Comparison of tissue damage, cleansing and cross-contamination potential during wound cleansing via two methods: lavage and negative pressure wound therapy with instillation

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Abstract

The use of lavage was compared to negative pressure wound therapy (NPWT) with instillation (NPWTi) to assess extent of soft tissue damage, debris removal and environmental cross-contamination susceptibility in three distinct models. Scanning electron microscopy in an ex vivo model showed increased visible tissue trauma from lavage treatment at low and high pressures versus NPWTi, with the degree of trauma relative to the pressure of the irrigant. These results were corroborated in granulating full-thickness excisional swine wounds coated with dextran solution to simulate wound debris. Both low-pressure lavage and NPWTi demonstrated effective cleansing in this model, reducing debris by >90%. However, using three-dimensional photography to evaluate tissue damage by measuring immediate tissue swelling (changes in wound volume and depth) showed significantly greater ($P<0.05$) swelling in low-pressure lavage-treated wounds compared with NPWTi-treated wounds. Lastly, bench top wound models were inoculated with fluorescent bacterial particles to assess environmental cross-contamination potential and collected at measured distances after treatment with low-pressure lavage and NPWTi. No evidence of cross-contamination was found with NPWTi, whereas one-half of the particles became ‘aerosolised’ during low-pressure lavage ($P<0.05$). Collectively, these studies demonstrate the effective wound cleansing capabilities of NPWTi without the tissue damage and environmental contamination associated with lavage.

Introduction

Wound cleansing, a crucial step in wound management and healing, uses fluid to remove loosely adherent cellular debris and necrotic tissue from the wound bed (1). There is also a general consensus that cleansing helps reduce bacterial counts (2,3). In essence, wound cleansing removes impediments to the healing process in order to create an optimal wound healing environment. Successful management of a wound also involves removing contaminants while inflicting minimal injury to the tissue.

Key Messages

- wound cleansing, a crucial step in wound management and healing, uses fluid to remove loosely adherent cellular debris and necrotic tissue from the wound bed
- various irrigation methods that are mainly categorised as either low-pressure lavage (4–15 psi) or high-pressure lavage (35–70 psi) exist
- studies have shown that lavage, specifically high-pressure lavage, can cause considerable trauma to bone tissue

• Lavage has also been reported to cause considerable disruption to soft tissue in both in vivo and in vitro studies.
• During wound lavage, the potential for spreading pathogens, especially resistant organisms, in the hospital environment is also a cause for concern.
• The first recognized hospital-acquired outbreak associated with pulsed lavage wound care as a novel mode of transmission for Acinetobacter baumannii was in 2004.
• NPWT in conjunction with instillation (NPWTi) has the advantage of timed, automated and controlled delivery of the chosen irrigant uniformly in the wound bed followed by a programmed ‘soak’ time and subsequent removal of solution through application of NPWT.
• Other clinical benefits have been reported, including efficacy in treating wounds with high levels of exudate, debris and slough content, and use in pain management.
• Three independent studies (ex vivo, in vivo and bench top) were conducted to evaluate the effects of NPWTi in comparison to lavage on tissue trauma, cleansing and environmental cross-contamination, respectively.
• The question that many of the authors of these reports pose, though, is ‘does the cleansing capacity of lavage outweigh the damaging effects to the soft and bone tissue caused by this treatment?’
• Several studies have shown that although there is better cleansing with higher lavage pressures, the consequences include further dissemination of cleansing fluid into the interstices of the soft tissue causing considerable gross and microscopic disruption that may lead to more cell death, higher infection rates and greater bacterial retention.
• While animal models are routinely used in wound studies, an ex vivo wound model provides a cost-effective, readily available venue for performing experiments pertaining to wound healing aspects, such as cleansing, tissue trauma and bacterial penetration in soft tissue.
• The digital and SEM wound images of the ex vivo wound model successfully illustrate the differences in traumatic effects of wound cleansing; there was visually less tissue disruption in NPWTi-treated soft tissue compared with lavage-treated tissue.
• The tissue trauma observed in the scanning electron microscopic (SEM) images was in the form of ruptures and fissures in the tissue, and the intensity of the trauma could be visually assessed to be relative to the pressure of the irrigant used to cleanse the wound.
• While lavage has been the conventional method for wound cleansing, there have been more than 30 NPWTi reports published since the first clinical application of instillation therapy with NPWT.
• Many of these publications report clinical benefits of instillation therapy, including NPWTi’s wound-cleansing attributes.
• Because the ex vivo model demonstrated that high-pressure lavage caused substantial soft tissue trauma, a full-thickness acute wound healing model in normal healthy swine was used to assess the effectiveness of debris and contaminant removal by NPWTi in comparison to that of low-pressure lavage.
• The molecular weight of the dextran used to simulate particulate debris was selected to imitate proteinaceous materials found in wound exudates.
• Both regimens cleaned >90% of the simulated debris.
• Upon gross examination of the wounds after cleansing treatments, it was noted that 83% of the wounds treated with low-pressure lavage exhibited blanching.
• There was no blanching or swelling detected in association with NPWTi.
• In addition to the abundant literature about lavage’s cleansing capacity and tissue disruption propensities as a possible ‘side effect’, there are also numerous reports of infection control issues regarding lavage that contributed to hospital-acquired infections (HAIs).
• The results of our study in comparing the susceptibility of bacterial cross-contamination between NPWTi and several low-pressure lavage cleansing regimens indicate that the lavage treatments using the cleansers caused significant cross-contamination, whereas NPWTi caused none.
• NPWTi’s sealed environment virtually eliminated chances of bacterial aerosolisation while the therapy was applied.

Currently, wound cleansing is often achieved through lavage, a procedure in which the wound is irrigated under pressure with one of several widely used solutions. Isotonic (normal) saline solution is physiologically compatible and non-toxic and, thus, can maintain cell viability. Normal saline does not damage tissue, cause allergies or sensitisation or alter the normal flora of the skin; therefore, it does not interfere with the normal healing process (4).

Various irrigation methods that are mainly categorised as either low-pressure lavage (4–15 psi) or high-pressure lavage (35–70 psi) exist; examples include the bulb syringe method, gravity flow, manual pump irrigation and mechanical/pulsed lavage. The Agency for Health Care Policy and Research (AHCPP) has suggested that cleansing with low-pressure lavage is sufficient to remove surface pathogens and debris without causing wound trauma or bacterial spreading. AHCPP has also suggested that pressures above 15 psi may cause tissue damage (5). Studies have shown that lavage, specifically high-pressure lavage, can cause considerable trauma to bone tissue (6,7). In 2002, Adili et al. illustrated that this macroscopic damage had an adverse effect on early fracture healing in rat femurs (7). Lavage has also been reported to cause considerable disruption to soft tissue in both in vivo and in vitro studies (8). Furthermore, healthy tissue traumatised by lavage cleansing may have impaired capabilities in resisting infection (8,9). In addition, high-pressure lavage has been shown to increase the depth of contaminant and bacterial penetration into soft tissue (9,10). High-pressure lavage can therefore lead to complications and/or delays in the wound healing process and may even be deleterious to the wound healing progression.
During wound lavage, the potential for spreading pathogens, especially resistant organisms, in the hospital environment is also a cause for concern. For instance, the rates of detection for methicillin-resistant *Staphylococcus aureus* or vancomycin-resistant enterococci on the protective gowns and/or gloves of health care workers have been reported to be as low as 18% and as high as 67% (11–13). Reports of emerging health-care-associated pathogens that are relatively resistant to the most common surface disinfectants and alcohol-based antiseptics, such as *Clostridium difficile*, are also disconcerting (14). Therefore, given that the pressure delivered to the wound by lavage must be high enough to remove the adhered microbes from the surface of the wound bed, it is reasonable to express concern regarding the possibility of bacterial cross-contamination due to splatter. During lavage, bacteria can become ‘aerosolised’ and particulate matter disseminated. In fact, Maragakis *et al.* (15) investigated and reported the first recognised hospital-acquired outbreak associated with pulsed lavage wound care as a novel mode of transmission for *Acinetobacter baumannii*. Also, the US Food and Drug Administration (FDA) issued a medical device safety article warning of the infection risks associated with pulsed lavage (16). The use of lavage can, therefore, be potentially hazardous for the patient and the health care provider by contributing to the spread of hospital-acquired infections (HAIs) in both patients and visitors.

Negative pressure wound therapy (NPWT) is widely recognised as a powerful tool in wound care. NPWT in conjunction with instillation (NPWTi) has the advantage of timed, automated and controlled delivery of the chosen irrigant uniformly in the wound bed followed by a programmed ‘soak’ time and the subsequent removal of solution through application of NPWT (17). Studies have shown NPWTi to be effective in instilling various wound treatment solutions, such as cleansers, antiseptics and analgesics (18).

NPWTi delivered by V.A.C. Instill® Wound Therapy (KCI USA, Inc., San Antonio, TX) was first introduced to clinicians in 2003 (19). Since then, other clinical benefits have been reported including efficacy in treating wounds with high levels of exudate, debris and slough content, and use in pain management (19,20). In a recent porcine study, use of NPWTi with saline as the instillant increased collagen deposition in granulation tissue for accelerated wound fill (21).

Three independent studies (ex vivo, in vivo and bench top) were conducted to evaluate the effect of NPWTi in comparison to lavage on tissue trauma, cleansing and environmental cross-contamination. The ex vivo model was used to qualitatively evaluate the effect of NPWTi and both high- and low-pressure lavage on tissue trauma. On the basis of the results from the ex vivo study, the ex vivo porcine study quantitatively compared the effect of NPWTi and low-pressure lavage on tissue trauma and cleansing. The bench top study using synthetic polymeric wound models quantitatively evaluated the environmental effects of NPWTi and low-pressure lavage.

**Materials and methods**

**Soft tissue damage in the ex vivo wound model**

**Experimental setup**

Raw porcine meat was purchased from a local butcher shop, stored at 4°C until ready for use and sliced into approximately 4.0 cm sections. These sections were then immediately processed for wound creation. A 4.5 cm × 4.5 cm stainless steel block of 1/3 cm thickness was placed underneath the centre of the pork loin section (Figure 1A). A stainless steel template with a 5 cm × 5 cm opening in the centre was placed over the pork loin such that the stainless steel block beneath caused the tissue to protrude through the opening in the template (Figure 1B). A 15 cm Padgett Dermatome (Integra, Plainsboro, NJ), set at the highest thickness (0.076 cm), was passed over the template five times to create an approximately 5 cm × 5 cm wound of uniform thickness via three-dimensional (3D) camera photographs and software analysis (Figure 1C). Two hundred and fifty microlitres of simulated wound fluid (0.14% bovine serum albumin and 0.01 M phosphate-buffered saline, pH 7.4–8) were then applied to the pork loin wound and allowed to adhere to the tissue at room temperature for 2 hours before cleansing treatment.

**Cleansing therapy: NPWTi**

Following simulated wound fluid application to the ex vivo wound model, the wound was dressed using the new reticulated open cell foam (ROCF) V.A.C. VeraFlo™ Dressing (ROCF-V; V.A.C. VeraFlo™ Medium Dressing; KCI USA, Inc.). The dressing was cut into 5 cm × 5 cm pieces, which were placed into the simulated wound bed. A semi-occlusive adhesive drape covered the wound and dressing. NPWTi (V.A.C. VeraFlo™ Therapy; KCI USA, Inc.), which was delivered using a prototype of the V.A.C.ULTA™ Therapy System (KCI USA, Inc.), consisted of six complete cycles of negative pressure at −125 mmHg for 5 minutes and instillation of normal saline followed by a 10 seconds soak time.

**Cleansing therapy: low- and high-pressure lavage**

Following simulated wound fluid application to the ex vivo wound model, the entire wound bed was irrigated with 200 ml of normal saline with either the low- (12.7 psi) or high-pressure (45 psi) pulsed lavage devices. A commercially available low-pressure lavage device with a soft-splash shield tip was used per manufacturer’s instructions. For the high-pressure pulsed lavage device, the low setting of a commercially available jet-tipped water flosser used in medical applications was utilized at a distance of 5 cm from the wound bed.

**Tissue damage evaluation: scanning electron microscopy**

After treatment, a 10 mm biopsy punch sample was cored from the wound and digitally photographed before placement on ice
Figure 1 Process of ex vivo wound model generation. (A) 4.5 cm × 4.5 cm stainless steel block placed underneath the pork loin. (B) Stainless steel template with a 5 cm × 5 cm opening placed over the pork loin and the block underneath it so that this portion rises through the opening. (C) Wound created utilising the wound template.

Debris cleansing and tissue trauma in porcine experimental wounds

Animals

The domestic swine Sus scrofa scrofa used in this study were obtained from Gentiporc, Inc. (Alexandria, MN), housed at an Association for the Assessment and Accreditation of Laboratory Animal Care (AAALAC)-accredited facility and maintained on a twice-daily constant-nutrient diet with water received ad libitum. The protocol and its amendments were reviewed by the Institutional Animal Care and Use Committee (IACUC) for compliance with regulations prior to study initiation or implementation of amended activities. All animals received humane care.

Wound creation and experimental setup

Three female swine (61.5–64.7 kg) were used for the study. Each animal was anaesthetised with a mixture of Tiletamine (Fort Dodge Veterinaria S.A., Fort Dodge, IA), Zolazepam (Fort Dodge Veterinaria S.A.) and Xylazine (Lloyd Laboratories, Shenadoah, IO) to create dorsal full-thickness excisional wounds. Four wounds of 5 cm diameter were created on each side of the spine for a total of eight wounds. The day of wound creation was termed day 0. In order to form a light bed of granulation tissue, all wounds were dressed with ROCF (V.A.C.® GranuFoam™ Round Dressing; KCI USA, Inc.) that was cut to size and placed in each wound cavity for direct contact with the wound bed. Semi-occlusive adhesive drape (V.A.C.® Drape; KCI USA, Inc.) was used to secure the dressings on the wounds. The wounds were bridged in pairs with a T.R.A.C.™ Pad (KCI USA, Inc.) connected to an NPWT unit (V.A.C. ATS®; KCI USA, Inc.) via T.R.A.C.™ tubing (KCI USA, Inc.). The wounds received continuous NPWT of −125 mmHg for a 4-day period to establish a lightly granulated wound bed. On day 4, the animals were anaesthetised and the dressings removed, showing a thin base layer of granulation tissue at the wound bed. The wounds were gently wiped with saline-soaked gauze prior to application of the simulated wound debris. A 10 mg/ml stock of fluorescein isothiocyanate (FITC)-conjugated dextran solution (average molecular weight 59 000–77 000 kDa; Sigma Aldrich, St Louis, MO) was prepared in dd H₂O and 0.45 ml of this solution applied to each wound prior to cleansing. The cleansing treatments were evaluated in contralateral wounds.
Cleansing therapy: NPWTi

Four wounds per pig received NPWTi. These wounds were dressed with ROCF-V and semi-occlusive adhesive drape connected to the therapy system according to its instructions for use. The system was programmed to deliver NPWTi with the following settings: negative pressure at $-125 \text{ mmHg}$ for 5 minutes and instillation of sterile normal saline followed by a 5-minute soak time. The duration of the negative pressure phase was reduced to 5 minutes to simulate 24 hours of typical use (ten therapy cycles) into a shorter 2-hour period.

Cleansing therapy: low-pressure lavage

Four wounds per pig received low-pressure lavage therapy utilising a commercially available low-pressure lavage device. Low-pressure lavage therapy was performed according to its instructions for use delivering 1000 ml of sterile normal saline in less than 2 minutes. Care was taken to ensure that the entire surface of the wound bed was lavaged.

Wound examination

Gross observations of the wounds included a visual inspection of the wound bed before and after cleansing treatment. The wound colour, visual presence of peripheral swelling and the amount and colour of wound exudates were amongst the observations noted.

Cleansing evaluation: digital fluorescent imaging and analysis

The cleansing abilities of the two systems were evaluated using digital fluorescence imaging of the FITC-dextran. A fluorescence camera system was assembled using a digital camera (Canon Rebel T1i; Canon U.S.A. Inc., Melville, NY), an ultraviolet light source and an opaque hood which covered the wound. Images of the wounds were collected at three time points: before application of the FITC-dextran (baseline), after application of the FITC-dextran (pre-cleansing) and after cleansing treatment (post-cleansing). Images were loaded into ImageJ (Version 1.44p; U. S. National Institutes of Health, Bethesda, MD) and the green channel isolated for analysis. A threshold value for background fluorescence was determined in each baseline image. The wound edges were outlined, and the total number of pixels and the number of green pixels above the threshold were counted within each wound.

Tissue damage/swelling evaluation: 3D imaging and wound dimension measurements

Prior to and post-cleansing, 3D images of the wounds were collected with the 60 cm focal length 3D LifeViz™ camera (Quantificare S.A., Sophia Antipolis Cedex, France) to assess tissue damage. Images were imported into the DermaPix® analysis software (Quantificare) for a 3D reconstruction of each wound, allowing for accurate determination of wound depth and wound volume. Changes in wound depth and wound volume were indicative of increases or decreases in wound bed swelling. The swelling ratio was calculated by dividing the pre-cleansing wound depth or volume by the post-cleansing wound depth or volume. The swelling ratio is interpreted as follows: ratio = 1 indicates no swelling; ratio > 1 indicates increased swelling and ratio < 1 indicates reduced swelling.

Oedema evaluation: histological analyses

After cleansing, a 12 mm full-thickness punch biopsy was sampled from the centre of each wound and fixed in 10% neutral buffered formalin for paraffin embedding. Tissue samples were sectioned and stained with haematoxylin and eosin (H&E) and Masson’s trichrome for evaluation by a board-certified pathologist. Oedema in the wound surface and subjacent subcutaneous adipose tissue was scored as follows: 0 = none, 1 = minimal, 2 = moderate, 3 = marked and 4 = severe.

Environmental cross-contamination using a bench top polymeric sacral pressure ulcer model

Experimental setup

The wound model used in this study (Seymour II – 0910 Wound Care Model; VATA Inc., Canby, OR) simulated a deep stage IV sacral pressure ulcer with exposed bones, undermining, tunnelling, subcutaneous fat, eschar and slough. The model was set on a surface at a height representing that of a patient bed. To collect any droplets/splashing or ‘aerosolisation’ resulting from the wound cleansing methods, 96-well microplates were arranged around the wound in two concentric levels (Figure 2). One set of 96-well microplates was placed such that the outer edges of the wound were 7.5 cm from the centre of each plate. The second concentric set of
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microplates was arranged similarly, with the centre of each plate 15 cm from the outer edges of the wound of the model. The simulated sacral pressure ulcer of the model was then inoculated with \(6 \times 10^7\) heat-killed, fluorescently-labelled bacterial particles \((3 \times 10^7 \text{ Escherichia coli} \text{ and } 3 \times 10^7 \text{ Staphylococcus aureus}\) particles) purchased from Invitrogen (Carlsbad, CA) in a 250 μl total volume of bacterial particles and simulated wound fluid.

**Cleansing therapy: NPWTi**

After inoculation with bacterial particles, NPWTi was applied to the model’s simulated sacral pressure ulcer. Two NPWTi dressings were evaluated: ROCF-V and ROCF-VC (V.A.C. VeraFlo Cleanse™ Dressing; KCI USA, Inc.). The experiment consisted of five cycles with each cycle comprised of 20 minutes of negative pressure at \(-125\) mmHg followed by instillation of normal saline solution and a 60 seconds soak time. Three replicates of the experiment were performed for each experimental (dressing) condition. The fluid collected in the NPWTi canisters was also quantified for bacterial particles.

**Cleansing therapy: low-pressure lavage**

After inoculation with bacterial particles, the simulated sacral pressure ulcer was irrigated with commonly used, commercially available wound cleansers or the low-pressure pulsed lavage device. The cleansers used were: normal saline solution and antiseptic wound cleansers as per manufacturer’s instructions for use. The wound cleansers deliver pressure in the optimal range of 4–15 psi depending on how the cleansers are packaged (pressurised can for the normal saline solution and spray bottles for two different antiseptic wound cleansers). Each wound cleanser was applied to independent simulated wound beds for approximately 1 minute. Normal saline (1000 ml) was used as the irrigant for the pulsed lavage device. Three replicates of the experiment were performed for each of the four low-pressure lavage treatments.

**Bacterial particle aerosolisation evaluation: fluorescence analysis**

After the cleansing treatments, the amount of fluorescent bacterial particles collected on the microplates from bacterial aerosolisation or splatter was quantified. The collector microplates were analysed with the Beckman Coulter DTX-880 Multimode Detector spectrophotometer (Fullerton, CA) using the Beckman Coulter Multimode Analysis Software. The sensitivity of the multimode detector used was 20 fmol fluorescein/200 μl of liquid sample. A standard curve was used to convert the fluorescence signal to number of particles in each plate. The cumulative number of particles in the inner and outer plates, respectively, was each normalised to the initial quantity of particulate inoculated in the simulated wound prior to treatment initiation.

**Statistics**

The data from the ex vivo study were qualitative in nature and, hence, were not statistically evaluated. Paired statistical analyses were used to analyse the data from the in vivo study. Statistical analyses were performed using JMP 7.0 software (SAS, Cary, NC). A distribution test was first performed to confirm that the data followed a normal distribution followed by the parametric paired \(t\)-test.

The data from the cross-contamination study were determined to have non-normal distribution. Hence, non-parametric analysis of variance (Kruskal–Wallis) was used to test for differences in bacterial particle recovery among the five groups. If differences were detected, the Tukey test was used to compare means between treatments. For all statistical analyses, \(\alpha\) was set at 0.05.

**Results**

**Soft tissue damage in ex vivo wound model**

Digital photography and SEM images provided a qualitative evaluation of the occurrence of soft tissue trauma caused by the cleansing treatments investigated in this study. A visual assessment of the wound surface before and after treatment showed no observable blanching with NPWTi treatment (Figure 3), whereas blanching was illustrated via digital images in both low- and high-pressure lavage treatments (Figure 3B and C). Compared with tissue that was not treated and NPWTi-treated samples (Figure 4A and B), SEM images...
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Debris cleansing and tissue trauma with porcine animal model

Wound examination – visual

During the course of establishing a lightly granulated wound bed, from day 0 to day 4, the wounds healed with normal progression. Mild erythema was observed only in few wounds after inoculation with dextran. Also, compared with before treatment (Figure 5A and C), no observable blanching of the wound bed was observed in NPWTi-treated samples (Figure 5B), whereas 10 of the 12 wounds that received low-pressure lavage treatment showed evidence of blanching (Figure 5D). No other differences were noted.

Cleansing evaluation

For the estimation of cleansing efficacy, the FITC-dextran signal was measured in fluorescence images collected of the wounds before and after cleansing (Figure 6). There was no statistically significant difference in fluorescence signal between treatment groups before introduction of the FITC-dextran and before cleansing therapy ($P = 0.5679$). Both therapies were effective at cleansing FITC-dextran from the wounds, significantly reducing the fluorescence signal by 99.2% (NPWTi, $P < 0.0001$) and 95.2% (low-pressure lavage, $P < 0.0001$) (Figure 7). The percent reduction in fluorescence of low-pressure lavage-treated wounds was less than that of NPWTi-treated wounds ($P = 0.0009$) (Table 1); however, these results must be interpreted with the caveat that lavage-induced blanching of the wound may cause an artificial increase in fluorescent signal unrelated to the amount of FITC-dextran present.

Tissue damage/swelling evaluation

Tissue damage was evaluated by measuring immediate tissue swelling via changes in wound volume and depth after cleansing treatments. Three-dimensional imaging analyses suggest that NPWTi-treated wounds exhibited a decrease in swelling (swelling ratio < 1), whereas low-pressure lavage-treated wounds exhibited an increase in swelling (swelling ratio > 1) (Figure 8). The difference in swelling between NPWTi and low-pressure lavage was significant when using either wound volume ($P = 0.0086$) or wound depth ($P = 0.0006$) as the basis for the determination of swelling.

Oedema evaluation

The punch biopsies collected for histology were used to qualitatively evaluate tissue damage, such as oedema, caused by the cleansing techniques. Histological scoring indicates minimal
Figure 5 Observed blanching of the porcine wound bed. (A) Before negative pressure wound therapy with instillation (NPWTi) treatment. (B) Post NPWTi treatment. (C) Before low-pressure lavage treatment. (D) Post low-pressure lavage treatment with blanched areas circled.

Figure 6 Fluorescence signal of porcine wound bed images before and after cleansing treatments. (A) Before negative pressure wound therapy with instillation (NPWTi) treatment with the wound circled for clarification. (B) Post NPWTi. (C) Before low-pressure lavage. (D) Post low-pressure lavage.

to moderate oedema of the wound resulting from either technique, but there was no significant difference between cleansing regimens (data not shown). However, it should be noted that determination of oedema via histology can be challenging because histological processing can induce artifacts that mask oedema.

Environmental cross-contamination
Fluorescence spectrometry quantified bacterial cross-contamination levels at two distances from the simulated wound: 7.5 and 15 cm from the edges of the wound on the sacral pressure ulcer model. There was a statistically significant difference between NPWTi treatments and wound
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Discussion

There is abundant literature about the cleansing effectiveness of lavage. The question that many of the authors of these reports pose, though, is ‘does the cleansing capacity of lavage outweigh the damaging effects to the soft and bone tissue caused by this treatment?’ In short, is the trauma caused to the healthy tissues a clinically relevant concern? Many researchers have found the answer to be ‘yes’. Several studies have shown that although there is better cleansing with higher lavage pressures, the consequences include further dissemination of cleansing fluid into the interstices of the soft tissue, causing considerable gross and microscopic disruption that may lead to more cell death, higher infection rates and greater bacterial retention (8–10). In contrast, others have found that not only is a higher pressure lavage more detrimental to bone structure and to healing but it is also no more efficacious in removing foreign material than a low-pressure lavage cleansing regimen (6,7).

While animal models are routinely used in wound studies, an ex vivo wound model provides a cost-effective, readily available venue for performing experiments pertaining to wound healing aspects, such as cleansing, tissue trauma and bacterial penetration in soft tissue (8,10). Raw, porcine meat has also been used successfully in creating an in vitro wound model for showing the bacterial growth kinetics in a contaminated wound (22). Our study also incorporated the use of an ex vivo bench top wound model for preliminary assessment and screening of high-pressure lavage, low-pressure lavage and NPWTi to qualitatively assess tissue disruption after cleansing treatments.

The digital and SEM wound images of the ex vivo wound model successfully illustrate the differences in traumatic effects of wound cleansing; there was visually less tissue disruption in NPWTi-treated soft tissue compared with lavage-treated tissue. The degree of blanching also increased in proportion to the pressure of lavage treatment. In addition, the tissue trauma observed in the SEM images was in the form of ruptures and fissures in the tissue, and the intensity of the trauma could be visually assessed to be relative to the pressure of the irrigant used to cleanse the wound. Within a field, the number of cavities and/or grooves formed on the wound bed because of the cleansing treatments was clearly visible and damage to the tissues even more apparent with high-pressure lavage treatment. High-pressure lavage appeared to cause the cross-contamination: up to 29% of the bacterial particles were recovered at 7.5 cm away and up to 34% at 15 cm (Table 2). The total quantity of recovered particles (sum of those recovered from both distances) ranged from 50% to 60% for each inoculated bacterial strain (Figure 9). Cleansing with the pulsed lavage device resulted in visible splatter a great deal further than the 7.5 and 15 cm collection plates – no bacterial particles were detected or quantified in the collection plates placed at these distances. The total number of particles transferred from the model to the NPWTi canister was also quantified (Table 2). At least 95% of the bacterial particles were captured in the canister during the five cycles of NPWTi.

**Table 1** Percent fluorescence on porcine wound bed before and after cleansing treatments

<table>
<thead>
<tr>
<th>Cleansing treatment</th>
<th>Pre-cleansing</th>
<th>Post-cleansing</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPWTi</td>
<td>65.6 ± 5.6</td>
<td>0.9 ± 0.3</td>
</tr>
<tr>
<td>LPL</td>
<td>60.7 ± 6.3</td>
<td>4.9 ± 0.9</td>
</tr>
</tbody>
</table>

LPL, low-pressure lavage, data shown as mean ± SEM.
most tissue trauma, whereas NPWTi appeared to cause the least damage out of all the cleansing regimens.

In order for a wound to properly heal, the wound bed must be free of excessive exudate, eschar and foreign matter, which are mitigating factors that impair wound healing (1). Therefore, wound cleansing is an integral part of good wound bed preparation. While lavage has been the conventional method for wound cleansing, there have been more than 30 NPWTi reports published since the first clinical application of instillation therapy with NPWT described by Fleischmann et al. in 1998 (23). Many of these publications report clinical benefits of instillation therapy (24), including NPWTi’s wound cleansing attributes. For instance, in 2008, Gabriel et al. stated that, in their experience, NPWTi actively removes exudate and microscopic debris ‘to an extent that we have not experienced with other wound-care modalities (20)’. Furthermore, in an agar model study, Rycerz et al. demonstrated that, compared with continuous irrigation, NPWTi provided significantly better wound instillant distribution and coverage of not only the simulated wound bed but also tunnels and areas with undermining, thereby facilitating a more thorough cleansing (25). As noted by Wolvos, NPWTi also helped to change the viscosity of wound exudate, thereby enabling easier removal of drainage (18).

Because the ex vivo model demonstrated that high-pressure lavage caused substantial soft tissue trauma, a full-thickness acute wound healing model in normal healthy swine was used to assess the effectiveness of debris and contaminant removal by NPWTi in comparison to that of low-pressure lavage. A thin layer of granulation tissue was first established in the created wounds before the application of simulated debris. The molecular weight of the dextran used to simulate particulate debris was selected to imitate proteinaceous materials found in wound exudate. Wound cleansing via normal saline solution was used in both cleansing treatments, because saline minimised the risk of damaging the newly formed granulation tissue. The results of our study support the previous findings that NPWTi was as effective at cleansing as low-pressure lavage; both regimens cleaned >90% of the simulated debris.

As with many of the reported findings and our own ex vivo results, the in vivo study also resulted in considerable tissue trauma to the wound bed after lavage cleansing. There was an immediate increase in blanching and swelling (as measured by wound volume and depth) associated with low-pressure lavage. Upon gross examination of the wounds after cleansing treatments, it was noted that 83% of the wounds treated with low-pressure lavage exhibited blanching. There was no blanching or swelling detected in association with NPWTi. Histological analyses showed no significant difference in tissue disruption, oedema and haemorrhage between NPWTi and low-pressure lavage. This is not entirely unexpected,

Table 2  Percent detected bacterial particles recovered from concentric microplates after the simulated sacral pressure wound cleansing

<table>
<thead>
<tr>
<th>Method/Products</th>
<th>E. coli 7.5 cm</th>
<th>S. aureus 7.5 cm</th>
<th>E. coli 15 cm</th>
<th>S. aureus 15 cm</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPWTi w/ROCF-V</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>NPWTi w/ROCF-VC</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>LPL w/Pressurised can</td>
<td>20.8 ± 7.3</td>
<td>25.8 ± 10.6</td>
<td>20.1 ± 9.8</td>
<td>32.1 ± 1.4</td>
</tr>
<tr>
<td>LPL w/Spray bottle #1</td>
<td>28.3 ± 3.2</td>
<td>29.8 ± 6.2</td>
<td>16.5 ± 1.4</td>
<td>27.9 ± 4.2</td>
</tr>
<tr>
<td>LPL w/Spray bottle #2</td>
<td>22.5 ± 9.0</td>
<td>22.8 ± 7.2</td>
<td>34.0 ± 6.6</td>
<td>34.3 ± 5.8</td>
</tr>
</tbody>
</table>

LPL, low-pressure lavage; ROCF, reticulated open cell foam; data shown as means ± SEM.
because the process of fixation and further tissue processing per se can introduce dehydration artifacts in the samples.

In addition to the abundant literature about lavage’s cleaning capacity and tissue disruption propensities as a possible ‘side effect’, there are also numerous reports of infection control issues regarding lavage that contributed to HAIs (15,26). HAIs are a foremost concern in many health care settings, not only because of the increased morbidity and mortality associated with them but also because of the economic implications. The Centers for Disease Control and Prevention (CDC) estimates about 100 000 deaths from the 1.7 million HAIs per year in the USA alone, resulting in approximately $4.5 billion in additional health care costs annually (27,28). One of the ways the US government has responded in an attempt to curb the health care spending was in 2008, when Centers for Medicare and Medicaid Services (CMS) no longer reimbursed hospitals for conditions that were not present on admission; in short, hospitals had to pay for HAIs. Reed and Kemmerly stress in their report about HAIs that ‘these infections, hospitalisations, intangibles, such as grief and anxiety, and dollars spent are all preventable (27)’.

Reports of environmental contamination of A. baumannii associated with low-pressure lavage have been published within the past decade emphasising significant morbidity and expense (15,26). In contrast to these previously published results, Ho et al. concluded from their study that low-pressure pulsed lavage was not associated with an increased rate of cross-contamination of A. baumannii (29). The authors did state, though, that their findings may have underestimated the true potential for contamination. They also mentioned that although many health care facilities have instituted infection control measures that are adequate in limiting the spread of cross-contamination, less stringent infection control measures may be followed during standard wound care that could contribute to environmental cross-contamination.

The results of our study in comparing the susceptibility of bacterial cross-contamination between NPWTi and several low-pressure lavage cleansing regimens indicate that the lavage treatments using the cleaners caused significant cross-contamination, whereas NPWTi caused none. Approximately half of the bacterial particles inoculated after use of the cleaners were detected at the distances measured, and it may be inferred that the other half were spread even further than the distances at which the collection plates were placed. Cleansing with the low-pressure pulsed lavage device spread bacterial particles at even further distances than the other lavage treatments. This was of some surprise because the device included an attached splash shield and, most especially, because the device also had conjoined suction capability. These aerosolised particles can hypothetically contribute to an increased risk of HAIs by cross-contamination of different wounds on the patient and various surfaces of the health care setting. This is a concern especially in the case of immunocompromised patients, particularly if skin microlesions exist. This study also showed that nearly all of the inoculated particles were recovered in the canisters, confirming that cross-contamination is decreased with the use of NPWTi with all of the foams tested, while also demonstrating that bacteria were actively removed from the wound bed during the instillation process. The sealed environment virtually eliminated chances of bacterial aerosolisation while the therapy was applied.

There are some assumptions and limitations to this study. The assumptions are that the dextran would behave with the same migration characteristics as particulate debris and that the bacterial particles would aerosolise similarly to live bacteria when treated with the cleansing regimens. Some may view the use of the stage IV sacral pressure wound model for the bacterial cross-contamination portion of the study as the ‘worst-case’ scenario and may consider this as a limitation. This was intended so that a thorough cleansing of the tunnelling and undermining present on the model was supposed to have been challenging, at the very least. Despite a relatively small sample size, significant differences were apparent between the low-pressure lavage and NPWTi groups in all areas assessed regarding this area of investigation.

A future direction regarding tissue disruption resulting from this study is the assessment of bacterial penetration after repeated lavage cleansing in comparison to NPWTi. Considering that traditional gauze dressings can be changed up to several times a day and lavage cleansing performed alongside the dressing changes, it would be of great clinical interest to determine if bacteria penetrate into tissue that is repeatedly challenged with trauma and disruption. Also, the coauthors of this study underestimated the degree of environmental cross-contamination the low-pressure pulsed lavage cleansing can cause and splatter collection plates should have been placed at further distances. Measurement of the distance of aerosolisation and splatter is a definite area of interest and is vital in clinical relevancy.

On the basis of the data obtained from these studies, it can be concluded that NPWTi has cleansing capacity comparable to low-pressure lavage but without causing the tissue trauma and disruption associated with the latter treatment. In addition, wound cleansing via NPWTi proved to be a very effective way to decrease bacterial aerosolisation and, thus, potentially reduce the risk of environmental cross-contamination that could lead to HAIs.

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