FILLING THE VACUUM
Role of negative pressure wound therapy in open wound management in cats

Mirja C Nolff

Introduction

Treatment of wounds is a common activity in clinical veterinary practice. Although the majority of wounds are minor, and can be closed after initial treatment (ie, healing by primary intention), more challenging cases require appropriate open wound therapy for successful management.\(^1\)\(^-\)\(^3\) Modern veterinary open wound treatment encompasses a broad range of indications, including extensive acute trauma and polytrauma, chronic non-healing wounds, burns and surgical site or wound-associated infections.\(^4\)\(^-\)\(^10\) Unfortunately, there is currently no study available that reports the most common causes, or the incidence, of wounds necessitating open wound treatment in cats. In the author’s experience, cats requiring open wound treatment are commonly polytrauma patients. There may be a history of high velocity trauma (eg, road traffic accident) causing degloving injuries, open fractures and crush/tear injuries. Other cases involve blunt trauma leading to fat tissue necrosis; high-rise syndrome associated with open fractures and impalement injuries; or bite wounds, resulting in crush/shearing injuries. Infection (either secondary wound infection after minor injury/bite or surgical site infection) is another common indication for open wound treatment in the author’s experience. Complications such as multidrug-resistant wound (or systemic) infections with pathogens of the so-called ESKAPE group (Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa and Enterobacter species) are frequent, especially in patients pretreated with antibiotics.\(^7\)\(^-\)\(^11\)

Effective open wound management options no longer involve simply covering a wound until the body has healed of its own accord. Rather, they should actively promote the healing process and improve immune function to prevent – or treat – local or systemic infection.\(^4\)\(^,\)\(^12\)\(^-\)\(^15\)

Practical relevance: Open wounds and their treatment present a common challenge in veterinary practice. Approaching 15 years ago negative pressure wound therapy (NPWT) started to be incorporated into clinical veterinary medicine, and its availability is becoming more widespread in Europe and the USA. Use of this therapy has the potential to significantly increase the healing rate of open wounds as well as free skin grafts in small animals, and it has been occasionally described for the management of feline wounds.

Aim: This review describes the mechanisms of action of, and indications for, NPWT, and offers recommendations for NPWT specific to feline patients.

Evidence base: The information presented is based on the current evidence and the author’s clinical experience of the technique gained over the past 12 years. Comparative studies of different treatment options are lacking and, since wound healing in cats and dogs differs, cat-specific studies are especially needed. Well-designed wound healing studies comparing different advanced techniques will improve open wound healing in cats in the future, and potentially allow better understanding of the role of NPWT in this setting.

Keywords: Open wound treatment; cat wound healing; skin grafts; negative pressure wound therapy

Effective open wound management should actively promote the healing process and improve immune function to prevent, or treat, local or systemic infection.
Basics of wound healing and special considerations in cats

Wound healing can be divided into three phases: inflammation, proliferation and maturation.15–17 Although these healing phases apply to all wounds, they are especially relevant to consider in open wound management, as treatment decisions can actively support or suppress them.1 For example, sufficient proliferation cannot happen if the inflammatory phase is not completed,3,4,13 and prolonged subinflammatory states with poor peaks of inflammation at the beginning of healing can be linked to poor granulation tissue formation and healing.18,19 The phase of proliferation starts with neovascularisation, followed by proliferation of fibroblasts and formation of myofibroblasts, collagen synthesis and subsequently epithelialisation and contraction.15,20,21 Contraction and epithelialisation happen simultaneously and to different extents depending on wound tension and location. Trunk wounds in cats, for example, heal to a large degree by contraction, but this effect is less important than epithelialisation at the distal extremities.9,22,23 Finally, the scar matures over several years.3,4,13

While the fundamental patterns and phases of healing are comparable in most species, there are major differences in the overall healing capacity, speed and the proportion of contraction and epithelialisation seen in individual species.22 This is of importance, especially when evaluating the clinical benefits of treatments, as different species might need different support of the species-specific weak points during healing.22 Unfortunately, studies establishing the precise mediators and progression of cellular events in wound healing in cats are not available. Bohling et al demonstrated that primary wound healing in cats differs substantially from that in dogs.24–26 While the inflammatory response in clean wounds in dogs is rapid and robust, inflammation in cats is weak.23 As a result, cats not only build up less granulation tissue, but when formed it is poorly vascularised and more fibrotic compared with that of dogs.24–26 In addition, healing in cats is relatively impaired in terms of rates of contraction (18% after 7 days in cats vs 41% in dogs) and epithelialisation (13% after day 14 in cats vs 44% in dogs).24–26 Finally, while resection of subcutaneous tissue slows healing in both species, this effect is much more pronounced in cats.26

These various factors have an important bearing on wound treatment techniques: it is not valid to assume that a technique that has proved superior in rats or pigs will also perform well in a cat.22 Given that veterinarians treat wounds commonly, this reveals a tremendous gap in the knowledge available to inform optimal decision-making in feline open wound management, as studies performed in the target species are lacking. Interestingly, among the few papers published on open wound healing in cats, negative pressure wound therapy (NPWT) is one of the most frequently mentioned techniques (Table 1).7,27–34

### Table 1 Overview of the current literature reporting on open wound management in cats

| Study                  | Publication type | Indication(s) for open wound management | Wound healing product(s)/technique | Outcome                                                                 |
|------------------------|------------------|----------------------------------------|-----------------------------------|-------------------------------------------------------------------------|
| Gemignani et al (2017)27 | Case report      | Unknown, contaminated wound            | Platelet-rich plasma and wet-to-dry bandage | Wound healed in 20 days                                                 |
| Nolff et al (2017)7    | Retrospective match-controlled clinical study (n = 20) | Trauma, infection (SSI), fat tissue necrosis | NPWT vs polymer foam           | Significantly faster closure under NPWT (mean 25.8 days [range 11–57]) than under foam (mean 39.5 days [range 28–75]) |
| Tsioli et al (2016)28   | Experimental controlled study (n = 10)                  | Not applicable                       | Hydrocolloid vs semiocclusive pad                       | No differences in planimetry; more oedema with hydrocolloid           |
| Nolff and Meyer-Lindenberg (2015)29 | Case series (n = 6) | Trauma                               | NPWT followed by polyurethane foam and NPWT-augmented skin grafting | Mean duration of open wound management was 21 days (range 3–43); Mean graft take rate was 97% (range 80–100%; 100% in 7/10 grafts) |
| Nolff and Meyer-Lindenberg (2015)29 | Case report       | Necrotising fasciitis                 | NPWT                               | Wound grafted after 29 days; 100% graft take                             |
| Jordan et al (2012)31  | Case report      | Peristomal urine-induced tissue necrosis | Polymer foam followed by NPWT       | Vancomycin-resistant Enterococcus faecalis infection under open therapy with foam for 5 days. Switched to NPWT for 9 days. Wound closed with flank skin fold flap |
| Owen et al (2009)32    | Case report      | Urine-induced skin and muscle necrosis | NPWT                               | 40.3% wound contraction and granulation after 8 days. Wound closed with flap |
| Guille et al (2007)33  | Case report      | Traumatic wound (RTA)                 | NPWT                               | Wound successfully grafted (NPWT assisted) 40 days after trauma          |
| Siegfried et al (2004)34 | Case series (n = 5) | Four traumatic wounds (degloving), one open treatment after sarcoma resection | Wet-to-dry bandages followed by skin grafts | Mean duration of open wound management until stable granulation achieved was 14 days (range 7–21); Mean graft take rate was 94% (range 90–100%) |

NPWT = negative pressure wound therapy; SSI = surgical site infection; RTA = road traffic accident
The presence of dressings and requirement for frequent dressing changes can cause substantial stress for feline patients, and this is an important consideration when choosing the right treatment regimen. Dressing changes should be performed in a quiet environment and with the least restraint possible. In cats with severe wounds, sedation and adequate procedural analgesia are mandatory for potentially painful dressing changes. Treatment options that require less frequent dressing changes owing to improved exudate management should be favoured over standard bandages that require daily or even more frequent changes.

**General principles of open wound treatment**

The three principal pillars of open wound treatment are debridement, lavage and dressing.4,12,17

**Debridement**

Debridement is needed to remove all necrotic tissue from the wound, thus helping the body to transition efficiently from the inflammatory phase of wound healing to the proliferative phase.13,15,35,36 If debridement is not carried out, or is performed poorly, necrotic tissue keeps the wound in an inflammatory state and provides a surface for biofilm formation.17

**Lavage**

Lavage is the process of decreasing the bioburden and cleaning the wound. Different lavage solutions are available and, once again, the evidence for ‘which to use in which indication’ in the veterinary literature is sparse,37,38 and also somewhat contentious in the light of recommendations in human medicine.39 In general, and based on current human guidelines, saline or lactated Ringer’s solution can be used in all settings (even when pockets are present or joints, body cavities or nerves might be involved),37,40 as long as the wound is not infected (ie, for contaminated or colonised wounds [Table 2]).

Where there is local infection, wound antiseptics may be warranted.39,40 In veterinary medicine, use of chlorhexidine 0.05% has traditionally been advocated, based on two experimental studies dating back to 1988 (Sanchez et al37 [iodine vs chlorhexidine, n = 6 dogs]) and 1992 (Lozier et al38 [chlorhexidine vs lactated Ringer’s solution, n = 6 dogs]). Since then, several new wound antiseptics have been released and are now recommended in human medicine due to improved performance and fewer issues with resistance formation compared with chlorhexidine.39,41 However, investigations of these new compounds in veterinary medicine are rare. A recent blinded clinically controlled study of dog bite wounds underlines the powerful effect of saline alone, but also demonstrates the superior performance of polyhexanide compared with saline;40 polyhexanide is a biguanide (part of the same pharmaceutical family as chlorhexidine), and has shown promising performance as a wound antiseptic.39,42,43 Feline studies and/or comparative studies in dogs or cats comparing chlorhexidine and new wound antiseptics are lacking, and should be a focus of future research in order to evaluate if current recommendations regarding chlorhexidine are still valid. The author’s preference is polyhexanide biguanide based on the evidence in human medicine.

**Choice of dressing**

The job of the dressing is to impart oncotic and thermal stability and keep the wound moist, while creating an environment that supports the cellular events of healing.12–15,17 This is mainly achieved by using ‘interactive’ wound dressings, which alter the wound environment and interact with the wound surface. A study in dogs was able to demonstrate that minor wounds healed faster under interactive dressings (polymer foam) than when passive dressings (silicone gauze and

**Table 2: Definition of wound bacterial status and recommendations for treatment**

| Bacterial status                     | Treatment recommendations          |
|--------------------------------------|-----------------------------------|
| Contamination                       | Debridement; potentially lavage   |
| Colonisation                         | Debridement; potentially lavage   |
| Critical contamination               | Debridement, lavage, antiseptics  |
| Infection                            | Debridement, antiseptics ± antibiotics |
| Biofilm                              | Debridement; no other options for effective treatment |

Based on recommendations by Kramer et al39
bandage) were used. There is a plethora of modern wound dressings available, but unfortunately there is insufficient data on the application of these in dogs and cats to make evidence-based recommendations. This becomes even more problematic given what little is known about the specifics of feline open wound care.

So far, only two studies are available that have investigated the effect of different dressings in cats. In a controlled experimental study in 10 cats, Tsioli et al compared healing of 2 × 2 cm wounds on the dorsum under hydrocolloid or semiocclusive cotton-polyester pads. These investigators were not able to demonstrate differences in healing rate; however, wounds treated with hydrocolloid showed more oedema than control wounds. In a retrospective clinical study comparing NPWT (n = 10 cats) and polyurethane foam dressings (n = 10 cats) for open wound management, the author’s own group was able to document that wounds treated with NPWT had a significantly shorter time to closure than wounds managed with an interactive dressing. Besides these two studies, all other published information on open wound care in cats consists of case reports or small non-controlled case series.

Owing to the lack of evidence, the selection of an ‘appropriate’ wound dressing is often based on personal preference and historical choices, rather than on evidence of effectiveness in the species in question. Nevertheless, among the many different treatment options, NPWT (an active dressing system) has proven especially interesting in treating complex wounds. Despite this form of therapy having until recently only been available in selected veterinary centres in Europe and the USA, there are now more experimental and clinical studies available regarding its efficacy in dogs than for any other open wound management technique. In cats, there is only one clinical study to date, but seven out of the nine available publications on wound care in cats, including case reports and case series, have involved NPWT.

The following sections of this review describe the technique, as well as the mode of action of NPWT, and seek to highlight indications for NPWT in open wound management in cats.

**NPWT: a cat-friendly technique**

NPWT is well tolerated and keeps the cat dry and the wound stable. Although the cat will have to be kept in a crate or cage, the therapy can be performed at home in stable patients. This, together with the fact that NPWT enables a significant reduction in the number of dressing changes (even in highly exudative wounds), relieves a lot of stress during treatment.

---

**The main indications for negative pressure wound therapy (NPWT) are acute traumatic, chronic or infected wounds requiring open wound treatment.**

**NPWT for open wound management**

The main indications for NPWT in people, dogs and also cats are acute traumatic, chronic or infected wounds requiring open wound treatment. The system is especially effective with regard to infection control and for wound bed preparation prior to reconstruction of large defects. This is an important consideration, as surgical site infections or acute infected wounds causing local tissue damage and sepsis represent a challenging situation.

**Mode of action**

The mode of action of NPWT is not yet completely understood, but there are different proven effects (mostly in mice and pigs or ex vivo) that lead to improved wound healing. The first (simple) effect is that the wound is protected against thermal and oncotic fluctuations (thus preventing fluid evaporation and scab formation) and is kept moist.

A more specific effect of the vacuum is that three-dimensional contraction of the wound is stimulated, thereby bringing the wound edges together, while antagonising the physiological mechanisms leading to skin retraction (macrodeformation). This contraction effect has been demonstrated in studies in pigs, as well as in dogs, and substantially aids secondary closure after NPWT in dogs. However, it is strongly dependent on wound geometry. While NPWT greatly increases contraction in deep, three-dimensional wounds, such as those involving the trunk (where the foam and vacuum are centred deep within the wound), this effect is not seen in superficial skin wounds at the extremities or very flat wounds with just skin loss (where the foam and vacuum are placed on top of the wound).

In addition, the vacuum decreases interstitial pressure, leading to decreased wound oedema and improved transport of exudate away from the wound. Owing to the interstitial pressure gradient, a mechanical deformation of the fibroblasts and collagen network occurs, inducing ion flux. The cells at the tissue/foam border undergo microdeformation, leading to mechanical stress (5–20% mechanical stress) that triggers ion flux mechanotransduction and cell stimulation. Recent studies in pigs and dogs have
even shown that NPWT modulates mediators of wound healing, increasing the concentration of cytokines (interleukin [IL]-10, IL-6, IL-8, vascular endothelial growth factor [VEGF] and fibroblast growth factor [FGF]-2)\textsuperscript{47,58,59} and decreasing the matrix-metalloproteinase (MMP) concentration (MMP-9, MMP-1 and MMP-13).\textsuperscript{60,61} The net effect of these processes is inhibition of apoptosis, increased intracellular signal transduction, changes in gene expression and, ultimately, increased cell proliferation and granulation.\textsuperscript{57} Depending on the wound, species and study design, increases in granulation rate of between 60% and 200% have been documented.\textsuperscript{58,62} Finally, NPWT increases tissue perfusion and neovascularisation, and speeds up tissue organisation and maturation within the wound.\textsuperscript{53,63,64}

**Technique**

Before application of the NPWT dressing (Figure 1), the wound needs to be debrided and lavaged. The area around the wound is generously clipped (wet shaving is not recommended due to risk of skin trauma), and the skin surrounding the wound is completely dried. Skin protective adhesives can be used to help fix the system in place (eg, Cavilon Skin Protective [3M] or Opsite [Smith and Nephew]). For challenging areas, the use of stoma rings or paste applied around the wound edges on the skin (eg, Brava series [Coloplast]) can help to seal the dressing. A piece of foam is cut to match the size of the defect and placed within the wound (avoiding overlap with the skin). It is important to ensure good contact between the foam and all areas of the wound. Finally, the foam is secured to the wound by applying NPWT foil. This is best achieved by using small overlapping pieces of the sticky foil, rather than...
As noted earlier, studies investigating open wound care in cats are rare, but among the body of publications, NPWT is the most frequently described technique. The author’s group performed a retrospective controlled clinical trial in 2017 to evaluate the effect of NPWT for open wound care in cats, and was able to demonstrate that NPWT led to significantly faster wound closure compared with use of an interactive dressing alone. In addition, NPWT was especially effective in terms of infection control and management of soft tissue necrosis. Given the known deficiencies of wound healing in cats (weak inflammatory response, impaired granulation, poor vascularisation and poor contraction), NPWT offers a number of benefits. Studies have shown that the device manages infection effectively, increases granulation and neovascularisation, and also aids wound contraction by active macro-contracture. The author’s clinical experience with cats treated by NPWT (in addition to the above-mentioned study cats), as well as the published case reports documenting treatment of conditions including major skin loss, urine-induced necrosis or malignant soft tissue infection such as necrotising fasciitis, strongly supports the beneficial effect of this system. Nevertheless, we need more well-designed wound healing studies comparing different advanced techniques to improve open wound healing in cats in the future, and potentially gain a better understanding of the underlying effects of NPWT in this setting.

The author’s clinic currently offers NPWT in cases requiring open wound care for large defects, severe de-gloving injuries involving bones and/or tendons (which provide a poor base for granulation) or for the management of wound infection. In the author’s opinion, NPWT is of most value in this last scenario. While the precise mechanisms are not clear, increased wound perfusion and mediator modulation are most likely involved. Even with severely infected wounds, and where other attempts (multiple antibiotics, repeated debridement and other forms of open management) have failed previously and led to worsening local and systemic infection (sepsis), the author and her colleagues have been able to implement successful treatment using NPWT. Surgical site infections are a very important indication for NPWT in cats. In such cases (frequently wounds that could be closed without reconstruction) the device is used until the wound infection and the cat’s general condition have stabilised, and the wound can then be closed.

Unfortunately, there are no concrete data to inform how long treatment should be continued in these cases. The author’s rule of thumb is to wait until the patient’s systemic status is improved and the wound is stable (i.e., with a steady reduction in exudation), does not need any further debridement, and displays well-vascularised granulation tissue. At this point, a further round of NPWT (3 days) is undertaken to prevent premature closure, and the wound is then closed. In general, and as more experience is gained, the trend is towards decreasing the duration of NPWT, with the average treatment comprising between three and five dressing changes over 9–15 days. Severe infection and fat tissue necrosis might require a longer treatment duration, however.

NPWT for skin graft augmentation

One of the earliest applications of NPWT in human medicine, besides open wound care, was skin graft augmentation, as recorded in a literature search by Azzopardi et al, this dates back 23 years. Indeed, in the first published case of NPWT usage in a cat, in 2007, the technique was applied to a grafted wound. NPWT facilitates graft fixation and encourages graft ‘take’. This effect becomes especially important for complex three-dimensional wounds that generally do not allow secure graft fixation by any other method. Increased graft take rates under NPWT compared with controls (standard bolster dressing) have been documented for humans and dogs. Mode of action

The mechanisms responsible for improved graft healing under NPWT can be divided into two broad categories:

- Active stimulation of cellular proliferation, revascularisation and microcirculation. A demonstration of these effects was provided by Saaiq et al’s 2010 study, which showed that pretreatment of the recipient bed using NPWT increased graft take in human patients.

NPWT can be used for skin graft augmentation, facilitating graft fixation and encouraging graft ‘take’.

Continued on page 829
Continued from page 828

Finally, it is important to note that epithelialisation does not occur under NPWT. So, once a sound granulation bed is achieved and infection is controlled (irrespective of potential residual bacterial contamination), wounds either have to be reconstructed or treatment has to be changed to dressings that facilitate epithelialisation (such as polymer foams).

Figures 3 and 4 show examples of the use of NPWT for management of traumatic wounds in cats.

Figure 3 (a) This cat was presented with an acute traumatic wound of unknown origin. (b) The full extent of the injury was evident after the area around the wound had been clipped and washed. (c) Dead areas of skin and foreign material were removed by surgical debridement and (d) negative pressure wound therapy was initiated. (e) At day 5 after initial presentation, a healthy granulation bed was evident and (f) the wound was reconstructed using a caudal epigastric flap. Healing proceeded uneventfully, apart from some minor suture dehiscence.

Figure 4 (a,b) Bilateral shearing injury in a cat, which underwent open wound management with negative pressure wound therapy (NPWT) for a total of 12 days. (c) The NPWT dressings were secured with soft bandages, and the cat tolerated treatment well. (d,e) Formation of granulation tissue had started by day 7. The cat subsequently underwent skin grafting (see Figure 5).

Avoidable complications, such as graft lift-off and fluid accumulation beneath the grafted skin, and neutralisation of shear forces. In contrast to conventional dressings, NPWT allows uniform pressure distribution across the entire graft surface, and thus is effective at neutralising shear forces.65–68

NPWT-augmented skin transplants exhibit improved healing, based on histological evaluation, as early as 3 days after grafting, when compared with controls (standard bolster dressing).70 Becker et al reported faster revascularisation and improved take under NPWT, although the researchers also found that oxygen tension within NPWT grafts was inferior to controls during the first 3 days.67

Technique

After skin transplantation is completed, the graft is covered with a protective, non-adhesive silicone dressing.77 Note that NPWT foam should never be placed in direct contact with the skin, since this can cause skin maceration and risk graft lift-off during dressing changes. The pre-cut foam is placed upon non-adhesive gauze and secured as described earlier for standard NPWT dressings. Both grey (polyurethane) and white (polyvinyl alcohol) foam can be used for graft augmentation, although the author favours grey foam, owing to its greater flexibility, at a pressure setting of ~125 mmHg. Some difference of opinion exists regarding...
NPWT for skin graft augmentation in cats

Reported indications for skin grafting in cats have included shearing/degloving injuries, iatrogenic bandage sloughing, crush injury and necrotising fasciitis, as well as wounds of unknown origin. In general, success rates for full-thickness free skin grafts in cats have been higher than in dogs. A first case series published in 2004 by Siegfried et al documented a take rate of 94% (five cats, range 90–100% take) with standard grafting and rigid post-graft bandaging (splint for 7–21 days followed by soft bandage for 7–21 days). In 2015, Riggs et al reported success rates of 77% for grafts placed in cats compared with 38% in dogs, with grafts applied to the antebrachium being especially challenging. The duration of post-graft bandage therapy ranged from 10 days to 4 weeks.

The only case series on NPWT-assisted skin grafting in cats (six cats, 10 grafts) reported a success rate of 97% (range 80–100%, with 7/10 grafts having 100% take); none of the cats needed a bandage for more than 10 days after grafting. Given the evidence in dogs and humans, it seems likely that NPWT augmentation leads to improved graft incorporation compared with the use of standard dressings in cats as well. However, as no direct comparative studies are available, this remains to be proven. Based on the author’s experience, two of the main advantages of NPWT-augmented grafts are the fact that no rigid splints are needed after grafting, and that the time a bandage is generally needed can be substantially decreased.

A sequence of images illustrating the use of NPWT for skin grafting in a cat, following open wound therapy for management of bilateral shear injuries, is shown in Figure 5.

NPWT for treatment of septic peritonitis

Another potential application of NPWT is in the treatment of septic peritonitis. In the human medical field, use of negative pressure has greatly reduced morbidity and mortality in patients managed with an open abdomen, and an initial veterinary case series has documented the use of NPWT to manage open abdomens in dogs and cats as well. Although the precise mode of action has yet to be determined, NPWT appears to be safe in these patients, and improves management of exudate, while keeping patients mobile, dry and comfortable. The device can either be placed directly within the abdominal cavity.
Although the majority of wounds are minor, and can be closed after initial treatment, more challenging cases require appropriate open wound therapy for successful management. There is a considerable gap, however, in existing knowledge of open wound care in cats.

While there are few papers published on open wound healing in cats, NPWT is one of the most frequently mentioned techniques, and its application for open wound treatment, as well as graft augmentation and indications such as septic peritonitis, seems promising.

Further controlled studies are required to investigate the full potential of NPWT in feline patients.

NPWT appears to be safe in patients with septic peritonitis, improving management of exudate, while keeping them mobile, dry and comfortable.

**Conflict of interest**

The author declared no potential conflicts of interest with respect to the research, authorship and/or publication of this article.

**Funding**

The author received no financial support for the research, authorship and/or publication of this article.
References

1. Balsa IM and Culp WT. Wound care. Vet Clin North Am Small Anim Pract 2015; 45: 1049–1065.
2. Corr S. Intensive, extensive, expensive. Management of distal limb shearing injuries in cats. J Feline Med Surg 2009; 11: 747–757.
3. Nolff MC. Wundmanagement. In: Modernes Wundmanagement bei Hund und Katze. Stuttgart: Thieme, 2020, pp 86–113.
4. Pavletic MM. Atlas of small animal wound management and reconstructive surgery. Hoboken, NJ: Wiley, 2018.
5. Davidson JR. Current concepts in wound management and wound healing products. Vet Clin North Am Small Anim Pract 2015; 45: 537–564.
6. Lascelles BD, Davison L, Dunning M, et al. Use of omental pedicle grafts in the management of non-healing axillary wounds in 10 cats. J Small Anim Pract 1998; 39: 475–480.
7. Nolff MC, Fehr M, Reese S, et al. Retrospective comparison of negative pressure wound therapy and silver-coated foam dressings in open-wound treatment in cats. J Feline Med Surg 2017; 19: 624–630.
8. Pope ER. Head and facial wounds in dogs and cats. Vet Clin North Am Small Anim Pract 2006; 36: 793–817.
9. Nolff MC, Albert R, Reese S, et al. Comparison of negative pressure wound therapy and silver-coated foam dressings in open wound treatment in dogs: a prospective controlled clinical trial. Vet Comp Orthop Traumatol 2018; 31: 229–238.
10. Nolff MC, Fehr M, Bolling A, et al. Negative pressure wound therapy, silver coated foam dressing and conventional bandages in open wound treatment in dogs. A retrospective comparison of 50 paired cases. Vet Comp Orthop Traumatol 2015; 28: 30–38.
11. Nolff MC, Reese S, Fehr M, et al. Assessment of wound bio-burden and prevalence of multi-drug resistant bacteria during open wound management. J Small Anim Pract 2016; 57: 255–259.
12. Fahie MA and Shettko T. Moisture and healing: beyond the jargon. Ostomy Wound Manage 2000; 46 1A Suppl: 51S–62S; quiz 3S-4S.
13. Gurtner GC, Werner S, Barrandon Y, et al. Wound repair and regeneration. Nature 2008; 453: 314–321.
14. Pavletic MM and Trout NJ. Bullet, bite, and burn wounds in dogs and cats. Vet Clin North Am Small Anim Pract 2006; 36: 873–893.
15. Nolff MC. Unterdrucktherapie. In: Modernes Wundmanagement bei Hund und Katze. Stuttgart: Thieme, 2020, pp 131–140.
16. Theorêt CL and Wilming JM. Abrant wound healing in the horse: naturally occurring conditions reminiscent of those observed in man. Wound Repair Regen 2013; 21: 365–371.
17. Wilming JM and van Weeren PR. Second-intention repair in the horse and pony and management of exuberant granulation tissue. Vet Clin North Am Equine Pract 2005; 21: 15–32.
18. Li B and Wang JH. Fibroblasts and myofibroblasts in wound healing: force generation and measurement. J Tissue Viability 2011; 20: 108–120.
19. Wong VW, Akaishi S, Longaker MT, et al. Pushing back: wound mechatronaduction in repair and regeneration. J Invest Dermatol 2011; 131: 2186–2196.
20. Volk SW and Bohling MW. Comparative wound healing – are the small animal veterinarian’s clinical patients an improved translational model for human wound healing research? Wound Repair Regen 2015; 23: 372–381.
21. Wilmink JM, Nederbragt H, van Weeren PR, et al. Differences in wound contraction between horses and ponies: the in vitro contraction capacity of fibroblasts. Equine Vet J 2001; 33: 499–505.
22. Bohling MW and Henderson RA. Differences in cutaneous wound healing between dogs and cats. Vet Clin North Am Small Anim Pract 2006; 36: 687–692.
23. Bohling MW, Henderson RA, Swaim SE, et al. Cutaneous wound healing in the cat: a macroscopic description and comparison with cutaneous wound healing in the dog. Vet Surg 2004; 33: 579–587.
24. Bohling MW, Henderson RA, Swaim SE, et al. Comparison of the role of the subcutaneous tissues in cutaneous wound healing in the dog and cat. Vet Surg 2006; 35: 3–14.
25. Gemignani F, Peruzzi A and Iacopetti I. Use of canine sourced platelet-rich plasma in a canine contaminated cutaneous wound. Canadian Vet J 2017; 58: 141–144.
26. Tiøli V, Gouletou SO, Galatos AD, et al. Effects of two occlusive, hydrocolloid dressings on healing of full-thickness skin wounds in cats. Vet Comp Orthop Traumatol 2016; 29: 298–305.
27. Nolff MC and Meyer-Lindenberg A. Negative pressure wound therapy augmented full-thickness free skin grafting in the cat: outcome in 10 grafts transferred to six cats. J Feline Med Surg 2015; 17: 1041–1048.
28. Nolff MC and Meyer-Lindenberg A. Necrotising fasciitis in a domestic shorthair cat – negative pressure wound therapy assisted debridement and reconstruction. J Small Anim Pract 2015; 56: 281–284.
29. Jordan CJ, Kulendra E, Perry KL, et al. Management of persistent tissue necrosis following prepubic urethrostomy in a cat. Vet Comp Orthop Traumatol 2012; 25: 433–437.
30. Owen L, Hobston-Moore A and Holt P. Vacuum-assisted wound closure following urine-induced skin and thigh muscle necrosis in a cat. Vet Comp Orthop Traumatol 2009; 22: 417–421.
31. Guille AE, Tseng LW and Orshey RJ. Use of vacuum-assisted closure for management of a large skin wound in a cat. J Am Vet Med Assoc 2007; 230: 1669–1673.
32. Siegfried R, Schmokel H, Rytz U, et al. Treatment of large distal extremity skin wounds with autogenous full-thickness mesh skin grafts in 5 cats. Schweiz Archiv Tierheilkd 2004; 146: 277–283.
33. Granick M, Boykin J, Gamelli R, et al. Toward a common language: surgical wound bed preparation and debridement. Wound Repair Regen 2006; 14 Suppl 1: S1–S10.
34. Granick MS, Tran BNN and Alvarez OM. Latest advances in wound debridement techniques. Surg Technol Int 2020; 36: 37–40.
35. Schaefer IR, Swaim SE, Nusbaum KE, et al. Effects of chlorhexidine diacetate and povidone-iodine on wound healing in dogs. Vet Surg 1988; 17: 291–295.
36. Lozier S, Pope E and Berg J. Effects of four preparations of 0.05% chlorhexidine diacetate on wound healing in dogs. Vet Surg 1992; 21: 107–112.
and morphological changes. Ann Plast Surg 2011; 66: 296–300.

57 Saxena V, Hwang CW, Huang S, et al. Vacuum-assisted closure: microdeformations of wounds and cell proliferation. Plast Reconstr Surg 2004; 114: 1086–1096; discussion 1097–1098.

58 Jacobs S, Simhaee DA, Marsano A, et al. Efficacy and mechanisms of vacuum-assisted closure (VAC) therapy in promoting wound healing: a rodent model. J Plast Reconstr Aesth Surg 2009; 62: 1331–1338.

59 Kilpadi DV, Bower CE, Reade CC, et al. Effect of vacuum assisted closure therapy on early systemic cytokine levels in a swine model. Wound Repair Regen 2006; 14: 210–215.

60 Greene AK, Puder M, Roy R, et al. Microdeformational wound therapy: effects on angiogenesis and matrix metalloproteinases in chronic wounds of 3 debilitated patients. Ann Plast Surg 2006; 56: 418–422.

61 Glass GE, Murphy GF, Esmaeili A, et al. Systematic review of molecular mechanism of action of negative-pressure wound therapy. Br J Surg 2014; 101: 1627–1636.

62 Morykwas MJ, Faler BJ, Pearce DJ, et al. Effects of varying levels of subatmospheric pressure on the rate of granulation tissue formation in experimental wounds in swine. Ann Plastic Surg 2001; 47: 547–551.

63 Ichiosa S, Watanabe H, Sekiya N, et al. A technique to visualize wound bed microcirculation and the acute effect of negative pressure. Wound Repair Regen 2008; 16: 460–465.

64 Wackenfors A, Gustafsson S, Sjogren J, et al. Blood flow responses in the peristernal thoracic wall during vacuum-assisted closure therapy. Ann Thorac Surg 2005; 79: 1724–1730; discussion 1730–1731.

65 Azzopardi EA, Boyce DE, Dickson WA, et al. Application of topical negative pressure (vacuum-assisted closure) to split-thickness skin grafts: a structured evidence-based review. Ann Plast Surg 2013; 70: 23–29.

66 Blume PA, Key JJ, Thakor P, et al. Retrospective evaluation of clinical outcomes in subjects with split-thickness skin graft: comparing VAC® therapy and conventional therapy in foot and ankle reconstructive surgeries. Int Wound J 2010; 7: 480–487.

67 Becker ST, Rennekampff HO, Alkatout I, et al. Comparison of vacuum and conventional wound dressings for full thickness skin grafts in the minipig model. Int J Oral Max Surg 2010; 39: 699–704.

68 Llanos S, Danilla S, Barraza C, et al. Effectiveness of negative pressure closure in the integration of split thickness skin grafts: a randomized, double-masked, controlled trial. Ann Surg 2006; 244: 700–705.

69 Saaq M, Hameed Ud D, Khan MI, et al. Vacuum-assisted closure therapy as a pretreatment for split thickness skin grafts. J Coll Physicians Surg Pak 2010; 20: 675–679.

70 Kim EK and Hong JP. Efficacy of negative pressure therapy to enhance take of 1-stage allogrums and a split-thickness graft. Ann Plast Surg 2007; 58: 536–540.

71 Riggs J, Jennings JL, Friend EJ, et al. Outcome of full-thickness skin grafts used to close skin defects involving the distal aspects of the limbs in cats and dogs: 52 cases (2005–2012). J Am Vet Med Assoc 2015; 247: 1042–1047.

72 Cioffi KM, Schmiedt CW, Cornell KK, et al. Retrospective evaluation of vacuum-assisted perineal drainage for the treatment of septic peritonitis in dogs and cats: 8 cases (2003–2010). J Vet Emerg Crit Care 2012; 22: 601–609.