





ORIGINAL ARTICLE

Mechanical and contact characteristics of foam materials within wound dressings: Theoretical and practical considerations in treatment

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Abstract

In the treatment of acute and chronic wounds, the clinical performance of a given foam-based dressing, and, ultimately, the wound healing and cost of care outcomes are strongly influenced by the mechanical performance of the foam material/s within that dressing. Most aspects of the mechanical performance of foam materials, for example, their stiffness, frictional properties, conformability, swelling characteristics and durability, and the overall mechanical protection provided by a foam-based dressing to a wound strongly depend on the microstructure of the foam components, particularly on their microtopography, density and porosity. This article, therefore, provides, for the first time, a comprehensive, self-inclusive compilation of clinically relevant theoretical and practical considerations, based on published analytical and experimental research as well as clinical experience related to the mechanical performance of foams in foam-based wound dressings. The current bioengineering information is useful for establishing understanding of the importance of mechanical properties of foams in foam-based dressings among clinicians and researchers in industry and academia, and other potential stakeholders in the wound care field, for example, regulators and buyers. This information is also particularly important for the development of standardised test methods for the evaluation of foam-based wound dressings and resulting standard mechanical performance metrics for these dressings.

KEYWORDS

foam dressing properties, bioengineering testing methods, stiffness and durability, strength of adhesion and adherence models, treatment of wounds

Abbreviations: COF, coefficient of friction; DFU, diabetic foot ulcer; ECMO, extracorporeal membrane oxygenation; MARS, medical adhesive-related skin injuries; PU/PI, pressure ulcer/injury; SEM, scanning electron microscopy; SF, shape factor; VLU, venous leg ulcer.

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Key Messages

- the mechanical performance of foams within dressings affects clinical outcomes
- many mechanical performance aspects of foams depend on their microstructure
- we compiled the relevant theoretical and practical considerations for foams
- this highlights the importance of mechanical properties of foams in dressings
- this information is also relevant to the development of standardised test methods

1 | INTRODUCTION: STRUCTURE-FUNCTION PRINCIPLES APPLIED TO DRESSINGS IN A MECHANICAL CONTEXT

Various materials are used innovatively in the making of modern wound dressings and each type has a unique microarchitecture and function that is determined by how they are composed, combined and constructed. How the dressings are affixed to the skin and wound, handled and removed, are the foundations for their clinical performance, contributing to the experience of the patients using the product and the cost-effectiveness of the treatment course.¹⁻⁸ Foam materials within dressings appear to be a particularly good case for demonstrating the above concept because of their popularity in clinical practice. The clinical outcomes of wound care are, of course, not only derived from the properties of the foam/s in the dressing per se, but the complexities of dressing designs also have a profound effect on dressing performance. Importantly, it is uncommon for a dressing alone to heal a chronic wound, and even the best dressings require a holistic approach that targets the underlying cause of the wound to achieve wound healing. However, the application of a dressing that is mismatched with the demands of the wound may make complete healing difficult, if not impossible. For example, the compressive and tensile stiffness and strength of a foam material within a dressing, affecting the ability of the foam-based dressing to protect the wound mechanically, and the bending stiffness of foams, which influences the conformability performance of foam-based dressings, all depend on the apparent density* and porosity of that foam material, which in turn, influence the absorbency performance of the foam while managing exudates, as demonstrated in the clinical case described in Figure 1. Likewise, the coefficient of friction (COF) of

the wound-facing surfaces of foam dressings is influenced by the microtopography of the surface. Hence foams used within wound dressings deserve specific attention in the literature. The interaction of these bioengineering factors and how they eventually shape the clinical outcome is currently unknown and was not thoroughly studied before. For example, in their Cochrane review work, Walker et al⁹ stated, specifically for foam dressings applied to treat a pressure ulcer/injury (PU/PI), that 'Clinicians need to carefully consider the lack of robust evidence in relation to the clinical and cost-effectiveness of foam dressings for treating PUs/Pis when making treatment decisions, particularly when considering the wound management properties that may be offered by each dressing type and the care context' (p. 2). The clinical authors of this article share this view and observe that many wound care specialists do not understand the differences between commercially available foam dressings, which likely leads to suboptimal care outcomes. The current article takes the first step in addressing these gaps, by identifying key characteristics of foams in wound dressings that should be studied and reported for standardisation, as suggested in clinical work.¹⁰

The well-established structure-function principle in material science provides insights into the properties of the microstructure (also often referred to as the microarchitecture) of the studied material that explain the resulting mechanical, thermal, fluid transport and retention metrics related to the function of each dressing.^{5,6,11-16} The characteristics of each of these material properties and their collective quality, together with the durability of the materials and sustained integrity of the dressing structure while in use under different real-world conditions and activities shape the clinical performance of wound dressings. Understanding material behaviours and how to quantitatively describe the material microarchitecture is therefore vital to achieve desirable clinical



FIGURE 1 A clinical case demonstrating the importance of the porosity property of foam dressings: A 56-year-old female after breast reconstruction developed a massive inflammatory response following trauma because of rejection of the surgical material. The highly viscous exudation provoked by the inflammation was misinterpreted by the clinical team to be related to infection. The exudate was, in fact, so viscous that regular absorbent and even superabsorbent dressings were not able to manage it. Only a high-specification foam dressing specifically characterised by large pores was able to absorb and retain the wound fluids without causing maceration of the perilesional skin. The photograph of the wounds and this case description were provided by clinical co-author PA

performance through appropriate design, processing and manufacturing of wound dressings.^{2,5} The purpose of this article is to review the contemporary knowledge and formulate the bioengineering theory that applies to porous materials in wound dressings, and primarily, to foams that are in extensive use in contemporary dressings.¹⁷ Foams in foam-based dressings are most commonly made of soft polyurethane, which is often treated in various ways to create a hydrophilic non-adherent membrane that allows the passage of exudate through to the insulating foam body of the dressing. Foams have many important advantages as a wound dressing material over traditional textile dressings (eg, cotton bleached gauze) or hydrocolloid dressings, in light of their superior fluid absorbency and lower risk to adhere to fragile wound bed tissues. Foams are also preferred for their mechanical cushioning properties, thermal insulation for the wound, flexibility to conform relatively well to curved body contours (and further mimic skin tones to minimise the negative effect of wounds on the body image, and therefore, improve compliance to the treatment plan), and maintenance of a moist wound healing environment. However, according to the aforementioned structure-function

principles, the quality of the clinical performance of a foam-based dressing in any of the above aspects stems from the microstructure and properties of the specific foam types used in the dressing.

It is noteworthy that most wound dressings contain different material types in addition to polymeric foams alone. Many foam-based dressings incorporate layers combining polymeric with cellulose or superabsorbent materials in multi-layered dressings.^{16,18-20} Since their invention in the 1950s, polymeric foams have received extensive attention from material scientists in numerous fields of engineering and technology (such as in the packing industry, seat and mattress industry, for sound management components and in building construction to mention a few), and analytical (mathematical) formulations describing their mechanical behaviour were developed in these diverse fields. Nevertheless, foams have not been carefully scrutinised in terms of their performance as a wound dressing material⁹ and there is no mathematical-analytical theory for foams in foam-based wound dressings in the literature yet. Moreover, the relatively simple structural and mechanical behaviours of foams with respect to other (such as hydrocolloid and superabsorbent) dressing materials allow closed-form mathematical-analytical formulations to be developed and utilised specifically for analyses of foams in wound dressings. Other wound dressing materials may require sophisticated computer codes and simulation tools for complex numerical computer analyses to achieve similar aims. *Foams are, therefore, a natural, solid starting point in the formulation of a rigorous scientific theory for the mechanical behaviour of wound dressings.*

The specific aims of this work are therefore: (a) to review, for the first time in the literature, the mechanics and contact characteristics of foam materials within foam-based or foam-made dressings; and (b) provide key theoretical and practical considerations, which are relevant to the design, efficacy research and development of new testing standards for foam-containing or foam-made wound dressings. As a first step towards the development of better, clinically relevant testing standards for foam-containing dressings, there is a need to understand and establish a scientific (bioengineering) theoretical framework around the importance of the mechanical properties of foams in dressings. This framework should ultimately take into consideration the perspectives of wound care researchers, clinicians, patients, manufacturers, regulators, payers and likely also other relevant stakeholders in the wound care arena such as professional societies, health economists and health policy makers. The current work is a cornerstone of this initiative.

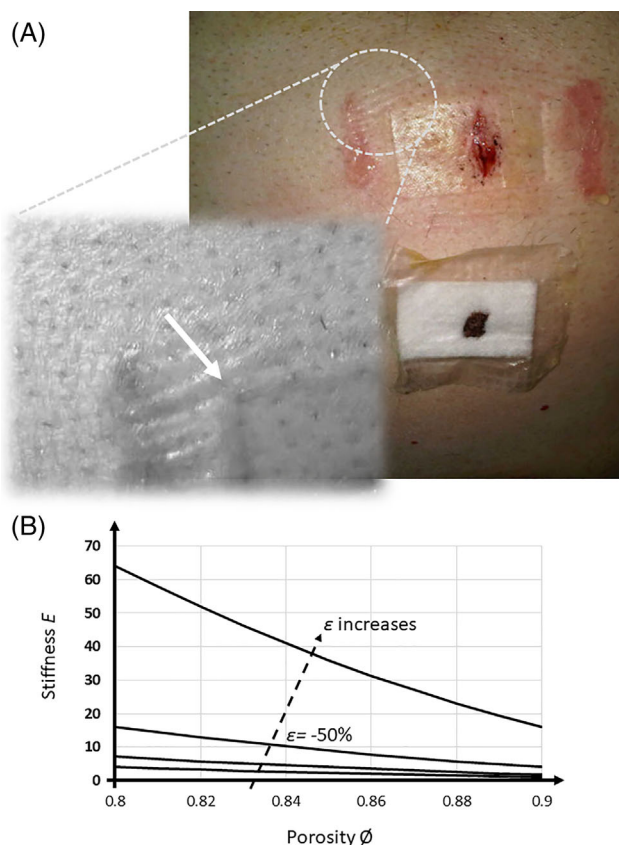


FIGURE 2 The potential implications of an overly stiff foam-made dressing: (A) an indentation mark caused by a foam dressing (marked by a white arrow; reproduced from Sanusi²¹ who reported this phenomenon around laparoscopic port-site wounds); (B) the theoretical effects of the porosity and deformation levels of a foam (with the same solid polymer phase) on its stiffness. The parameters of Equation (A8) were set as follows: $E_0 = 100$ kPa, $B = 2$, and the strain level ϵ was increased from zero (ie, no deformation) to 75% strain at 25% intervals. The non-linear increase in stiffness of the foam-made dressing E with the decrease in the porosity \emptyset and increase in the strain level ϵ is plotted, demonstrating that the influence of the strain level is more dominant (as strains effectively decrease the porosity by narrowing and collapsing the air voids). Of note, a decreased stiffness with an increased porosity, as shown here, is expected as long as the compared foams are made of the same solid polymer material. Different solid polymer materials with different porosities can theoretically obtain the same compressive stiffness

2 | THE STIFFNESS OF FOAMS AND HOW THEY RELATE TO FOAM DENSITY AND POROSITY

Foams in wound dressings are composed of a solid polymer phase and a dispersed air phase. In addition to offering storage space for excess exudate in the voids of the foam, from a mechanical perspective, foams are lightweight and have the ability to absorb mechanical energy

through deformation, which promotes mechanical protection and cushioning capabilities. Foams also facilitate good thermal insulation of the wound bed against heat loss to the ambient environment in the absence of native skin. Foam dressings in the wound care industry can theoretically be designed to optimise these mechanical properties, for example, balancing between stiffness and conformability determined by the foam density. Ideally, a foam dressing should have a stiffness level that is similar to that of native skin, to avoid sharp stiffness gradients between the dressing and peri-wound skin which may imprint the skin, that is, create localised, sustained soft-tissue distortions under the dressing and particularly at the perimeter or borders of the applied dressing, which may compromise cell and tissue viability over time (Figure 2A).^{5,22,23} Concurrently with this requirement, the borders of a foam dressing should have the capability to prevent any leakage of exudate to the peri-wound skin, to prevent inflammation or maceration and subsequent skin breakdown.^{24,25} McKee et al²⁶ reviewed the range of stiffnesses reported for human skin evaluated by means of indentation testing, which is the relevant loading mode to the interaction with dressings. They noted a wide range of reported skin stiffness data and decided to conduct a statistical outlier analysis of the reported empirical data. They concluded that the elastic moduli of human skin are within the 6 to 222 kPa range, averaging at approximately 85 kPa. Overly stiff dressings that substantially exceed the aforementioned skin stiffness can be an outcome indicator of an inadequate design and/or material selection (ie, excessively low porosity). More commonly, however, a foam dressing with a suitable stiffness under no mechanical strain or moderate strain can become exceedingly stiff if it is compressed to produce strains that exceed the strain domain for the intended use. For example, excessive dressing strain may develop under compression bandaging or medical devices, or when subjected to the bodyweight as in a non-offloaded diabetes-related foot ulcer (DFU).^{5,22,23,27-29} Orlov and Gefen^{23,30} recently reported that the compressive stiffness of certain foam dressings compressed to 50% strain can reach approximately 300 kPa, that is, ~ 3.5 times the mean compressive skin stiffness reported by McKee et al,²⁶ which indicates that currently, the biomechanical compatibility of foam dressings with skin is not always optimal.²³

The chemistry of the raw foam material and the details of the manufacturing process of foaming eventually determine the strength and stiffness of the final foam material. In particular, these factors influence both the properties of the solid polymer phase and the microarchitectural characteristics of the porosity of the foam, that is, the percentage and shape of the microscopic void spaces

in the microstructure designed to collect and store the wound exudate when the foam is included in a wound dressing (please see Figure 1 for a clinical case description demonstrating the importance of this porosity property). The denser a foam is, the less air that it contains in its pores, and so the work of compressive forces will mostly be invested in deforming the solid polymer, that is, the foam stiffness increases when density increases. Of course, the density and porosity of foams are interchangeably affected by external mechanical forces that act to compress the foam and narrow or collapse the spaces in the microarchitecture (which further reduces the absorbency capacity of the dressing subjected to the loads³¹). The foam densities reported by Lee et al³² for polyurethane foams used in commercially available dressings were between 0.06 and 0.26 g/cm³, hence there is relatively large variability in this property among existing products. The detailed derivation of how the strength and stiffness of foam materials depend on their porosity and density is provided in Appendix A.1 for readers interested in the technical aspects of foam material science. The theoretical influence of the porosity and compressive strain level on the stiffness of a foam dressing is further illustrated in Figure 2B. Generally, for foams made of the same solid polymer phase, there is a non-linear increase in the stiffness with the decrease in the porosity and increase in the level of the applied external loading (or the non-dimensional deformation, ie, strain level), as depicted in Figure 2B.

2.1 | Foams in wound dressings under large mechanical deformations and repetitive loading

Despite best effort and intention to ensure proper offloading, many wounds are subjected to mechanical loads, unintentionally or deliberately, and such loads may be sustained (static) or dynamic/cyclic in nature.^{30,33} For example, acute care patients requiring cardiopulmonary support through extracorporeal membrane oxygenation (ECMO) but develop a sacral PU/PI during the supine ECMO support period, which is then treated by applying a dressing to their non-offloaded sacral wound are difficult to reposition as the postural changes may dislodge the device. Likewise, patients with spasticity, those with a severe gastroesophageal reflux disease, patients with dementia or delirium, and those with psychiatric disorders are challenging to reposition and often apply their bodyweight on their wounds, and on their dressings. Another example is the gold standard treatment of a venous leg ulcer (VLU) because of venous insufficiency, which typically includes the application of a dressing on

the VLU, and graduated compression therapy over the applied dressing, so that the dressing is deformed in compression and must function under the sustained compression.³⁴⁻³⁶ Patients with a DFU will preferably have their wounds partially or completely offloaded by means of an offloading (plantar pressure redistribution) device, for example, a total contact cast, but ambulatory patients are sometimes non-adherent with either offloading or immobilisation, and apply their bodyweight on the DFU.^{29,37-40} Moreover, even when offloading devices are used by patients with DFUs as indicated, and in combination with foam dressings (which is a common clinical practice^{41,42}), the dressing may move under the device, for example, because of minor misalignments of the device causing shear forces on the dressing that lead to migration of the dressing during walking. Prior studies suggest that individuals with a DFU walk over 4000 steps per day on average, in particular those who use removable offloading devices for whom the estimated average of daily steps is approximately 4500 steps.⁴³⁻⁴⁵ Moreover, the duration of standing, as another important measure of weight-bearing activities, is estimated to be three-folds longer than the duration of walking for these persons.⁴⁶ Thus, extra attention should be paid to the resilience of wound dressings applied to DFUs against repetitive pressure and shear loading. This phenomenon, observed by the clinical authors of this work, not only challenges the absorbency of the applied dressing but also its capacity to stay-on, and while a dressing migrates, it shears the peri-wound and may deteriorate it. For repetitive or impact loads such as those occurring in a wound dressing attached to the plantar surface of the foot to treat a DFU, Equation (A8) in Appendix A.1 may not be a sufficient description and specific experimental characterisation will be required.

All the above clinical scenarios necessitate the understanding of how foams within foam-based wound dressings behave under compressive mechanical loading. The stress-strain curve of a soft polyurethane foam undergoing repetitive or impact compressive loading can be divided into four typical regions: elastic, plateau, densification and unloading regions (Figure 3).^{27,49-51} These regions correspond to the variation in the apparent mechanical properties, because of the changes in the microstructure of the foam. Specifically, in the elastic (also called the linear) region, the stress linearly increases with the strain and reaches an initial peak stress. This region is generally observed at a strain of 5% to 10%, and when the compressive load is removed, the foam specimen returns to its original shape. Of note, foams in foam dressings, particularly when subjected to bodyweight loads or under compression therapy, may be deformed to substantially larger strains. While the elastic region

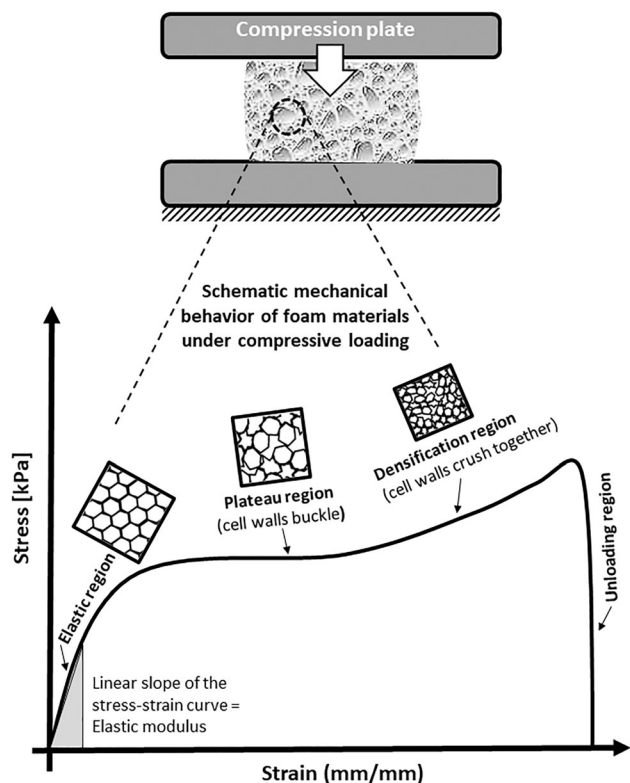


FIGURE 3 Schematic compressive stress-strain behaviour of flexible foams, including soft polyurethane foams, when subjected to large deformations. According to Gibson and Ashby,^{47,48} three compression regions can be identified in the curve: elastic (also known as the linear region), plateau (also termed the buckling region), and densification. Each of these three regions corresponds to a different compression mechanism, as follows. The elastic (linear) region is associated with bending of the cell walls, and corresponds to a first deformation of the foam material struts under the increasing load. The plateau (buckling) region corresponds to the collapse of the foam cells by elastic buckling. In this phase, the struts bend with little increase in the compressive force. Consequently, the apparent foam stiffness decreases with respect to the elastic/linear region as the material becomes effectively softer. The densification region corresponds to a state of completely collapsed cells, where opposite cell walls touch each other such that further strain compresses the solid itself. The stress then increases rapidly with the increased compression, and the foam becomes stiffer with respect to the former two regions. Detailed analytical descriptions and example experimental data for this stress-strain behaviour for flexible foams can be found in Alzoubi et al²⁷ and Elfarhani et al⁴⁹ and in the references cited therein

corresponds to a relatively intact cell structure that begins to deform with the bending and stretching of the cell walls, in the plateau (also called the buckling) region, the cells are already substantially deformed. The apparent foam stresses in the plateau region may fluctuate slightly with the strain, but overall, the stress maintains the average stress level despite an increase in the strain,

corresponding to the altered cell structure and collapse of the cells in an elastic buckling mode. The densification region forms when a substantial load is applied to the foam. This region corresponds to a structure in which most of the cells are distorted to such an extent that the cell walls are already in contact with each other (Figure 3). In the unloading region, the stress rapidly decreases to zero.⁵⁰ The strain that decreases because of the rebounding of the specimen during unloading is the elastic strain (Figure 3).

The above theoretical and experimental formulations clearly demonstrate that the stiffness properties of foam dressings may vary considerably for different clinical usage scenarios, primarily depending on the magnitude of the mechanical loads to which the dressing is subjected, from zero for a completely offloaded wound to very large strains as occurring under compression bandaging. Of note is that the compressive strain ε of a foam dressing may not necessarily remain constant over time. While in the context of a non-offloaded PU/PI, or a VLU treated by dressings under compression bandaging, the strain level in the dressing materials may be approximately steady for long time periods, dressings applied to non-offloaded plantar DFUs may be subjected to repetitive and rapid deformations. For dressings applied to such plantar DFUs, the compressive strain can, therefore, be theoretically simplified as cyclic (sinusoidal), that is, $\varepsilon(t) = \varepsilon_0 \sin(\omega t)$ where ε_0 is the strain amplitude, t indicates time and ω indicates the step frequency for the relevant activity; the frequency of human walking, which is relevant in this regard, is within the range of 1.7 to 2.1 Hz.^{52,53}

2.2 | Additional micro-morphological foam characteristics related to foam function in wound dressings

Scanning electron microscopy (SEM) is a useful imaging modality for determining the morphological differences between foam materials used in wound dressings, as reported by Lee et al.³² They studied the uniformity and pore size of foams used in commercial wound dressings and reported that some foams have a substantially more uniform microstructure than others, for example, the pore diameters in the surface of the foam of one dressing were at the range of 25 to 75 μm while pore sizes in the surfaces of foams of other dressings were spread over a much wider range of 32 to 1000 μm . Similar non-uniformity was also observed at the depth of the foams. Other than the diameter of the pores, their shape can be characterised as well, such as by means of the aspect ratio, a function of the largest diameter and the smallest

diameter orthogonal to it. The shape factor (SF) is another important dimensionless morphological parameter, which depends on the cross-sectional pore area (A) and the pore perimeter (P), and determines the degree of circularity of the pores⁵⁴:

$$\text{SF} = \frac{4\pi A}{P^2}. \quad (1)$$

Of note, the SF of a perfectly circular pore is 1, and elongated pores have lower SFs. The work of Yunus et al⁵⁴ demonstrated that the SF of open-cell polyurethane foams (often used in wound dressings) is typically substantially below unity, and may approach 0.5,⁵⁵ which is indeed visualised by the elliptical shapes of pores in the SEM micrographs obtained by Lee et al.³²

Another important morphological parameter is the thickness of the walls that make the pores, also known as the struts, which was investigated by Heit et al,⁵⁶ also using SEM. They found that in foams used in wound healing products, the wall thickness increases with the size of the pores, and is approximately 70 μm for small pore sizes (720 μm), 130 μm for medium pore sizes (1300 μm) and 300 μm for large pore sizes (3100 μm) in the foams examined in their study. Both the pore size and the wall thickness affect the stiffness and strength of the foam, as both influence the porosity value and, thereby, the level of material resistance to mechanical forces through deformation (Equation A8 in Appendix A.1). Indeed, the work of Lee et al³² demonstrated that different foam materials used in wound dressings have strength properties that span over orders of magnitudes, from 0.011 to 0.248 kg/mm^2 (108 kPa to 2432 MPa), and likewise, the elongation at failure of the tested foams ranged between 180% and 1101%.

Lastly, the interconnections between cells in the microstructure of foams, also known as the interconnectivity, is a critical morphological feature in foams used in wound dressings, as the interconnections ultimately facilitate fluid transfer between pores in the foam. The interconnectivity of foams can be quantified simply as the percentage of the cells that are connected to each other in their vicinity or to the surface, out of the total cell count. Closed-cell foams have zero connectivity, whereas open-cell foams have up to 100% connectivity. Rodriguez-Perez et al⁵⁷ indicated that most open-cell polyurethane foams have connectivity of approximately 99%. Tortuosity (φ) is a morphological parameter related to connectivity, measuring the distance that a fluid needs to travel in order to transport from one pore to an adjacent pore. For a three-dimensional pore microarchitecture in the microstructure of foams, the tortuosity can be defined as the Euclidean distance sums of the centroids of the

interconnected pores divided by the length of the pores. Because tortuosity is challenging to measure in practice, correlations are often used to obtain a first approximation from the porosity \emptyset , such as by means of the Bruggeman equation:

$$\varphi^2 = C_1 \emptyset^{1-C_2}, \quad (2)$$

where C_1 and C_2 are empirical constants that are normally taken as 1 and 1.5, respectively.⁵⁸ Note that when the porosity approaches unity (ie, when theoretically, there are no solid barriers for fluid flow), then the tortuosity φ approaches 1, which is the minimal possible value (ie, the dimensionless shortest route, of a straight line, for a fluid to transport between two points). Rodriguez-Perez et al⁵⁷ examined a polyurethane open-cell foam for tortuosity by measuring the electrical resistance of the foam specimen while an electrical conducting liquid was transported through the foam. They found that for a foam with a cell size of 356 μm (which is in the midrange of dressing foams) the tortuosity was 1.7. In addition, the tortuosity tends to decrease when the cell size increases. Tortuosity is not merely a morphological property; the tortuosity of foams used in wound dressings plays an important role in the context of the potential swelling of foams, and the associated mechanical pressures that they may apply on their surroundings such as the wound bed and peri-wound skin, as further discussed below.

2.3 | Mechanical durability of foams in wound dressings

The mechanically vulnerable element in foams is the struts, and hence, fatigue-related mechanical microdamage in foams is associated with bending, elastic buckling and ultimately, plastic (irreversible) collapse of the struts.⁵⁹ Accordingly, the durability of foams also originates at the microstructure and is linked with the sizes and shapes of the pores and the thickness of the walls that separate them, as the micro-failure (microdamage) accumulation is influenced by the extent of wall bending and stretching.⁴⁷ In terms of the apparent (bulk) properties, the mechanical stresses leading to microdamage in foams, therefore, strongly depend on the foam density or porosity, as well as on the strength of the solid phase of the polymer material (Equations A2 and A5 in Appendix A.1).⁶⁰ In addition, any existing defects such as edge cracks or internal holes in the foam can compromise its mechanical durability, with edge defects being more likely to compromise the strength of foams with respect to internal defects.⁶¹ The latter point has practical

implications on the function of foams in foam-based wound dressings. A dressing with a visible physical defect caused by mechanical forces, such as because of intense rubbing, sustained indentation or scratching, is more prone to further damage from (subsequent) mechanical forces, which may lead to disintegration of the dressing and, thereby, a compromised healing of the wound.^{14,15,62}

3 | THE BENDING STIFFNESS OF FOAM DRESSINGS AND THE ASSOCIATED WOUND BED PROTECTION

The bending stiffness of foams used in wound dressings is highly important in the context of the mechanical protection provided by the dressing to the wound bed and the peri-wound tissues, as well as in terms of the conformability of the applied dressing to curved body surfaces, and its ease of use (ie, how much a dressing would mechanically ‘resist’ to applying it on convex body surfaces). Park et al⁶³ measured the bending stiffness of polyurethane foams in the context of flexion of shoe insoles, and as expected, their results demonstrated that the bending stiffness of foams depends on the thickness and elastic modulus† of the specific foam material. This aligns with the fundamental theory of flexural bending, which determines that the bending stiffness (also known as the flexural rigidity) of an elastic material is the product of the elastic modulus (Figure 3) and the moment of inertia of the cross-section. For a rectangular cross-section (which is characteristic to the vast majority of wound dressings), the moment of inertia of the cross section is $w t^3/12$ where w is the width and t is the thickness of the dressing, hence the thickness of the dressing plays a major role in the bending stiffness (because of the third power in the latter formula). For example, to illustrate the effect of the thickness on the bending stiffness, and hence on the conformability of foams in foam-based dressings, in order to double the bending stiffness of a single-layer foam dressing having a unit thickness, it is possible to increase its elastic modulus two-fold through lowering the porosity (Equation A3 in Appendix A.1) without changing its thickness, or alternatively, to use the same material but increase its thickness by just 25%. A thicker dressing not only increases the bending stiffness but also increases the absorbency reservoir of the foam for wound fluid uptake. On the other hand, a greater bending stiffness may cause discomfort to patients if the dressing is placed on a body part that is relatively mobile such as over a joint (eg, in flexion/extension).

A higher bending stiffness of the dressing can also make it more difficult to apply the dressing on curved body surfaces (because the dressing will resist in shape change to achieve conformability, and it will be more challenging for the adhesive mechanism of the dressing to maintain the curved shape due to an elastic recoil). As the body is not a flat plane and most of it has convex surfaces that often move substantially (such as the feet, elbows, knees, hips, or neck), some manufacturers have developed specialised flexible dressings with a deliberately low bending stiffness achieved, for example, using patterned perforations in the material, to match anatomy-specific or patient-specific needs of conformability and/or body movements with the dressing.^{64,65} The added flexibility of the dressing by means of such treatments of the foam materials contributes to decreasing dressing wastage through this enhanced capacity to remain in place for as long as needed.⁶⁴

4 | ADHESION OF THE DRESSING BORDERS TO THE PERI-WOUND AND IMPLICATIONS OF NON-ADHERENCE

Optimising the adhesiveness of dressings has been a persistent challenge to the wound care industry. In modern dressings, an adhesive mechanism is integrated in at least a border area around the wound pad so that no further fixation (with secondary adhesive tape or polyurethane film layer) is needed (though for some highly convex or moving anatomical regions, such as the female breast and the knees or axilla, respectively, an adhesive border of a standard-shape dressing is often not suitable or needs to be cut to shape⁶⁶). Repeatedly changing these adhesive wound dressings frequently causes pain and stripping damage, which is removal of a large amount of stratum corneum from the newly formed epithelium and/or the peri-wound skin with the adhesive.^{67,68} With the introduction of silicone adhesives, the extent of this problem decreased‡,^{69,71} yet peri-wound skin trauma caused by the repetitive removals of adhesive dressings still occurs and is known to increase the size of wounds and the risk of infections and delays healing, and thereby, adversely affect the quality of life of patients and have cost implications for health care providers.⁷⁰ In the more recent literature, this is referred to as medical adhesive-related skin injury (MARS), which is associated with a history of contact dermatitis or prolonged bed rest periods and is considered preventable, including through the selection of appropriate dressing products, in addition to suitable procedures for application, monitoring, reapplication and removal of the dressings.⁷²⁻⁷⁵

Despite that foam materials are generally considered non-adhesive, there are many foam-based dressings with an adhesive skin contact layer that enable the dressings to be secured in place without the need for secondary support or fixation. An overly aggressive adhesiveness to the wound or to the fragile peri-wound skin may result in trauma and pain upon removal, and may also cause stripping of the stratum corneum, leading to an inflammatory response and compromised barrier function.⁷⁶⁻⁷⁸ However, it is important to balance this consideration with the need to adequately secure the dressing in place for the intended period of use.⁶⁶ An insufficiently strong (or durable) adhesiveness leads to frequent dressing detachments, therefore requiring more dressing changes, which may also cause skin irritation or MARSIs, leading to discomfort and pain, in addition to increased costs of wound care. Importantly, the frequency of dressing changes, which is strongly affected by the quality of the adhesiveness of the applied dressing over time, directly impacts the quality of life of patients and the overall cost of care.^{8, 4,70,82} Dykes et al⁷⁷ indicated that foam dressings with an acrylic adhesive require the greatest separation forces to detach from the forearm skin (2.2 N), followed by polyurethane adhesives (1.7 N), and the least force to peel the dressing occurred when the dressing had a soft silicone adhesive (1.1 N) or a hydrocolloid adhesive (1 N). Their findings regarding a soft silicone adhesive agree with visual analog scale-based and McGill questionnaire-based pain evaluations conducted by Woo et al⁸³ who compared a soft silicone foam dressing (Mepilex[®] Border; Mölnlycke Health Care AB, Gothenburg, Sweden) with an adhesive hydrocellular polyurethane foam dressing (Allevyn Adhesive; Smith & Nephew, Hull) and reported that the foam dressings with the soft silicone adhesive were associated with less pain, both before and during dressing changes. Surprisingly, the Dykes⁷⁷ work highlighted that the level of force needed to separate the dressing from the skin does not always correlate to the damage to the stratum corneum, which was greatest for the acrylic adhesive (95%) and negligible for the soft silicone adhesive. However, the hydrocolloid adhesive associated with relatively low peeling forces caused stratum corneum damage which was equivalent to that of a polyurethane adhesive (85%-90%). An important observation described by Waring et al⁷⁸ is an overall wide inter-subject variation in the levels of adhesion to human skin (ie, variability in inter-subject peeling forces), and likewise, product-related differences in attachment to dry versus moist or oily skin.

Differences in peel force levels are reported in the literature, for example, by Dykes et al⁷⁷ (approximately 1 to 2 N) and Waring et al⁷⁸ (approximately 0.5 N), which may be associated with anatomical site differences and

test method settings, such as the dressing removal technique, angle and the speed of removal, which plays a crucial role. As the skin is viscoelastic, the state of contact stresses between the dressing and the skin strongly depends on the speed of removal: the faster the removal, the greater the skin (and wound) stresses are.⁵ The latter point has important clinical implications, in the sense that the dressing removal technique, as taught, trained and implemented in practice, is of critical importance in order to minimise the risk of MARSIs. Indeed, current clinical MARSIs prevention guidelines recommend a 'low and slow' removal technique, that is, to slowly pull back the adherent dressing at a low horizontal angle (approximately parallel to the skin), away from the corner or edge so that it is gradually separated from the skin, which not only provides greater control of the removal action, but also reduces the localised wound and peri-wound skin stresses as indicated above.^{73,84}

Standard peel test methods utilising standardised substrates, in particular steel and glass, are commonly used by the wound dressing industry, as reviewed by Bernatchez and Bichel.⁸⁵ A few experimental synthetic viscoelastic substrata, which mimic adhesion on human skin, were developed in an academic research context to measure adherence properties,^{86,87} or for surgical simulations.⁸⁸ However, from a standardisation point of view, repeatable and robust methods are required, hence complex substrates run the risk of increasing variability substantially. In addition, information is needed regarding how the adhesiveness performance may be affected by the wound fluid over time for different exudate types, compositions and viscosities.

A great promise for research in this area of wound dressing science is in dynamic computer models to simulate the behaviour of both the viscoelastic human skin during dressing detachment and the adhesive technology under investigation. Computer modelling and simulation work conducted at the group of the lead author (AG) indicate that the adhesion forces, angle of removal and speed of removal of the dressing all affect the state of mechanical stresses of the peri-wound skin, and thereby the risk for developing a MARSIs. The pattern by which peri-wound stress concentrations change in location and magnitude throughout the removal process for a dressing peeled from a planar wound is demonstrated in such a computer simulation in Figure 4, and further bioengineering analysis is provided in Appendix A.2. Specifically, the formulation in Appendix A.2 indicates that the most sensitive region of the peri-wound skin to stripping damage is where the wound width is maximal (assuming that there is no adhesion between the wound pad and the wound bed). However, the equations in Appendix A.2 also demonstrate that the level of the applied force and

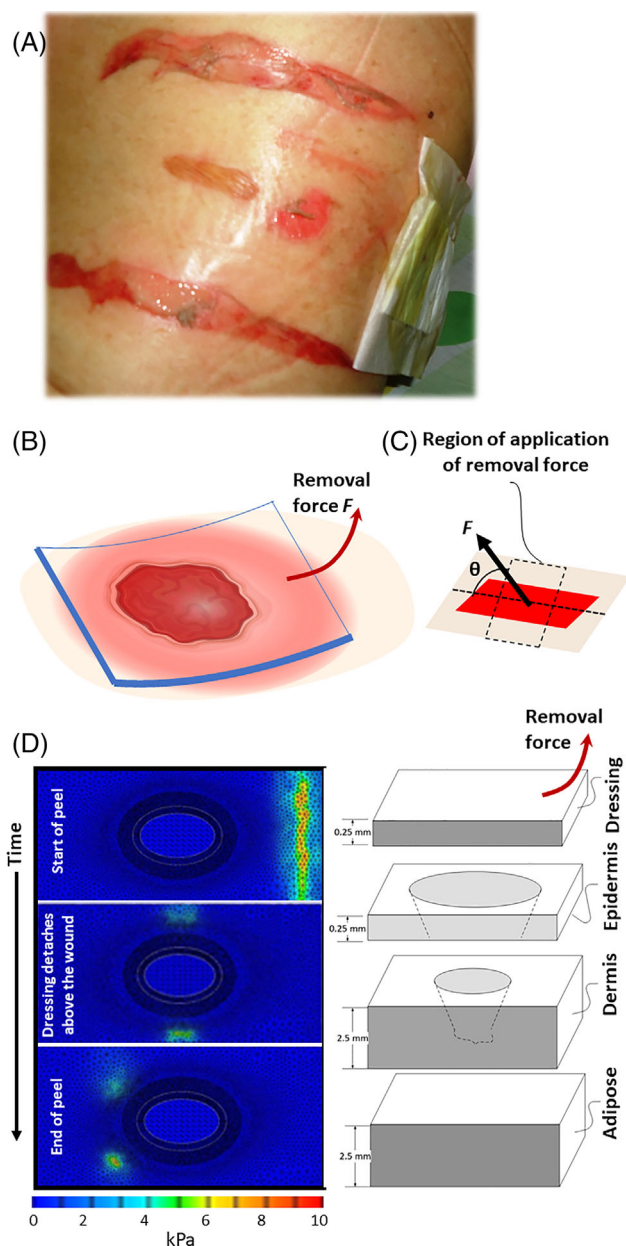


FIGURE 4 A medical adhesive-related skin injury (MARS), photograph courtesy of Mr Paulo Ramos (Northern Regional Health Administration, ARS Norte Vila do Conde, Portugal) (A), and example bioengineering in silico research of the aetiology of MARS conducted at the laboratory of author AG, using computational (finite element) simulations of the process of removal of a dressing (B) by applying a removal force F at an angle θ (C). Both the applied force and the angle of the force application may change over time, resulting in the detachment of the dressing from the peri-wound skin in the computer model (D; right frames). The location of the effective skin stress concentrations associated with the detachment action moves over the peri-wound skin while the dressing detaches (D; left frames)

the angle of application are important factors affecting the skin stress concentration magnitudes, and can be controlled by a trained clinician. As mentioned above,

under real-world conditions, this interaction becomes more complicated as it also depends on the speed of the removal process, which cannot be kept constant as this is a manual manoeuvre. Another factor affecting the intensity of the tissue stress concentrations during the peeling is the convexity of the wound and peri-wound surfaces, which is not taken into account in the analyses detailed in Figure 4B-D and Appendix A.2; stress concentrations are typically amplified on curved surfaces.⁸⁹ Importantly, the current theory indicates that faster dressing removals create larger skin reaction forces because of the viscoelasticity of the skin. Accordingly, the dressing removal rate is a critical factor to consider for avoiding the occurrence of MARS, as recent relevant guidelines and clinical practice indeed highlight.^{73,75}

Lastly, it is important to note that the laboratory tests of peeling forces reviewed above, or the current computer modelling of the peel process of dressings (Figure 4D), do not account for the patient-specific and wound-specific, time-dependent biological interactions of the wound and dressing.^{2,28} For example, an important mechanism that increases the adhesion forces in real-world clinical practice (though it is not very common for foam dressings), is potential ingress of new granulation tissue into the wound-facing aspect of the dressing (^{5,6,90,91}). Another biological influence that may appear in real-world scenarios is if the exudate within the dressing dries out and fibrin-containing scab forms, which binds the dressing to the wound and result in greater peel forces when the dressing needs to be removed.⁵

5 | THE COF OF THE OUTER AND INNER DRESSING SURFACES

A wound dressing can be subjected to frictional forces causing shearing in numerous clinical scenarios, such as when the region of the wound is rubbing against the bedsheets, clothing or any other object (eg, the safety rails of a hospital bed), or during head-of-bed elevation, repositioning, turning, bathing, or transferring patients, or while a patient spontaneously moves in their bed or chair. During such events, the wound and peri-wound tissues are exposed to single-instance or repetitive shear deformations. Repeated frictional distortions may possibly cause breaks in the wound surface that allow pathogens to penetrate to the vasculature, and eventually generate secondary infections.⁹² These friction-related tissue distortions can be minimised if the COF of the dressing with the contacting elements is optimal.⁹³ Specifically, a high COF at the outer dressing surface with, for example, the bedsheets or clothing, will cause delivery of greater frictional forces to the dressing (the

frictional force is the normal force times the COF), resulting in more shear within the dressing structure, which may further distort the wound bed or peri-wound in shear if the dressing is not effective in absorbing the shear internally.^{33,94} Likewise, excess friction at the dressing-skin interface may compromise the peri-wound tissues, by inducing continuous or repetitive shear loads on them. Hence, typically, low COFs are desired for dressing materials in contact with either an external object (ie, at the outer dressing surface) or at the wound-facing aspect of the dressing. In addition, adequate attachment of the dressing to the skin is required to minimise frictional sliding movements of the dressing over the skin/wound.⁹⁵

The COF of polyurethane foams for potential use as a skin-contacting material in wound dressings varies from approximately 0.15 for the harder foams up to 1.5 for the softest grades.^{93,96-98} Specifically, Vilhena and Ramalho⁹³ measured the COF of different materials used in a hospital setting, including a polyurethane foam, against the ventral forearm for natural dry adult skin, and obtained a range of COFs between 0.25 and 0.47 with a midrange of 0.36, which is similar to the COFs of polyester-fabric or cotton-made bed protectors (being around 0.4⁹⁹), but greater than for specialised hospital fabrics (0.27) or diapers (0.28). The above foam-skin COF values are at the lower end of those that can be expected in a moist wound environment, which is constantly exposed to exudate fluids, and/or where there is substantial sweating from the peri-wound region. Typically, COFs of synthetic materials with skin largely increase in the presence of moisture⁹⁷ because of the hydrosensitive mechanical behaviour of the stratum corneum that softens in the presence of water (which is known as the plasticizing effect). Exposure to moisture and wetness may further smoothen the skin roughness asperities profile, and consequently, increase the de-facto contact area of materials with skin, thereby inducing more frictional adhesion, that is, a correlated adhesion and friction behaviour.^{93,100} Indeed, Schwartz et al⁹⁷ measured the COFs of porcine skin, dry and wet, with saline, sweat or urine, when rubbing against a single-layer polyurethane foam dressing using a tilting-table tribometer, and reported that the wet contact had a 1.3-fold greater COF with biological fluids (sweat and urine), overall increasing the COF more than saline (pointing to the importance of testing dressing materials with realistic biological fluids, not just saline). Of note, some body regions such as the axilla are more densely populated with sweat glands,¹⁰¹ hence the above effect of perspiration increasing the COF of polyurethane foams with skin is affected by the anatomical site. To reduce the COFs of wound-contacting foams, surface polishing or treatment with lubricants can be applied⁹³;

however, such surface modifications will likely be more needed if the dressing is indicated for highly perspiring peri-wound or largely exuding wound conditions where greater dressing-skin COFs are expected because of the continuous fluid exposure.⁹⁷

The COF of the external surface of the dressing with potentially contacting materials such as the clothing and bed linen is also highly important, as high COFs will lead to high external frictional forces, shearing of the dressing and potential transmissibility of the shear stresses forming in the dressing to the wound bed and peri-wound region. Related to this, Ohura et al¹⁰² measured the COF between the outer layers of wet dressings and clothing, and found, as may be expected, that these COFs were sensitive to the level of wetness in the dressing. They also found that a foam-based dressing exhibited a lower COF with the clothing with respect to a hydrocolloid for simulated highly exuding conditions.¹⁰²

6 | SHAPE CONFORMABILITY OF DRESSINGS TO THE WOUND BED AND TO BODY SURFACE CONTOURS

The curved and asymmetrical surfaces of the human body are challenging for application of foam dressings, specifically bordered ones.^{66,103} In particular, the clinical demand for placing dressings on irregular body surfaces at various anatomical regions to treat different wound aetiologies is often contradictory to manufacturers' interests in standardisation of the shapes (including symmetry) and sizes of dressings to optimise the processes and costs of production. Conformability of a foam material to the wound shape is warranted but needs to not overstress the wound bed because of swelling and dilatation as the dressing absorbs exudate fluids. The shape conformability of dressing materials and structures results from the interaction of the foam material characteristics such as the elastic modulus, the bending/flexural modulus and the Poisson's ratio, and the swelling behaviour of the foam. Theoretically, a dressing material with a high-volume capacity storage (which may intuitively be perceived as beneficial for fluid handling) could have the potential negative effect of dilatation in a confined wound bed space to an extent causing localised pressure increases on the delicate wound bed tissues. Excessively high pressures may lead to tissue stress concentrations in the wound bed and possible wound deterioration, by mechanically damaging an ongoing epithelialization and/or by inducing pain and psychological stress to the patient. With that said, low-level mechano-modulation has been shown to be linked to the stimulation of inflammatory, fibroblast migratory and fibrosis-related tissue

remodelling responses, and hence theoretically, gentle, moderate pressures on the wound bed may, in fact, contribute positively to the wound healing process.^{28,104,105}

The theory and derivation of the swelling pressure of foams in foam-based dressings are detailed in Appendix A.3. The formulation in this Appendix demonstrates that the swelling pressure of a foam-based dressing in the wound bed depends on the elapsed time post-application of the dressing and the effective diffusion coefficient of the foam, which in turn depends on the porosity and tortuosity of the foam. Specifically, the derivation in Appendix A.3 indicates that the swelling pressure will increase faster for a greater porosity \emptyset and lower tortuosity φ in the microarchitecture of the foam-based dressing. In other words, to control the compression and the rate of compression applied by a foam material onto the wound bed when the foam-based dressing continuously absorbs exudate fluids, the swelling pressure can be regulated through appropriate selection of the microstructural features of the foam, specifically, by limiting the porosity \emptyset , increasing the tortuosity φ or doing both (Equation A13 in Appendix A.3). Similarly, to the other dressing characteristics reviewed here, the extent of swelling differs notably across manufacturers and products, and this can be associated, at least partially, to the above theory, as the foam microstructural features also differ considerably between commercially available foam-based dressings.¹⁶

7 | SUMMARY AND CONCLUSIONS

All the key aspects of the mechanical performance of foams within foam-based wound dressings strongly depend on the microstructure of the specific foam. For example, the compressive and tensile stiffness and strength of foams, affecting the ability of the dressing to protect the wound mechanically, and the bending stiffness of foams, which influences the conformability performance of foam-based dressings, all depend on the relative density and porosity of the foam materials. Likewise, the COFs of foam dressings with the wound bed and peri-wound skin depend on the microtopography of the surfaces. Although there are not many published works focusing on the microarchitecture of foams in foam-based wound dressings, the variability in microstructural properties reflected in the published studies reviewed here, with even the most fundamental property of the pore size ranging over an order of magnitude across manufacturers and products (ie, 100–1000 μm), reflects that an optimal porosity range of values for foam dressings is unknown at this time. As evident from the

theoretical formulations in this article, the variability in pore sizes then reflects on high variabilities in foam stiffness, strength, conformability performance and durability of the foam—all of which are critical for delivering adequate mechanical protection to the wound bed and peri-wound, consistently over the intended period of use. In addition, multiple balances and optimizations are needed in the chosen mechanical properties of foam materials used in foam dressings, and the structure of foam dressings as a construct, for achieving an adequate and clinically efficacious dressing design; relevant examples are summarised in Table 1.

Currently, no test standards exist that characterise the microstructure of foam-based dressings as part of their performance evaluation, and of course, no target values exist to guide manufacturers regarding the foam porosity, shape of pores and the variations in pore size and shape that would provide superior performance in the mechanical aspect of wound protection. Likewise, COFs and peeling forces are not standardised in the wound dressing industry, despite the critical impact of both on potential wound bed and peri-wound mechanical damage related to movements of the dressing against the skin or against elements in the outer environment, dressing removals and repetitive changes. Testing methods such as the ASTM 903,¹⁰⁶ which is a standard test method for peel or stripping strength of adhesive bonds, are used in the absence of specific peeling tests for wound dressings. A particular challenge in this regard is to standardise a material substrate to represent the skin in such testing, although a range of options potentially exists, such as elastomers (silicones), ethylene/methyl acrylate films, epoxy resins, textiles and metals, or alternatively, excised porcine skin.^{86,107,108} Preliminary research towards the standardisation of testing of adhesive bandage tapes, which are, of course, much less complex than advanced foam-based wound dressings, already indicated that the peeling forces depend on the width of the tape and storage conditions, two example properties that are typically not considered in the evaluation of adhesiveness of wound dressings.¹⁰⁹ Importantly, theoretical and computational analyses such as those reported here can be used to identify influential factors, for example, as in Equations (A9 and A10) (Appendix A.2), which point to the role of the dressing width in the skin loading state during removals, but details of theoretical-computational frameworks for wound dressings are also lacking in the literature. This article, therefore, provides, for the first time, a scientific compilation of theoretical considerations and experimental results related to the mechanical performance of foam materials in foam-based wound dressings, in the light of their frequent usage and

TABLE 1 Examples of balances and optimizations needed in the mechanical properties of adequately designed foam dressings

Mechanical property	If excessively low, will result:	If excessively high, will result:
Compressive material stiffness (elastic modulus) ^a	<ul style="list-style-type: none"> Compromised physical-mechanical protection to the wound bed against external forces and impacts Extreme deformations under external loads leading to potentially low durability and dressing disintegration, which compromises the wound healing 	<ul style="list-style-type: none"> Risk of imprinting the peri-wound skin, increasing the inflammatory response and damaging the peri-wound viability, which then compromises wound healing Low conformability to curved body surfaces and increased likelihood of dressing detachments
Structural (bending) stiffness	<ul style="list-style-type: none"> Compromised physical-mechanical protection to the wound bed against external forces and impacts Extreme deformations under external loads leading to potentially low durability and dressing disintegration, which compromises the wound healing 	<ul style="list-style-type: none"> Risk of imprinting the peri-wound skin, increasing the inflammatory response and damaging the peri-wound viability, which then compromises wound healing Low conformability to curved body surfaces and increased likelihood of dressing detachments
Adhesive strength	<ul style="list-style-type: none"> Frequent dressing detachments exposing the wound bed surface and leading to a greater likelihood of wound infections, thereby compromising the wound healing 	<ul style="list-style-type: none"> Risk of developing a medical adhesive-related skin injury that increases peri-wound inflammation and damages the peri-wound viability, thereby compromising the wound healing

^aPlease see Figure 3.

clinical relevance, en route to the development of standard, clinically relevant mechanical performance metrics for wound dressings.

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DATA AVAILABILITY STATEMENT

Data sharing not applicable to this article as no datasets were generated or analysed during the current study.

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ENDNOTES

* The apparent density refers to the ratio of the mass and the volume of the foam test specimen (the volume is that occupied by the solid polymer phase including all the hollow spaces therein).

† Slope of the linear part of the stress–strain curve (Figure 3).

‡ Though often, additional adhesive tapes are still needed to keep a bordered foam dressing securely in place, especially if the dressing is cut to fit curved or small body regions⁶⁶; adhesives are also less effective on some skin types, depending, for example, on sebum levels, skin dryness and sweating.^{69,70}

§ Dressing products and related supplies (eg, for wound cleansing) generally represent 15% to 20% of the total wound management costs, whereas more substantial costs are of the nursing time (30%–35% of total cost) and hospitalisation (>50%).^{79–81}

REFERENCES

- Corbett LQ, Ennis WJ. What do patients want? Patient preference in wound care. *Adv Wound Care*. 2014;3(8):537–543.
- Cullen B, Gefen A. The biological and physiological impact of the performance of wound dressings. *Int Wound J*. 2022; in press. doi:10.1111/iwj.13960
- Davies P, McCarty S, Hamberg K. Silver-containing foam dressings with Safetac: a review of the scientific and clinical data. *J Wound Care*. 2017;26(Sup6a):S1–S32.
- Fletcher J, Wasek S. Clinician perspectives on time and resources related to dressing changes. *Wounds Int*. 2016;7(4):28–32.
- Gefen A, Alves P, Beeckman D, et al. How should clinical wound care and management translate to effective engineering standard testing requirements from foam dressings?

- Mapping the existing gaps and needs. *Adv Wound Care*. 2022a; in press. doi:[10.1089/wound.2021.0173](https://doi.org/10.1089/wound.2021.0173)
6. Gefen A, Sibbald G, Woo K. So many options, so little time: dressing selection tips from the experts. *Wound Care Canada*. 2022b;20(1):18-21.
 7. Payne WG, Posnett J, Alvarez O, et al. A prospective, randomized clinical trial to assess the cost-effectiveness of a modern foam dressing versus a traditional saline gauze dressing in the treatment of stage II pressure ulcers. *Ostomy Wound Manage*. 2009;55(2):50-55.
 8. Scanlon E, Karlsmark T, Leaper DJ, et al. Cost-effective faster wound healing with a sustained silver-releasing foam dressing in delayed healing leg ulcers—a health-economic analysis. *Int Wound J*. 2005;2(2):150-160.
 9. Walker RM, Gillespie BM, Thalib L, Higgins NS, Whitty JA. Foam dressings for treating pressure ulcers. *Cochrane Database Syst Rev*. 2017;10(10):CD011332.
 10. Walker RM, Gillespie BM, Thalib L, Higgins NS, Whitty JA. Foam dressings for treating pressure injuries in patients of any age in any care setting: an abridged Cochrane systematic review. *Int J Nurs Stud*. 2018;87:140-147.
 11. Duncan G, Andrews S, McCulloch W. Issues in clinical practice: dressings. *Primary Intent*. 2002;10(1):29-35.
 12. Fontenot KR, Edwards JV, Haldane D, et al. Structure/function relations of chronic wound dressings and emerging concepts on the interface of nanocellulosic sensors. In: Filpponen I, Peresin MS, Nypelö T, eds. *Lignocellulosics*. Amsterdam: Elsevier; 2020:249-278 (Chapter 9). doi:[10.1016/B978-0-12-804077-5.00014-2](https://doi.org/10.1016/B978-0-12-804077-5.00014-2)
 13. Ghomi ER, Khalili S, Khorasani SN, Re N, Ramakrishna S. Wound dressings: current advances and future directions. *J Appl Polym Sci*. 2019;136(27):47738. doi:[10.1002/app.47738](https://doi.org/10.1002/app.47738)
 14. Lustig A, Gefen A. The performance of gelling fibre wound dressings under clinically relevant robotic laboratory tests. *Int Wound J*. 2022b;19(Suppl 1):3-21. doi:[10.1111/iwj.13761](https://doi.org/10.1111/iwj.13761) in press.
 15. Orlov A, Lustig A, Grigatti A, Gefen A. Fluid handling dynamics and durability of silver-containing gelling fiber dressings tested in a robotic wound system. *Adv Skin Wound Care*. 2022; 35(6):326-334. doi:[10.1097/01.ASW.0000823972.16446.ff](https://doi.org/10.1097/01.ASW.0000823972.16446.ff)
 16. Uzun M, Shah AT. In vitro characterisation and evaluation of different types of wound dressing materials. *J Biomed Eng Technol*. 2013;1(1):1-7.
 17. Thomas S. Foam dressings (chapter 10). *Surgical Dressings and Wound Management*. Cardiff: Medetec Publications; 2010a:181-194.
 18. Gefen A, Krämer M, Brehm M, Burckardt S. The biomechanical efficacy of a dressing with a soft cellulose fluff core in prophylactic use. *Int Wound J*. 2020;17(6):1968-1985. doi:[10.1111/iwj.13489](https://doi.org/10.1111/iwj.13489)
 19. Gefen A. Not all superabsorbent wound dressings are born equal: theory and experiments. *J Wound Care*. 2021;30(9):738-750.
 20. Tudoroiu EE, Dinu-Pîrvu CE, Albu Kaya MG, et al. An overview of cellulose derivatives-based dressings for wound-healing management. *Pharmaceuticals*. 2021;14(12):1215.
 21. Sanusi AL. Severe wound traction-blisters after inadequate dressing application following laparoscopic cholecystectomy: case report of a preventable complication. *Patient Saf Surg*. 2011;5(1):4.
 22. Gefen A. The selection of cushioning and padding materials for effective prophylaxis of medical device-related pressure ulcers: clinical intuition does not always work. *Wounds Int*. 2022a;13(1):10-19.
 23. Orlov A, Gefen A. Differences in prophylactic performance across wound dressing types used to protect from device-related pressure ulcers caused by a continuous positive airway pressure mask. *Int Wound J*. 2022a; in press. doi:[10.1111/iwj.13942](https://doi.org/10.1111/iwj.13942)
 24. Raepsaet C, Alves P, Cullen B, et al. Clinical research on the use of bordered foam dressings in the treatment of complex wounds: a systematic review of reported outcomes and applied measurement instruments. *J Tissue Viability*. 2022a;31(3):514-522.
 25. Raepsaet C, Alves P, Cullen B, et al. Study protocol for the development of a core outcome set (COS) for clinical effectiveness trials of bordered foam dressings in the treatment of complex wounds. *J Tissue Viability*. 2022b; in press. doi:[10.1016/j.jtv.2022.09.004](https://doi.org/10.1016/j.jtv.2022.09.004)
 26. McKee CT, Last JA, Russell P, Murphy CJ. Indentation versus tensile measurements of Young's modulus for soft biological tissues. *Tissue Eng Part B Rev*. 2011;17(3):155-164.
 27. Alzoubi MF, Al-Hallaj S, Abu-Ayyad M. Modeling of compression curves of flexible polyurethane foam with variable density, chemical formulations and strain rates. *J Solid Mech*. 2014;6(1):82-97.
 28. Gefen A. The influence zone: a critical performance measure for negative pressure wound therapy systems. *Br J Nurs*. 2022b;31(15):S8-S12.
 29. Lustig A, Gefen A. Fluid management and strength postsimulated use of primary and secondary dressings for treating diabetic foot ulcers: robotic phantom studies. *Int Wound J*. 2022a;19(2):305-315.
 30. Orlov A, Gefen A. The fluid handling performance of the curea P1 multipurpose dressing against superabsorbent and foam dressing technologies. *Int Wound J*. 2022b;19(4):945-956.
 31. Thomas S. Laboratory findings on the exudate-handling capabilities of cavity foam and foam-film dressings. *J Wound Care*. 2010b;19(5):192-199.
 32. Lee SM, Park IK, Kim YS, et al. Physical, morphological, and wound healing properties of a polyurethane foam-film dressing. *Biomater Res*. 2016;20:15.
 33. Schwartz D, Gefen A. The biomechanical protective effects of a treatment dressing on the soft tissues surrounding a non-offloaded sacral pressure ulcer. *Int Wound J*. 2019;16(3):684-695.
 34. Hall L, Adderley U. Active debridement of venous leg ulcers: a literature review to inform clinical practice. *Br J Community Nurs*. 2019;24(Sup6):S24-S29.
 35. Hampton S. Venous leg ulcers: short-stretch bandage compression therapy. *Br J Nurs*. 1997;6(17):990-998.
 36. Johnson S. Compression hosiery in the prevention and treatment of venous leg ulcers. *J Tissue Viability*. 2002;12(2):67-74.
 37. Ababneh A, Finlayson K, Edwards H, Lazzarini PA. Factors associated with adherence to using removable cast Walker treatment among patients with diabetes-related foot ulcers. *BMJ Open Diabetes Res Care*. 2022;10:e002640. doi:[10.1136/bmjdr-2021-002640](https://doi.org/10.1136/bmjdr-2021-002640)
 38. Crews RT, Armstrong DG, Boulton AJ. A method for assessing off-loading compliance. *J Am Podiatr Med Assoc*. 2009; 99(2):100-103.

39. Jeffcoate WJ, Harding KG. Diabetic foot ulcers. *Lancet*. 2003; 361(9368):1545-1551.
40. Kimawi AA, Snyder RJ, Cala MA, Cuffy C. When traditional offloading is not an option, could an external fixator be a solution?: A case report. *Wounds*. 2017;29(2):46-50.
41. Baker N, Osman IS. The principles and practicalities of off-loading diabetic foot ulcers. *Diabet Foot J*. 2016;19(4):172-181.
42. Vaseenon T, Thitiboonsuwan S, Cheewawattanachai C, Pimchoo P, Phanphaisarn A. Off-loading total contact cast in combination with hydrogel and foam dressing for management of diabetic plantar ulcer of the foot. *J Med Assoc Thai*. 2014;97(12):1319-1324.
43. Fernando ME, Woelfel SL, Perry D, et al. Dosing activity and return to preulcer function in diabetes-related foot ulcer remission. *J Am Podiatr Med Assoc*. 2021;111(5). doi:10.7547/20-166
44. Jarl G, van Netten JJ, Lazzarini PA, Crews RT, Najafi B, Mueller MJ. Should weight-bearing activity be reduced during healing of plantar diabetic foot ulcers, even when using appropriate offloading devices? *Diabetes Res Clin Pract*. 2021; 175:108733. doi:10.1016/j.diabres.2021.108733
45. Najafi B, Grewal GS, Bharara M, Menzies R, Talal TK, Armstrong DG. Can't stand the pressure: the association between unprotected standing, walking, and wound healing in people with diabetes. *J Diabetes Sci Technol*. 2017;11(4):657-667.
46. Najafi B, Crews RT, Wrobel JS. Importance of time spent standing for those at risk of diabetic foot ulceration. *Diabetes Care*. 2010;33(11):2448-2450.
47. Gibson LJ, Ashby MF. The mechanics of three-dimensional cellular materials. *Proc R Soc London Ser A Math Phys Sci*. 1982;382:43-59.
48. Gibson LJ, Ashby MF. *Cellular Solids: Structure and Properties*. 2nd ed. Cambridge: Cambridge University Press; 1997.
49. Elfarhani M, Jarraya A, Abid S, Haddar M. Fractional derivative and hereditary combined model of flexible polyurethane foam viscoelastic response under quasi-static compressive tests. *Cell Polymers*. 2016;35(5):235-270.
50. Hwang BK, Kim SK, Kim JH, Kim JD, Lee JM. Dynamic compressive behavior of rigid polyurethane foam with various densities under different temperatures. *Int J Mech Sci*. 2020; 180(15):105657.
51. Park KB, Kim HT, Her NY, Lee JM. Variation of mechanical characteristics of polyurethane foam: effect of test method. *Materials*. 2019;12(17):2672.
52. Pachi A, Ji T. Frequency and velocity of people walking. *Struct Eng*. 2005;3:36-40.
53. Uematsu A, Inoue K, Hobara H, et al. Preferred step frequency minimizes veering during natural human walking. *Neurosci Lett*. 2011;505(3):291-293.
54. Yunus S, Sefa-Ntiri B, Anderson B, Kumi F, Mensah-Amoah P, Sonko SS. Quantitative pore characterization of polyurethane foam with cost-effective imaging tools and image analysis: a proof-of-principle study. *Polymers*. 2019;11(11):1879.
55. Takashimizu Y, Iiyoshi M. New parameter of roundness R: circularity corrected by aspect ratio. *Prog Earth Planet Sci*. 2016;3:2.
56. Heit YI, Dastouri P, Helm DL, et al. Foam pore size is a critical interface parameter of suction-based wound healing devices. *Plastic Reconstr Surg J*. 2012;129:589-597.
57. Rodriguez-Perez MA, Álvarez-Láinez M, de Saja JA. Microstructure and physical properties of open-cell polyolefin foams. *J Appl Polym Sci*. 2009;114:1176-1186.
58. Tjaden B, Brett DJL, Shearing PR. Tortuosity in electrochemical devices: a review of calculation approaches. *Int Mater Rev*. 2018;63:2.
59. Belda R, Palomar M, Marco M, Vercher-Martínez A, Giner E. Open cell polyurethane foam compression failure characterization and its relationship to morphometry. *Mater Sci Eng C Mater Biol Appl*. 2021;120:111754.
60. Pugna A, Negrea R, Linul E, Marsavina L. Is fracture toughness of PUR foams a material property? A statistical approach. *Materials*. 2020;13(21):4868.
61. Kidane A. On the failure and fracture of polymer foam containing discontinuities. *Int Sch Res Notices*. 2013;408596. doi:10.1155/2013/408596
62. Gefen A, Ousey K. Safe and effective wound care during the COVID-19 pandemic. *J Wound Care*. 2020;29(11):622-623. doi:10.12968/jowc.2020.29.11.622
63. Park CC, Choi WS, Lee JN. Effects of hardness and thickness of polyurethane foam midsoles on bending properties of the footwear. *Fibers Polymers*. 2007;8(2):192-197.
64. Rook S, Davies P, Frenthof E, Würfel T. Mepilex® border flex—results of an observational study in German specialist wound care centres. *Wounds Int*. 2019;10(1):40-43.
65. Serena TE, Chadwick P, Davies P, et al. Multifunctional and patient-focused Mepilex border flex: an exploration of its holistic clinical benefits. *J Wound Care*. 2019;28(Sup6b): S1-S31.
66. Fletcher J. Dressings: cutting and application guide. *World Wide Wounds*. 2007. <http://www.worldwidewounds.com/2007/may/Fletcher/Fletcher-Dressings-Cutting-Guide.html>.
67. Klode J, Schöttler L, Stoffels I, Körber A, Schadendorf D, Dissemond J. Investigation of adhesion of modern wound dressings: a comparative analysis of 56 different wound dressings. *J Eur Acad Dermatol Venereol*. 2011;25(8): 933-939.
68. McNichol L, Lund C, Rosen T, Gray M. Medical adhesives and patient safety: state of the science: consensus statements for the assessment, prevention, and treatment of adhesive-related skin injuries. *J Wound Ostomy Continence Nurs*. 2013; 40(4):365-380.
69. Matsumura H, Imai R, Ahmatjan N, et al. Removal of adhesive wound dressing and its effects on the stratum corneum of the skin: comparison of eight different adhesive wound dressings. *Int Wound J*. 2014;11(1):50-54.
70. Rippon M, White R, Davies P. Skin adhesives and their role in wound dressings. *Wounds UK*. 2007;3(4):76-86.
71. Matsumura H, Ahmatjan N, Ida Y, Imai R, Wanatabe K. A model for quantitative evaluation of skin damage at adhesive wound dressing removal. *Int Wound J*. 2013;10(3): 291-294.
72. de Faria MF, Ferreira MBG, Felix MMS, Bessa RMV, Barbosa MH. Prevention of medical adhesive-related skin injury during patient care: a scoping review. *Int J Nurs Stud Adv*. 2022;4:100078. doi:10.1016/j.ijnsa.2022.100078
73. Fumarola S, Allaway R, Callaghan R, et al. Overlooked and underestimated: medical adhesive-related skin injuries. *J Wound Care*. 2020;29(Sup3c):S1-S24.

74. Hitchcock J, Haigh DA, Martin N, Davies S. Preventing medical adhesive-related skin injury (MARSI). *Br J Nurs*. 2021; 30(15):S48-S56.
75. Kim J, Shin Y. Medical adhesive-related skin injury associated with surgical wound dressing among spinal surgery patients: a cross-sectional study. *Int J Environ Res Public Health*. 2021; 18(17):9150.
76. Cutting KF. Impact of adhesive surgical tape and wound dressings on the skin, with reference to skin stripping. *J Wound Care*. 2008;17(4):157-162.
77. Dykes PJ, Heggie R, Hill SA. Effects of adhesive dressings on the stratum corneum of skin. *J Wound Care*. 2001;10:7-10.
78. Waring M, Rippon M, Bielfeldt S, Brandt M. Cell attachment to adhesive dressings: qualitative and quantitative analysis. *Wounds UK*. 2008;4(3):35-47.
79. Drew P, Posnett J, Rusling L. The cost of wound care for a local population in England. *Int Wound J*. 2007;4(2):149-155.
80. Lindholm C, Searle R. Wound management for the 21st century: combining effectiveness and efficiency. *Int Wound J*. 2016;13:5-15.
81. Tiscar-González V, Menor-Rodríguez MJ, Rabadán-Sainz C, et al. Clinical and economic impact of wound care using a polyurethane foam multilayer dressing. *Adv Skin Wound Care*. 2021;34(1):23-30.
82. Tickle J. Positive clinical and patient outcomes with a next generation foam dressing. *Wounds UK*. 2016; EWMA Special Issue:56-62.
83. Woo KY, Coutts PM, Price P, Harding K, Sibbald RG. A randomized crossover investigation of pain at dressing change comparing 2 foam dressings. *Adv Skin Wound Care*. 2009; 22(7):304-310.
84. Taroc AM. A guide for adhesive removal: principles, practice, and products. *Am Nurse Today*. 2017;12(10):24-26.
85. Bernatchez SF, Bichel J. The science of skin: measuring damage and assessing risk. *Adv Wound Care*. 2022; in press. doi:10.1089/wound.2022.0021
86. Krueger EM, Cullum ME, Nichols TR, Taylor MG, Sexton WL, Murahata RI. Novel instrumentation to determine peel force in vivo and preliminary studies with adhesive skin barriers. *Skin Res Technol*. 2013;19(4):398-404.
87. Renvoise J, Burlot D, Marin G, Derail C. Adherence performances of pressure sensitive adhesives on a model viscoelastic synthetic film: a tool for the understanding of adhesion on the human skin. *Int J Pharm*. 2009;368(1-2):83-88.
88. Williams TP, Snyder CL, Hancock KJ, et al. Development of a low-cost, high-fidelity skin model for suturing. *J Surg Res*. 2020;256:618-622.
89. Pilkey WD, Pilkey DF, Bi Z. *Peterson's Stress Concentration Factors*. 4th ed. Hoboken, NJ: Wiley; 2020:640.
90. Borgquist O, Gustafson L, Ingemansson R, Malmsjö M. Tissue ingrowth into foam but not into gauze during negative pressure wound therapy. *Wounds*. 2009;21(11):302-309.
91. Malmsjö M, Ingemansson R. Green foam, black foam or gauze for NWPT: effects on granulation tissue formation. *J Wound Care*. 2011;20(6):294-299.
92. Kohta M, Nakamura Y, Yunoki S. The effectiveness of topical aid sliding sheet potentially used for pressure injury treatment. *Chronic Wound Care Manage Res*. 2021;8:1-11. doi:10.2147/CWCMR.S284180
93. Vilhena L, Ramalho A. Friction of human skin against different fabrics for medical use. *Lubricants*. 2016;4:6. doi:10.3390/lubricants4010006
94. Gefen A, Alves P, Creehan S, Call E, Santamaria N. Computer modeling of prophylactic dressings: an indispensable guide for healthcare professionals. *Adv Skin Wound Care*. 2019; 32(7S):S4-S13.
95. Grigatti A, Gefen A. What makes a hydrogel-based dressing advantageous for the prevention of medical device-related pressure ulcers. *Int Wound J*. 2022;19(3):515-530.
96. Chessin N, Driver WE. Compression and friction properties of rigid polyurethane foams. *J Cellular Plastics*. 1967;3: 185-191.
97. Schwartz D, Magen YK, Levy A, Gefen A. Effects of humidity on skin friction against medical textiles as related to prevention of pressure injuries. *Int Wound J*. 2018;15(6):866-874.
98. Zhang ZX, Zhang T, Wang D, Zhang X, Xin Z, Prakashan K. Physicomechanical, friction, and abrasion properties of EVA/PU blend foams foamed by supercritical nitrogen. *Polymer Eng Sci*. 2018;5:673-682.
99. Rotaru GM, Pille D, Lehmeier FK, et al. Friction between human skin and medical textiles for decubitus prevention. *Tribol Int*. 2013;65:91-96.
100. Zeng H, Pesika N, Tian Y, et al. Frictional adhesion of patterned surfaces and implications for gecko and biomimetic systems. *Langmuir*. 2009;25(13):7486-7495.
101. Evans RL, Marriott RE, Harker M. Axillary skin: biology and care. *Int J Cosmet Sci*. 2012;34(5):389-395.
102. Ohura N, Ichioka S, Nakatsuka T, Shibata M. Evaluating dressing materials for the prevention of shear force in the treatment of pressure ulcers. *J Wound Care*. 2005;14(9):401-404.
103. Waring M, Butcher M. An investigation into the conformability of wound dressings. *Wounds UK*. 2011;7(3):14-24.
104. Kuehlmann B, Bonham CA, Zucal I, Prantl L, Gurtner GC. Mechanotransduction in wound healing and fibrosis. *J Clin Med*. 2020;9(5):1423.
105. Orlov A, Gefen A. Effective negative pressure wound therapy for open wounds: the importance of consistent pressure delivery. *Int Wound J*. 2022c; in press. doi:10.1111/iwj.13879
106. ASTM D903-98. *Standard Test Method for Peel or Stripping Strength of Adhesive Bonds*. West Conshohocken, PA: ASTM International, Book of Standards Volume 15.06; 2017. doi:10.1520/D0903-98R17
107. Dąbrowska AK, Rotaru GM, Derler S, et al. Materials used to simulate physical properties of human skin. *Skin Res Technol*. 2016;22:3-14.
108. Kheyfets VO, Thornton RC, Kowal M, Finol EA. A protocol for measuring pull-off stress of wound-treatment polymers. *J Biomech Eng*. 2014;136(7):745011-745015.
109. El Gholmy SH. Performance and testing of adhesive bandage tape. *J Eng Fibers Fabrics*. 2019;14:1-5. doi:10.1177/1558925019843712
110. Gunashekar S, Pillai KM, Church BC, Abu-Zahra NH. Liquid flow in polyurethane foams for filtration applications: a study on their characterization and permeability estimation. *J Porous Mater*. 2015;22:749-759.
111. Saint-Michel F, Chazeau L, Cavaillé JY, Chabert E. *Mechanical Properties of High Density Polyurethane Foams: I. Effect of the Density*. *Composites Science and Technology*. Vol 66. Amsterdam: Elsevier; 2006:2700-2708.

112. Goods SH, Neuschwanger CL, Whinnery LL. Mechanical properties of a structural polyurethane foam and the effect of particulate loading. *Mater Res Soc Symp Proc.* 1998;521:15-20. doi:[10.1557/proc-521-15](https://doi.org/10.1557/proc-521-15)
113. Dawson MA, Germaine JT, Gibson LJ. Permeability of open-cell foams under compressive strain. *Int J Solids Struct.* 2007;44(16):5133-5145.
114. Dubrovskii SA, Lagutina MA, Kazanskii KS. Method of measuring the swelling pressure of superabsorbent gels. *Polymer Gels Netw.* 1994;2:49-58.
115. Allen R, Sun S. Computing and comparing effective properties for flow and transport in computer-generated porous media. *Geofluids.* Special Issue on Flow and Transport in Porous Media: Multiscale Focus; 2017:4517259.
116. Clark MM. *Transport Modeling for Environmental Engineers and Scientists.* 2nd ed. Hoboken, NJ: John Wiley & Sons; 2009.
117. Shen L, Chen Z. Critical review of the impact of tortuosity on diffusion. *Chem Eng Sci.* 2007;62:3748-3755.

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APPENDIX A: FORMULATION AND DERIVATION OF THE THEORY OF FOAM MECHANICAL PERFORMANCE

A.1 | The relationship between the porosity of foams and their strength and stiffness

Porosity in a foam dressing material is defined as the ratio of the volume of the pores in the foam sample to the total sample volume. The latter is determined by dividing the mass of the material sample by the density of the solid (ie, the specific polymer material type) used. The difference in the porous sample and solid volumes yields the void or pore volume in the foam sample, which is then used to determine the porosity of the said foam sample. The porosity, \emptyset , is hence determined using the equation:

$$\emptyset = \frac{V_d - V_s}{V_d}, \quad (\text{A1})$$

where V_d is the volume of the foam sample, and V_s is the volume of the solid within (ie, of the un-foamed polymer). Hence, $0 < \emptyset < 1$ defines a material with varying levels of porosity; a material with no pores at all has $\emptyset = 0$.

There are several factors that shape the microarchitecture, including the porosity of foams, and thereby the mechanical properties of foam materials such as the chemistry of the solid and the foaming process defined by the manufacturing protocol.¹¹⁰ Specifically, in the production of polyurethane foams, the foaming and gelling reactions are critical for cell nucleation, growth, distribution and collapse and will therefore eventually determine the porosity \emptyset of the finished foam product.¹¹⁰ Indeed, porosity values of foam dressings differ remarkably, as demonstrated by Lee et al³² who investigated the microstructure of 11 commercial foam-based dressing types. For example, Lee et al³² demonstrated that dressings have non-homogeneous pore sizes and morphologies, and that the pore sizes in a cross-section range from approximately 100 to 1000 μm , that is, by an order of magnitude across products, which is a surprisingly immense variation.

Both the strength σ and the stiffness E of foam materials tend to decrease with an increase in the foam porosity level \emptyset , and this can be described phenomenologically, for example, using power laws:

$$\sigma = \sigma_0(1 - \emptyset)^A, \quad (\text{A2})$$

$$E = E_0(1 - \emptyset)^B, \quad (\text{A3})$$

where σ_0 and E_0 are the strength and stiffness properties of the fully dense material, respectively (ie, where $\emptyset = 0$), and $A > 1$ and $B > 1$ are empirical constants. It is

shown that when the porosity approaches unity $\emptyset \rightarrow 1$, the strength and stiffness both diminish to zero, that is, the denser the foam, the stiffer and stronger it becomes.¹¹¹ Now recalling that porosity is also related to the density of the porous material (ie, the foam in the dressing), such that:

$$\emptyset = 1 - \frac{\rho_d}{\rho_s}, \quad (\text{A4})$$

where ρ_d is the volume of the dressing sample and ρ_s is the volume of the solid (so that if, hypothetically, $\rho_d = \rho_s$ then $\emptyset = 0$). Lee et al³² reported that ρ_d is in the range of 0.06 to 0.26 g/cm^3 (for the 11 different products that they had investigated). Considering that the density of the solid polyurethane material is 1.20 g/cm^3 , foam dressings have porosities \emptyset in the 0.78 to 0.95 range, with a mean porosity of 0.87, although Heit et al⁵⁶ reported that slightly greater foam porosities (up to 0.975) can also be found in foam dressing materials.^{54,56} By substituting Equation (A4) in Equations (A2 and A3), we obtain¹¹²:

$$\sigma = \sigma_0 \left(\frac{\rho_d}{\rho_s} \right)^A. \quad (\text{A5})$$

$$E = E_0 \left(\frac{\rho_d}{\rho_s} \right)^B. \quad (\text{A6})$$

Now further considering that a foam dressing may also be subjected to sustained deformations, for example, because of bodyweight forces or under compression bandaging, and also assuming that a dry dressing material can be described as isotropic, the apparent foam density under uniaxial compression ρ_d^* , where the strain ε is taken to be negative in compression, is given as follows¹¹³:

$$\rho_d^* = \frac{\rho_d}{(1 + \varepsilon)(1 - \nu\varepsilon)^2}, \quad (\text{A7})$$

where ν is Poisson's ratio of the foam (note that for zero strain the apparent density is unaffected). For large compressive strains, the cells in open-cell flexible foams buckle and collapse without expanding laterally, so that their Poisson's ratio in the large deformation regime is approximately zero. Since the current analyses primarily concern the behaviours of low-density (high-porosity), open-cell, flexible foams (typical to wound dressings), and under high compressive strains, $\nu\varepsilon$ is taken here to be zero. Hence, the apparent density of a compressed foam is non-linearly proportional to the strain magnitude, and so will double when the foam is subjected to

50% strain ($\varepsilon = -0.5$), or triple when the foam is subjected to 67% strain ($\varepsilon = -0.67$). Substituting the effective density in a foam dressing subjected to deformations, ρ_d^* , in Equation (A6) for large strains, yields:

$$E = E_0 \left(\frac{\rho_d}{(1 + \varepsilon)\rho_s} \right)^B = E_0 \left(\frac{1 - \emptyset}{1 + \varepsilon} \right)^B. \quad (\text{A8})$$

A.2 | Factors affecting the state of peri-wound stresses during removal of dressings

Consider a dressing with width D that is removed from above a wound with width W , using a force F applied at an angle θ relative to the wound surface through a strip of width d (Figure 4B-D). If we assume that forces are not directly transferred to the wound bed (as they should in good wound care practice), so that the force only distributes on the peri-wound skin, the effective area for transfer of the peeling force will be $(D - W)d$. The normal (σ) and shear (τ) stress concentrations that this generate on the peri-wound skin are therefore as follows:

$$\sigma = \frac{F \sin(\theta)}{(D - W)d}. \quad (\text{A9})$$

$$\tau = \frac{F \cos(\theta)}{(D - W)d}. \quad (\text{A10})$$

It is natural to first pull the edge of the dressing upwards to detach it (ie, so that $\theta \rightarrow 90^\circ$), which maximises the normal stress on the peri-wound as the edge of the dressing detaches (Figure 4D; top frame of computational simulation). At that region of the edge of the dressing, $W = 0$, and therefore, the normal stress concentration is simply F/Dd . However, to progress the peeling process further, the force must take a lower angle, and hence, over the wound, the shear stress τ increases and maximises when $(D - W)$ minimises, that is, at the widest region of the wound.

A.3 | Swelling of foam in a confined space

Dubrovskii et al¹¹⁴ demonstrated that the swelling pressure p during swelling of a porous material in a confined space depends on the time t according to the following equation:

$$p = C \left[1 - \exp\left(\frac{-t}{\tau}\right) \right], \quad (\text{A11})$$

where C is an empirical constant and the characteristic swelling time $\tau = \frac{R^2}{2\pi^2 D_{\text{eff}}}$ where R is the radius of the material specimen and D_{eff} is the effective diffusion coefficient in the porous material. The effective diffusion coefficient D_{eff} depends on the porosity ($0 < \emptyset < 1$) and tortuosity ($\varphi > 1$) of the dressing material such that¹¹⁵⁻¹¹⁷:

$$D_{\text{eff}} = \frac{D_{\text{pore}}\emptyset}{\varphi}, \quad (\text{A12})$$

where D_{pore} is the diffusion coefficient of the fluid flowing through a pore, \emptyset is the porosity of the dressing material ($0 < \emptyset < 1$) and φ is the tortuosity of the connections between the pores ($\varphi \geq 1$). Intuitively, the more tortuous the porous foam dressing material, the slower the rate of the effective diffusion (ie, the material has a lower D_{eff}). Equation (A11) indicates that in confined swelling, the pressure plateaus to a steady state. By substituting the explicit term for the characteristic swelling time and the effective diffusion coefficient (Equation A12) into Equation (A11) we obtain:

$$p = C \left[1 - \exp\left(\frac{-2\pi^2 D_{\text{pore}}\emptyset t}{\varphi R^2}\right) \right]. \quad (\text{A13})$$

The latter Equation (A13) demonstrates that the swelling pressure of a foam-based dressing in the wound bed depends on the elapsed time post-application of the dressing and the effective diffusion coefficient of the foam, which in turn depends on the porosity and tortuosity of the foam.