempach, CH) for all patients in the control group. Since November 2018, a single-use closed-wound negative pressure wound therapy (NPWT) device (PICO, SMITH AND NEPHEW INC, Hull, UK) was applied systematically in all patients instead of surgical glue as part of an institutional SSI prevention quality improvement initiative (NPWT group) (Fig. 1B). The NPWT device was left in place until post-operative day 5 or the day of discharge if earlier.

Results. Primary outcome was iSSI within 30 days post-operatively. SSI was defined in accordance with the CDC classification, with iSSI regrouping superficial (CDC—A) and deep (CDC—B) incisional SSI. Data was retrieved from the institutional enhanced recovery after surgery (ERAS) database, with standardized complication assessment by trained abstractors for entire study period. An appointment was scheduled 6 weeks after surgery. Suggesting that there is a 30 days follow up for all patients. Secondary outcomes were length of hospital stay (LOS), return to operating room (OR) and postoperative complications at 30 days, classified according to the CDC classification.
the validated Clavien classification, with grade ≥ 3b defining major morbidity. Anastomotic leak was defined as a clinically symptomatic leak requiring return to OR.

Statistical analysis. Continuous variables were summarized as median (interquartile range: IQR) or mean ± standard deviation (SD) and categorical variables as frequencies and percentages.

Chi-square and 'Student’s t-test were used for comparison of categorical and continuous variables, respectively. Statistical significance was defined as $p \leq 0.05$. Both matching (similarity of baseline characteristics in both comparative groups) and multivariable analysis (low event rate) were not performed.

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Results

In total, 337 patients were included. The NPWT group (study group) included 85 patients (25%), while 252 patients (75%) had conventional primary skin closure (control group). Baseline demographics were comparable between both groups (all $p > 0.05$, Table 1).

Surgical details are depicted in Table 1. The proportion of ileostomy and colostomy closure was similar in both groups ($p = 0.22$). The majority of anastomoses were handsewn in both groups (overall 84%). Wound contamination class II (clean-contaminated) was reported in 94% of patients in the NPWT group, compared to 100% in the control group. Median surgical duration was around 10 min longer in the NPWT group ($p = 0.004$).

Postoperative outcomes are detailed in Table 2.

Median time to NPWT removal was 4 days (IQR 3–5). Device malfunction was observed in two patients (2.3%), requiring early removal of the NPWT dressing. Median hospital stay was 4 (IQR 3–6) days in both groups.
Incisional surgical site infections were observed in 4 patients (4.7%) in the NPWT group and in 27 patients (10.7%) in the control group (p = 0.097). Overall postoperative morbidity, major complications, return to OR and anastomotic leak rates were all similar in both groups (Table 2). Overall, one patient died in the control group due to malignant arrhythmia at POD 8.

Discussion

The present study suggests a potential benefit associated with the systematic application of a closed-wound NPWT device after loop ostomy closure. Technical issues were very uncommon. Adequate powered randomized trials are needed to confirm these preliminary findings.

NPWT has been established as a SSI preventing measure for several different indications, especially in colorectal surgery. However, the NEPTUNE trial did not describe NPWT as superior to classical wound closure.

Nevertheless, a few studies assessed the effect of NPWT application after loop ostomy closure. A recently published randomized controlled study of NPWT in primarily closed wounds after loop ileostomy closure described an ISS rate of 5.7% in the NPWT group vs. 19% without NPWT. The present series revealed a similar rate of ISS in the NPWT group, but lower rates in the control group, which may explain the lack of significant difference in the present comparison. However, the low SS rate may also be related to underreporting as a potential bias of retrospective studies. A recent observational study from Japan evaluating an open-wound VAC therapy system customized for closed wounds did not report any SSI in 50 patients after ileostomy closure.

While their results were promising, the customized technique of NPWT application on open wounds may be difficult to apply on a large scale.

In this present series, NPWT dressings were well-tolerated and easy to use, with only exceptional technical issues. Surgical duration was slightly longer, which however may also be related to constitutional factors (higher BMI and comorbidity indices in the NPWT group). The main advantage probably consists in simplified post-operative wound care, unlike the time-consuming purse-string closure. While several reports revealed lower SSI rates after purse-string closure compared to primary closure, wounds require daily care until discharge and specialist wound care for about 35 days; in our institution, the median duration of JPMA was 4 days. In contrast, primarily closed ostomy wounds without SS occurrence typically heal within 7–24 days. The subcutaneous suture is further convenient since no follow-up care is needed after device removal providing uneventful wound healing.

The present study has several limitations related to its retrospective nature without dedicated preset defined data, the small sample size of the study group and the uncontrolled setting. However, consecutive patients were included to limit any potential selection bias. Based on the positive preliminary experience of other series, our group decided to implement closed-wound NPWT therapy as a new standard of care (practice change) and to compare outcomes to the unselected pre-implementation cohort. This design was chosen given the consistency of surgical and perioperative care and standardized, prospective SSI surveillance based on our institutional ERAS protocol.

An adequately powered randomized controlled multicentric trial comparing different techniques and considering patient preference might probably be the most appropriate way to further optimize ostomy wound management. The ongoing SR-PICO randomized study (KCT0004063) may confirm our preliminary results.

In conclusion, additional closed-wound NPWT dressings after primary skin closure of ostomy wounds seems beneficial in reducing ISS. This strategy challenges the purse-string closure method in ease of management, reduction of resources and time to complete wound healing.

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**Author contributions**

P.C. and D.C. have equal contribution to the main manuscript text. All authors reviewed the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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