1 2	Clinical Practice Guideline:	Wound Care
3	Date of Implementation:	October 18, 2012
4 5	Product:	Specialty
6		

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1	PRACTITIONER SCOPE AND TRAINING
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3	
4	GUIDELINES
5	Wound Debridement
6	Wound care is defined as the care of wounds that are refractory to healing or have
7	complicated healing cycles either because of the nature of the wound itself or because of
8	complicating metabolic and/or physiological factors. This definition excludes management
9	of acute wounds, the care of wounds that normally heal by primary intention such as clean,
10	incised traumatic wounds, surgical wounds that are closed primarily and other
11	postoperative wound care not separately payable during the surgical global period.
12	
13	American Specialty Health – Specialty (ASH) would expect that wound care may be
14	medically necessary for the following types of wounds as indicated by appropriate
15	documentation in support of medical necessity:
16	 Second- and third-degree burn wounds Surgical wounds that must be left open to head by secondary intention
17	• Surgical wounds that must be left open to heal by secondary intention
18	 Infected open wounds induced by trauma or surgery Wounds associated with complicating outsimersurger metabolic secondaries and the secondaries of the secondar
19 20	 Wounds associated with complicating autoimmune, metabolic, vascular or pressure factors
20	
21 22	• Open or closed wounds complicated by necrotic tissue and eschar
22	Documentation to support selective debridement (CPT® Codes 97597 and 97598) must
23 24	include the following to support medical necessity:
25	 Clear description of instruments used for debridement (e.g., high-pressure waterjet,
26	scissors, scalpel, forceps)
27	• Thorough objective assessment of the wound including drainage, color, texture,
28	temperature, vascularity, condition of surrounding tissue, and size of the area to be
29	targeted for debridement
30	• Description of adjunctive measures to support debridement procedures, if indicated
31	(e.g., management of pressure (e.g., off-loading, padding, appropriate footwear),
32	infection, vascular insufficiency, metabolic disorder, and/or nutritional deficiency)
33	• Documentation of complexity of skills required by treating practitioner indicated
34	in medical record
35	
36	Documentation to support non-selective debridement (CPT® 97602) must include the
37	following to support medical necessity:
38	• Type of technique utilized (i.e., wet-to-moist, enzymatic, abrasion)
39 40	• Thorough objective assessment of the wound including drainage, color, texture,
40	temperature, vascularity, condition of surrounding tissue, and size of the area to be
41	targeted for debridement

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• Description of adjunctive measures to support debridement procedures, if indicated (i.e., management of pressure (i.e., off-loading, padding, appropriate footwear), infection, vascular insufficiency, metabolic disorder, and/or nutritional deficiency)

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- Documentation of complexity of skills required by treating practitioner indicated in medical record
- 5 6

If there is no documented evidence (e.g., objective measurements) of ongoing significant 7 benefit, then the medical record documentation must provide other clear evidence of 8 9 medical necessity for treatments. Physicians and qualified non-physician practitioners, licensed physical therapists and licensed occupational therapists acting within their scope 10 of practice and licensure may provide debridement services and use the Physical Medicine 11 and Rehabilitation codes including CPT® 97597, 97598 and 97602. Removal of non-tissue 12 integrated fibrin exudates, crusts, biofilms, or other materials from a wound without 13 removal of tissue does not meet the definition of any debridement code and may not be 14 reported as such. 15

16

Debridement of the wound(s) when indicated must be performed discriminately and at appropriate intervals. Prolonged, repetitive debridement services require adequate documentation of complicating circumstances that reasonably necessitated additional services. ASH expects that with appropriate care, wound volume or surface dimension should decrease by at least 10 percent per month or wounds will demonstrate margin advancement of no less than 1 mm/week. ASH expects the wound-care treatment plan to be modified in the event that appropriate healing is not achieved.

24

Medically necessary chronic wound care must be performed in accordance with accepted 25 standards for medical and surgical treatment of wounds. Eventual wound closure with or 26 without grafts, skin replacements or other surgery (such as amputation, wound excision, 27 etc.) should be the goal of most chronic wound care. Isolated wound care, when other 28 adjunctive measures are indicated, is not considered to be medically necessary. With 29 appropriate management, it is expected that, in most cases, a wound will reach a state at 30 which its care should be performed primarily by the patient and/or the patient's caregiver 31 with periodic physician assessment and supervision. Wound care that can be performed by 32 the patient or the patient's caregiver will be considered to be maintenance care and not 33 medically necessary. 34

35

ASH considers CPT® code 17250 (Chemical cauterization of granulation tissue (proud flesh, sinus, or fistula)) an integral service as part of a health care provider's medical or surgical care and not separately billable with debridement CPT® codes in the table below.

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1 Evaluation/Re-assessment

2 Other than an initial evaluation, wound assessment is an integral part of all wound care 3 service codes and, as such, these assessments are not separately billable.

- Initial wound assessments that are medically necessary may be reimbursable as a separately identifiable Evaluation and Management (E/M) service or i.e., physical therapy evaluation CPT® 97161-97163.
- Re-assessments/re-evaluations of a wound (which may be completed with a dressing change) are considered to be a non-covered routine service. An exception would require documentation clearly supporting that there had been a significant improvement, decline, or change in the patient's condition or functional status that was not anticipated in the plan of care and required further evaluation.

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13 **CPT® Codes and Descriptions**

CPT® Code	CPT® Code Description
97597	Debridement (e.g., high pressure waterjet with/without suction, sharp selective debridement with scissors, scalpel and forceps), open wound, (e.g., fibrin, devitalized epidermis and/or dermis exudate, debris, biofilm), including topical application(s), wound assessment, use of a whirlpool, when performed and instructions (s) for ongoing care, per session, total wound(s) surface area; first 20 sq cm or less
97598	Debridement (e.g., high pressure waterjet with/without suction, sharp selective debridement with scissors, scalpel and forceps), open wound, (e.g., fibrin, devitalized epidermis and/or dermis, exudate, debris, biofilm), including topical application(s), wound assessment, use of a whirlpool, when performed and instruction(s) for ongoing care, per session, total wound(s) surface area; each additional 20 sq cm, or part thereof (List separately in addition to code for primary procedure)
97602	Removal of devitalized tissue from wound(s), non-selective debridement, without anesthesia (e.g., wet-to-moist dressings, enzymatic, abrasion, larval therapy), including topical application(s), wound assessment, and instruction(s) for ongoing care, per session
17250	Chemical cauterization of granulation tissue (i.e. proud flesh)

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1	Woun	nd Care Modalities
2	A.	Whirlpool
3		• If the patient uses whirlpool for treatment of a wound prior to receiving
4		selective debridement services for the wound during the same visit, then the
5		whirlpool is not separately reimbursable and should not be billed with modifier
6		59 unless two separate wounds are treated with different modalities.
7		• If the patient uses whirlpool for treatment of a wound prior to receiving non-
8		selective debridement services for the wound during the same visit, then the
9		whirlpool is separately reimbursable and may be billed with modifier 59.
10		• Whirlpool can also be completed during the same visit for non-wound care-
11		related purposes. It is appropriate to separately bill CPT® 97022 when the
12		whirlpool is used for other purposes not involving wound care, e.g., facilitation
13		of range of motion activities.
14		
15	В.	Electrical Stimulation Therapy
16		Care of chronic Stage III and Stage IV pressure ulcers, arterial ulcers, diabetic
17		ulcers and/or venous stasis ulcers through use of Electrical Stimulation (ES)
18		(electrical current via electrodes placed directly on the skin in close proximity to
19		the ulcer; CPT®/HCPCS codes G0281, 97014, 97032) may be covered as
20		medically necessary when the following criteria are met:
21		• Patient is a Medicare beneficiary; AND
22		• Failure to demonstrate measurable signs of healing (e.g., signs of
23		epithelialization and reduction in ulcer size) with a 30-day trial of conventional
24		wound management, including optimization of nutritional status, moist
25		dressings, and debridement. ES would not be medically necessary as an initial
26		treatment modality.
27		Other considerations.
28		Other considerations:
29 20		• If after 30 days of ES therapy no measurable signs of healing (e.g., decrease in wound size/ourface or volume, decrease in amount of available and decrease in
30 31		wound size/surface or volume, decrease in amount of exudates and decrease in
		amount of necrotic tissue) are demonstrated, ES should be discontinued.
32 22		• ES treatment sessions are not medically necessary beyond one hour. Prolonged
33		treatments using ES do not provide additional benefit.
34 25		• ES also must be discontinued when the wound demonstrates a 100 percent
35 36		epithelialized wound bed.
36 37		• ASH considers ES therapy for chronic ulcers unproven when these criteria are not met (e.g., not a Medicare beneficiary)
37		not met (e.g., not a Medicare beneficiary).
38 39		• Additionally, comprehensive wound treatments must include optimization of nutritional status, debridement to remove devitalized tissue, maintenance of a
39 40		clean, moist bed of granulation tissue with appropriate moist dressings, and
		necessary care to resolve any infection that may be present. Specific wound
41		necessary care to resolve any infection that may be present. Specific wound

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care based on type of wound includes frequent repositioning of a member with pressure ulcers (usually every 2 hours); off-loading of pressure and good glucose control for diabetic ulcers; establishment of adequate circulation for arterial ulcers and the use of a compression system for members with venous ulcers.

C. Electromagnetic Therapy

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Care of chronic Stage III and Stage IV pressure ulcers, arterial ulcers, diabetic ulcers and/or venous stasis ulcers through use of Electromagnetic (EM) therapy (pulsed magnetic field to induce current) may be covered as medically necessary when the following criteria are met:

- Patient is a Medicare beneficiary; AND
- Failure to demonstrate measurable signs of healing (e.g., signs of epithelialization and reduction in ulcer size) with a 30-day trial of conventional wound management, including optimization of nutritional status, moist dressings, and debridement. EM would not be medically necessary as an initial treatment modality.
 - Other considerations:
- If after 30 days of EM therapy no measurable signs of healing (e.g., decrease in wound size/surface or volume, decrease in amount of exudates and decrease in amount of necrotic tissue) are demonstrated, EM should be discontinued.
 - EM treatment sessions are not medically necessary beyond one hour. Prolonged treatments using EM do not provide additional benefit.
- EM also must be discontinued when the wound demonstrates a 100 percent epithelialized wound bed.
- ASH considers EM therapy for chronic ulcers unproven when these criteria are not met (e.g., not a Medicare beneficiary).
- Additionally, comprehensive wound treatments must include optimization of 29 • nutritional status, debridement to remove devitalized tissue, maintenance of a 30 clean, moist bed of granulation tissue with appropriate moist dressings, and 31 necessary care to resolve any infection that may be present. Specific wound 32 care based on type of wound includes frequent repositioning of a member with 33 pressure ulcers (usually every 2 hours); off-loading of pressure and good 34 glucose control for diabetic ulcers; establishment of adequate circulation for 35 arterial ulcers and the use of a compression system for members with venous 36 ulcers. 37

1 D .	Ultraviolet (UV) Light
2	ASH considers the treatment of decubitus ulcers with CPT® code 97028 – UV light
3	NOT medically necessary, except in the following circumstance where it may be
4	reasonable and necessary:
5	• For Medicare beneficiaries requiring the application of a drying heat, such as
6	for the treatment of severe psoriasis where there is limited range of motion.
7	• Supportive Documentation Requirements (required at least every 10
8	visits)
9	 Area(s) being treated
10	 Objective clinical findings/measurements to support the need for
11	ultraviolet
12	 Minimal erythema dosage
13	
	Low-Frequency, Non-Contact, Non-Thermal Ultrasound
15	CPT® code 97610 [low frequency, non-contact, non-thermal ultrasound, including
16	topical application(s) when performed, wound assessment, and instruction(s) for
17	ongoing care, per day] describes a system that uses continuous low-frequency
18	ultrasonic energy to produce and propel a mist of liquid and deliver continuous low- frequency ultrasound to the wound had. This modelity is often referred to as MIST .
19 20	frequency ultrasound to the wound bed. This modality is often referred to as 'MIST Therapy.'
20 21	merapy.
21	Low-frequency, non-contact, non-thermal ultrasound (MIST Therapy) may be
22	covered as medically necessary wound therapy for Medicare beneficiaries for any
24	of the following clinical conditions:
25	• Wounds, burns and ulcers meeting ASH medical necessity criteria for
26	debridement, but which are too painful for sharp or excisional debridement and
27	described in the medical record
28	• Wounds, burns and ulcers meeting ASH medical necessity criteria for
29	debridement but with documented contraindications to sharp or excisional
30	debridement
31	• Wounds, burns and ulcers meeting ASH medical necessity criteria for
32	debridement but with documented evidence of no signs of improvement after
33	30 days of standard wound care
34	•
35	Other considerations:
36	• Low-frequency, non-contact, non-thermal ultrasound (MIST Therapy) must be
37	provided two to three times per week to be considered medically necessary
38	• The length of individual treatments will vary per wound size

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1	• Observable, documented improvements in the wound(s) should be evident after
2	six treatments. Improvements include documented reduction in pain, necrotic
3	tissue, or wound size or improved granulation tissue
4	• Continuing treatments are not covered for wounds demonstrating no
5	improvement after 6 treatments
6	• MIST therapy is considered unproven and not a covered service for non-
7	Medicare patients
8	
9	F. Ultrasound
10	ASH considers care of chronic wounds through use of therapeutic Ultrasound;
11	CPT® code 97035) medically necessary based on the following criteria:
12	• Failure to demonstrate measurable signs of healing (e.g., signs of
13	epithelialization and reduction in ulcer size) with a 30-day trial of conventional
14	wound management, including optimization of nutritional status, moist
15	dressings, and debridement. Ultrasound would not be medically necessary as
16	an initial treatment modality.
17	C. Low Lovel Leger Thereny
18	G. Low Level Laser Therapy
19 20	ASH considers Low Level Laser Therapy unproven for treatment of chronic
20 21	wounds. There is insufficient evidence to support its use.
21 22	Dressing Use and Change
22	Application of wound dressing continues to be the standard of care for wound treatment;
23 24	however, the literature is inconclusive as it relates to standardized topical preparations and
25	types of dressings. Documentation must support the use of the type of dressing for bandage.
23 26	Dressing size must be based on and appropriate to the size of the wound. For wound covers,
20	the pad size is usually about 2 in. greater than the dimensions of the wound. For example,
28	a 5 cm x 5 cm (2 in. x 2 in.) wound requires a 4 in. x 4 in. pad size.
29	
30	The quantity and type of dressings dispensed at any one time must consider the status of
31	the wound(s), the likelihood of change, and the recent use of dressings. Dressing needs
32	may change frequently (e.g., weekly) in the early phases of wound treatment and/or with
33	heavily draining wounds. Suppliers are also expected to have a mechanism for determining
34	the quantity of dressings that the patient is using and to adjust their provision of dressings
35	accordingly. No more than a one month's supply of dressings may be provided at one time
36	unless there is documentation to support the necessity of greater quantities in the home
37	setting in an individual case. An even smaller quantity may be appropriate in the situations
38	described above.
39	
40	Surgical dressings must be tailored to the specific needs of an individual patient. When
41	surgical draggings are provided in kits, only those components of the kit that most the

- surgical dressings are provided in kits, only those components of the kit that meet the 41 42
 - definition of a surgical dressing, that are ordered by the physician, and that are medically

1 necessary are covered. Most compression bandages are reusable. Usual frequency of

replacement would be no more than one per week unless they are part of a multi-layercompression bandage system.

4

Multi-layered, sustained, graduated, high compression bandage systems are used primarily
 to treat lymphedema and venous or stasis leg ulcers. Several graduated, high-compression
 bandage systems products have been developed, including Profore®, Dyna-Flex®,

- 8 Surepress[®], Setopress[®], and other similar product systems.
- 9

HCPCS/ CPT® Code	HCPCS/ CPT® Code Description
	Light compression bandage, elastic, knitted/woven, width less than 3 inches, per yard
	Light compression bandage, elastic, knitted/woven, width greater than or equal to 3 inches and less than 5 inches, per yard
14.6/150	Light compression bandage, elastic, knitted/woven, width greater than or equal to 5 inches, per yard
	Application of multi-layer compression system; leg (below knee), including ankle and foot

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11 A dressing change may not be billed as either a debridement or other wound care service 12 under any circumstance (e.g., CPT® 97597, 97598, 97602).

Medicare does not separately reimburse for dressing changes or patient/caregiver training in the care of the wound. These services are reimbursed as part of a billable
 E/M or procedure code that, commonly but not necessarily, occurs on the same date of service as the dressing change. If not included in another service, the costs associated with dressing changes may be reported as not separately payable.

19 20

21 Surgical Debridement

•

22 Debridement, Subcutaneous Tissue, Muscle and/or Fascia

in the payment for the procedure codes.

ASH considers services consisting of CPT® Codes 11042, 11043, 11045, and 11046 to be medically necessary for the debridement of muscle and/or subcutaneous tissue upon meeting **ALL of** the following criteria (1, 2, and 3) below:

- 26
- 27
- 28
- 1. Conditions that may require debridement include at least one of the following:

All topical applications (e.g., medications, ointments, and dressings) are included

ICD-10 Code	ICD-10 Code Description
170.232, 170.242	Atherosclerosis of native arteries of leg with ulceration of calf

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ICD-10 Code	ICD-10 Code Description
170.233, 170.243	Atherosclerosis of native arteries of leg with ulceration of ankle
170.234, 170.244	Atherosclerosis of native arteries of leg with ulceration of heel and midfoot
170.235, 170.245	Atherosclerosis of native arteries of leg with ulceration of other part of foot
I70.238 - I70.239, I70.248 - I70.249	Atherosclerosis of native arteries of leg with ulceration of other part of lower leg or unspecified site
170.25	Atherosclerosis of native arteries of other extremities with ulceration
170.332, 170.342, 170.432, 170.442, 170.532, 170.542, 170.632, 170.642, 170.732, 170.742	Atherosclerosis of bypass graft(s) of the leg with ulceration of calf
I70.333, I70.343, I70.433, I70.443, I70.533, I70.543, I70.633, I70.643, I70.733, I70.743	Atherosclerosis of bypass graft(s) of the leg with ulceration of ankle
I70.334, I70.344, I70.434, I70.444, I70.534, I70.544, I70.634, I70.644, I70.734, I70.744	Atherosclerosis of bypass graft(s) of the leg with ulceration of heel and midfoot
I70.335, I70.345, I70.435, I70.445, I70.535, I70.545, I70.635, I70.645, I70.735, I70.745	Atherosclerosis of bypass graft(s) of the leg with ulceration of other part of foot

ICD-10 Code	ICD-10 Code Description
I70.338 - I70.339, I70.348 - I70.349, I70.438 - I70.439, I70.438 - I70.439, I70.538 - I70.539, I70.548 - I70.549, I70.638 - I70.639, I70.648 - I70.649, I70.738 - I70.739, I70.748 - I70.749	Atherosclerosis of bypass graft(s) of the leg with ulceration of other part of lower leg or unspecified site
170.35, 170.45, 170.55, 170.65, 170.75	Atherosclerosis of bypass graft(s) of other extremity with ulceration
L02.415 - L02.419, L03.115 - L03.119, L03.125 - L03.129	Cutaneous abscess, cellulitis, and acute lymphangitis of lower and unspecified part of limb
L02.611 - L02.619	Cutaneous abscess of foot
L08.81, L08.89	Pyoderma vegetans - Other specified local infections of the skin and subcutaneous tissue
L08.9	Local infection of the skin and subcutaneous tissue, unspecified
L89.200, L89.210, L89.220, L89.300, L89.310, L89.320, L89.500, L89.510, L89.520, L89.600, L89.610, L89.620, L89.890, L89.95	Pressure ulcer of hip, buttock, ankle, heel, other site, and unspecified site; unstageable
L89.204, L89.214, L89.224, L89.304, L89.314, L89.324, L89.504, L89.514, L89.524, L89.604, L89.614, L89.624, L89.894, L89.94	Pressure ulcer of hip, buttock, ankle, heel, other site, and unspecified site; stage 4

ICD-10 Code	ICD-10 Code Description
L89.209, L89.219, L89.229, L89.309, L89.319, L89.329, L89.509, L89.519, L89.529, L89.609, L89.619, L89.629, L89.899, L89.90	Pressure ulcer of hip, buttock, ankle, heel, other site, and unspecified site; unspecified stage
L89.500 - L89.529	Pressure ulcer of ankle
L89.600 - L89.629	Pressure ulcer of heel
L89.890 - L89.899	Pressure ulcer of other site
L89.90 - L89.95	Pressure ulcer of unspecified site
L97.201 - L97.229	Non-pressure chronic ulcer of calf
L97.301 - L97.329	Non-pressure chronic ulcer of ankle
L97.401 - L97.429	Non-pressure chronic ulcer of heel and midfoot
L97.501 - L97.529	Non-pressure chronic ulcer of other part of foot
L97.801 - L97.829	Non-pressure chronic ulcer of other part of lower leg
L97.901 - L97.929	Non-pressure chronic ulcer of unspecified part of lower leg
L98.411 - L98.419	Non-pressure chronic ulcer of buttock
L98.491 - L98.499	Non-pressure chronic ulcer of skin of other sites
M72.6	Necrotizing fasciitis

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2. All significant relevant comorbid conditions are addressed that could interfere with optimal wound healing.

- 3. If there is no necrotic, devitalized, fibrotic, or other tissue or foreign matter present that would interfere with wound healing, the debridement service is not medically necessary. The presence or absence of such tissue or foreign matter must be documented in the medical record.
- 7 8

9 The number of debridement services required is variable and depends on numerous 10 intrinsic and extrinsic factors. Debridement of the wound(s) when indicated must be 11 performed discriminately and at appropriate intervals. ASH expects fewer than five 12 debridement sessions involving removal of muscle to be required for management of most

wounds. Prolonged, repetitive debridement services require adequate documentation of 1 complicating circumstances that reasonably necessitated additional services. 2

3

Local infiltration, metacarpal/digital block or topical anesthesia are included in the 4 reimbursement for debridement services and are not separately payable. Anesthesia 5 administered by or incident to the provider performing the debridement procedure is not 6 7 separately payable.

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Exclusion criteria: CPT[®] codes 11042, 11043, 11045, and 11046 are **NOT** appropriate 9 for the following conditions: 10

- Skin breakdown under a dorsal corn is not considered an ulcer and generally does • not require debridement. These lesions typically heal without significant surgical intervention beyond removal of the corn and shoe modification.
- Removing a collar of callus (hyperkeratotic tissue) around an ulcer is not debridement of skin or necrotic tissue.
- It is expected that, with appropriate care, and no extenuating medical or surgical 17 complications or setbacks, wound volume or surface dimension should decrease over time. 18 It is also expected the wound care treatment plan is modified in the event that appropriate 19 healing is not achieved. It is expected that co-morbid conditions that may interfere with 20 normal wound healing have been addressed; the etiology of the wound has been determined 21 and addressed as well as addressing patient compliance issues. This may include, for 22 23 example, evaluation of pulses, ABI and/or possible consultation with a vascular surgeon.
- 24

Debridement, Bone 25

ASH considers services consisting of CPT® Codes 11044 and 11047 to be medically 26 necessary for the debridement of bone upon meeting ALL of the following criteria (1, 2, 27 and 3) below: 28 1. Conditions that may require debridement include at least one of the following:

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30	

ICD-10 Code	ICD-10 Code Description
A18.03	Tuberculosis of other bones
M86.00, M86.10, M86.20	Acute hematogenous, other acute, and subacute osteomyelitis; unspecified site
M86.061 - M86.069, M86.161 - M86.169, M86.261 - M86.269	Acute hematogenous, other acute, and subacute osteomyelitis; tibia and fibula
M86.071 - M86.079, M86.171 - M86.179, M86.271 - M86.279	Acute hematogenous, other acute, and subacute osteomyelitis; ankle and foot

ICD-10 Code	ICD-10 Code Description
M86.08, M86.18, M86.28	Acute hematogenous, other acute, and subacute osteomyelitis; other site
M86.09, M86.19, M86.29	Acute hematogenous, other acute, and subacute osteomyelitis; multiple sites
M86.30, M86.40, M86.50, M86.60	Chronic multifocal, with draining sinus, other chronic hematogenous, and other chronic osteomyelitis; unspecified site
M86.361 - M86.369, M86.461 - M86.469, M86.561 - M86.569, M86.661 - M86.669	Chronic multifocal, with draining sinus, other chronic hematogenous, and other chronic osteomyelitis; tibia and fibula
M86.371 - M86.379, M86.471 - M86.479, M86.571 - M86.579, M86.671 - M86.679,	Chronic multifocal, with draining sinus, other chronic hematogenous, and other chronic osteomyelitis; ankle and foot
M86.38, M86.48, M86.58, M86.68	Chronic multifocal, with draining sinus, other chronic hematogenous, and other chronic osteomyelitis; other site
M86.39, M86.49, M86.59, M86.69	Chronic multifocal, with draining sinus, other chronic hematogenous, and other chronic osteomyelitis; multiple sites
M86.8X0, M86.8X6, M86.8X7, M86.8X8, M86.8X9	Other osteomyelitis; unspecified sites, lower leg, ankle and foot, other site, and multiple sites
M86.9	Osteomyelitis, unspecified
M90.861 - M90.869	Osteopathy in diseases classified elsewhere, lower leg
M90.871 - M90.879	Osteopathy in diseases classified elsewhere, ankle and foot
M90.88	Osteopathy in diseases classified elsewhere, other site
M90.89	Osteopathy in diseases classified elsewhere, multiple sites

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- 2. All significant relevant comorbid conditions are addressed that could interfere with optimal wound healing.
- 3. If there is no necrotic, devitalized, fibrotic, or other tissue or foreign matter present that would interfere with wound healing, the debridement service is not medically necessary. The presence or absence of such tissue or foreign matter must be documented in the medical record.

The number of debridement services required is variable and depends on numerous intrinsic and extrinsic factors. Debridement of the wound(s) when indicated must be performed discriminately and at appropriate intervals. ASH expects fewer than five debridement sessions involving removal of bone to be required for management of most wounds. Prolonged, repetitive debridement services require adequate documentation of complicating circumstances that reasonably necessitated additional services.

7

8 Local infiltration, metacarpal/digital block or topical anesthesia are included in the 9 reimbursement for debridement services and are not separately payable. Anesthesia 10 administered by or incident to the provider performing the debridement procedure is not 11 separately payable.

12

Exclusion criteria: CPT® codes 11044 and 11047 are <u>NOT</u> appropriate for the following
 conditions:

- Skin breakdown under a dorsal corn is not considered an ulcer and generally does not require debridement. These lesions typically heal without significant surgical intervention beyond removal of the corn and shoe modification.
- 18 19
- Removing a collar of callus (hyperkeratotic tissue) around an ulcer is not debridement of skin or necrotic tissue.
- 20 Debridement for osteomyelitis is covered for chronic osteomyelitis and osteomyelitis 21 associated with an open wound. It is expected that, with appropriate care, and no 22 extenuating medical or surgical complications or setbacks, wound volume or surface 23 dimension should decrease over time. It is also expected the wound care treatment plan is 24 modified in the event that appropriate healing is not achieved. It is expected that the 25 etiology of the wound has been determined and addressed as well as addressing patient 26 compliance issues. This may include, for example, evaluation of pulses, ABI and/or 27 possible consultation with a vascular surgeon. 28
- 29

ASH considers CPT® code 17250 (Chemical cauterization of granulation tissue (proud flesh, sinus, or fistula)) an integral service as part of a health care provider's medical or surgical care and not separately billable with surgical debridement CPT® codes listed in the table below.

34 35

<u>CPT®</u> Codes and Descriptions

CPT® Code	CPT® Code Description
11042	Debridement, subcutaneous tissue (includes epidermis and dermis, if performed); first 20 sq cm or less

CPT® Code	CPT® Code Description
11043	Debridement, muscle and/or fascia (includes epidermis, dermis, and subcutaneous tissue, if performed); first 20 sq cm or less
11044	Debridement, bone (includes epidermis, dermis, subcutaneous tissue, muscle and/or fascia, if performed); first 20 sq cm or less
11045	Debridement, subcutaneous tissue (includes epidermis and dermis, if performed); each additional 20 sq cm, or part thereof (List separately in addition to code for primary procedure)
11046	Debridement, muscle and/or fascia (includes epidermis, dermis, and subcutaneous tissue, if performed); each additional 20 sq cm, or part thereof (List separately in addition to code for primary procedure)
11047	Debridement, bone (includes epidermis, dermis, subcutaneous tissue, muscle and/or fascia, if performed); each additional 20 sq cm, or part thereof (List separately in addition to code for primary procedure)
17250	Chemical cauterization of granulation tissue (i.e. proud flesh)

2 Powered Negative Pressure Wound Therapy / Vacuum-Assisted Closu

ASH considers powered negative pressure wound therapy (NPWT)/vacuum-assisted closure (VAC) CPT® code 97605, 97606) (HCPCS code A6550, E2402) medically necessary upon meeting **ALL of** the criteria (1, 2, 3, and 4) below:

6	1.	Individual is 12.0 years of age or older; and
7	2.	A complete wound care program, which meets ALL of the requirements below,
8		has been tried:
9		• Documentation in the individual's medical record of evaluation, care, and
10		wound measurements by a licensed medical professional; and
11		 Application of dressings to maintain a moist environment; and
12		 Debridement of necrotic tissue if present; and
13		• Evaluation of and provision for adequate nutritional status; and
14		• Underlying medical conditions (e.g., diabetes, venous insufficiency) are
15		being appropriately managed; and

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1	3. A	n eligible condition is documented (individual must meet one or more of the
2		following):
3		a. Stage III or IV pressure ulcers (see key terms below) at initiation of vacuum
4		assisted wound therapy, in individuals who meet ALL of the following:
5		i. The individual has been appropriately turned and positioned; and
6		ii. The individual has used a group 2 or 3 support surface for pressure
7		ulcers on the posterior trunk or pelvis (no special support surface is
8		required for ulcers not located on the trunk or pelvis); and
9		iii. The individual's moisture and incontinence have been appropriately
10		managed; or
11		b. Neuropathic ulcers in individuals who meet BOTH of the following:
12		i. The individual has been on a comprehensive diabetic management
13		program; and
14		ii. Reduction in pressure on a foot ulcer has been accomplished with
15		appropriate modalities; or
16		c. Ulcers related to venous or arterial insufficiency, in individuals who meet
17		ALL of the following:
18		i. Compression bandages and/or garments have been consistently applied;
19		and
20		ii. Reduction in pressure on a foot ulcer has been accomplished with
21		appropriate modalities; and
22		iii. For initiation of therapy in the home setting, presence of the ulcer for at
23		least 30 days; or
24		d. Dehisced wounds or wound with exposed hardware or bone; or
25		e. Post sternotomy wound infection or mediastinitis; or
26		f. Complications of a surgically created wound where accelerated granulation
27		therapy is necessary and cannot be achieved by other available topical
28		wound treatment.
29	4.	The wound to be treated is free from ALL of the following absolute
30		contraindications to vacuum assisted wound therapy:
31		a. Exposed anastomotic site; or
32		b. Exposed nerves; or
33		c. Exposed organs; or
34		d. Exposed vasculature; or
35		e. Malignancy in the wound; or
36		f. Necrotic tissue with eschar present; or
37		g. Non-enteric and unexplored fistulas; or
38		h. Untreated osteomyelitis.

h. Untreated osteomyelitis.

1 2	Continued use of electrically powered vacuum assisted wound therapy is considered medically necessary when:
3	• Weekly assessment of the wound's dimensions and characteristics by a licensed
4	health care professional is documented; and
5	 Progressive wound healing is demonstrated.
6	1 rogressive would nearing is demonstrated.
7	Continued use of electrically powered vacuum assisted wound therapy is considered not
8 9	medically necessary when the continuation of treatment criteria above have not been met.
10	NPWT is considered NOT medically necessary for one or more of the following situations:
11 12	• An appropriate health care provider is not supervising or performing weekly wound measurement and assessment functions and documentation, as well as the dressing
13	changes required.
14	• Wound healing has occurred to the extent that NPWT is no longer needed.
15	• The depth of the wound is less than 1 mm, as wounds of this depth cannot
16	accommodate the sponge.
17	• Uniform granulation tissue has been obtained.
18	• The individual cannot tolerate the use of NPWT.
19	• The wound is infected.
20	• There is no progression of healing of the wound on two successive dressing changes
21	and/or up to 30 days.
22	
23	Unproven and Not Medically Necessary:
24	• Electrically powered vacuum assisted wound therapy is considered unproven and
25	not medically necessary for all other applications not meeting the medical necessity
26	criteria above, including when any absolute contraindications to vacuum assisted
27	wound therapy are present.
28	• Non-electrically powered vacuum assisted wound therapy (for example, the
29	SNaP TM Wound Care Device) is considered investigational and not medically
30	necessary for all conditions.
31	• Portable, battery powered, single use (disposable) vacuum assisted wound therapy
32	devices (for example, the PICO [™] Single Use Negative Pressure Wound Therapy
33	System or the V.A.C. Via [™] Negative Pressure Wound Therapy System) are
34	considered investigational and not medically necessary for all conditions.

1 **CPT®/HCPCS Codes and Descriptions**

CPT®/HCPCS Code	CPT® Code Description
97605	Negative pressure wound therapy (e.g., vacuum assisted drainage collection), utilizing durable medical equipment (DME) including topical application(s), wound assessment, and instruction(s) for ongoing care, per session; total wound(s) surface area less than or equal to 50 square centimeters
97606	Negative pressure wound therapy (e.g., vacuum assisted drainage collection), utilizing durable medical equipment (DME) including topical application(s), wound assessment, and instruction(s) for ongoing care, per session; total wound(s) surface area greater than 50 square centimeters
A6550	Wound care set, for negative pressure wound therapy electrical pump, includes all supplies and accessories
E2402	Negative pressure wound therapy electrical pump, stationary or portable

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3 Hyperbaric Oxygen (HBO)

ASH considers Hyperbaric oxygen therapy medically necessary for the treatment of diabetic wounds of the lower extremities in patients who meet **ALL of** the following criteria:

1. Patient has type I or type II diabetes and has a lower extremity wound that is due to diabetes;

2. Patient has a wound classified as Wagner grade III or higher; and

3. Patient has failed an adequate course of standard wound therapy.

11 12 The use of HBO therapy is covered as adjunctive therapy only after there are no measurable signs of healing for at least 30 –days of treatment with standard wound therapy and must 13 be used in addition to standard wound care. Standard wound care in patients with diabetic 14 15 wounds includes assessment of a patient's vascular status and correction of any vascular problems in the affected limb, if possible, optimization of nutritional status, optimization 16 of glucose control, debridement by any means to remove devitalized tissue, maintenance 17 of a clean, moist bed of granulation tissue with appropriate moist dressings, appropriate 18 off-loading, and necessary treatment to resolve any infection that might be present. Failure 19 to respond to standard wound care occurs when there are no measurable signs of healing 20 21 for at least 30 consecutive days. Wounds must be evaluated at least every 30 days during administration of HBO therapy. Continued treatment with HBO therapy is not covered if 22 measurable signs of healing have not been demonstrated within any 30-day period of 23

24 treatment.

- · · · · ·	c Oxygen Therapy (HBOT):
99183	d if selection criteria are met: Physician or other qualified health care professional attendance and supervision of hyperbaric oxygen therapy, per session
HCPCS codes cover	ed if selection criteria are met:
G0277	Hyperbaric oxygen under pressure, full body chamber, per 30 minute interval
ICD-10 codes cover	ed if selection criteria are met
E08.51 - E08.59, E09.51 - E09.59	Diabetes mellitus due to underlying condition with peripheral circulatory disorders
E08.618 - E08.69, E09.618 - E09.69	Diabetes mellitus due to underlying conditions with other specified manifestations
E11.51 - E11.59, E13.51 - E13.59	Diabetes with peripheral circulatory disorders
E11.618 - E11.69, E13.618 - E13.69	Diabetes with other specified manifestations
I83.201 - I83.229	Varicose veins of lower extremities with ulcer and inflammation

2 Skin Substitutes and Soft Tissue Grafts

ASH considers the Skin Substitutes and Soft Tissue Grafts for wound care medically necessary according to the criteria **indicated below**:

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6 Application of a skin substitute graft/Cellular and Tissue-based Products (CTP) in the 7 treatment of diabetic foot ulcers (DFUs) and venous leg ulcers (VLUs) is considered 8 reasonable and necessary if the patient meets all criteria as outlined here:

- 9
 1. The presence of a chronic, non-infected DFU having failed to achieve at least 50%
 ulcer area reduction with documented standard of care (SOC) treatment* for a
 minimum of 4 weeks with documented compliance.
- The presence of a chronic, non-infected VLU having failed to respond to
 documented SOC treatment* for a minimum of 4 weeks with documented
 compliance.
- An implemented treatment plan to be continued throughout the course of treatmentdemonstrating all the following:
 - Debridement as appropriate to a clean granular base.
 - Documented evidence of offloading for DFUs.
 - Documented evidence of sustained compression dressings for VLUs.
 - Infection control with removal of foreign body or focus of infection.
 - Management of exudate with maintenance of a moist environment.

Documentation of smoking history, and counselling on the effect of 1 smoking on wound healing. Treatment for smoking cessation and outcome 2 of counselling (if applicable). 3 4. The skin substitute graft/CTP is applied to an ulcer that has failed to heal or has 4 stalled in response to documented SOC treatment*. Documentation of response to 5 treatment requires measurements of the initial ulcer, pre-SOC ulcer measurements, 6 weekly SOC ulcer measurements, post-completion SOC ulcer measurements 7 following (at least) 4 weeks of SOC treatment, ulcer measurements at initial 8 placement of the skin substitute graft/CTP, and before each subsequent placement 9 of the skin substitute graft/CTP. Failure to heal or stalled response despite standard 10 of care measures must have preceded the application for a minimum of 4 weeks 11 and established SOC treatment must continue for the course of therapy. Continuous 12 compression therapy for VLUs must be documented for the episode of care. 13 5. The medical record documentation must include the interventions having failed 14 during prior ulcer evaluation and management. The record must include an updated 15 medication history, review of pertinent medical problems diagnosed since the 16 previous ulcer evaluation, and explanation of the planned skin replacement with 17 choice of skin substitute graft/CTP. The procedure risks and complications must 18 also be reviewed and documented. 19 20 6. The patient is under the care of a qualified provider for the treatment of the systemic disease process(es) etiologic for the condition (e.g., venous insufficiency, diabetes, 21 neuropathy) and documented in the medical record. 22 23 24 *SOC treatment includes: 25 Comprehensive patient assessment (e.g., history, exam, vascular assessment) and diagnostic tests as indicated as part of the implemented treatment plan. 26 For patients with a DFU: assessment of type 1 or type 2 diabetes and management 27 • history with attention to certain co-morbidities (e.g., vascular disease, neuropathy, 28 osteomyelitis), review of current blood glucose levels/hemoglobin A1c (HbA1c), 29 diet and nutritional status, activity level, physical exam that includes assessment of 30 skin, ulcer, and vascular perfusion, and assessment of off-loading devices or use of 31 appropriate footwear. 32 For patients with a VLU: assessment of clinical history (that includes prior ulcers, 33 • body mass index, history of pulmonary embolism or superficial/deep venous 34 thrombosis, number of pregnancies, and physical inactivity), physical exam (e.g., 35 edema, skin changes and vascular competence), evaluation of venous reflux, 36

edema, skin changes and vascular competence), evaluation of venous reflux,
 perforator incompetence, and venous thrombosis. The use of a firm strength
 compression garment (>20 mmHg) or multi-layered compressive dressing is an
 essential component of SOC for venous stasis ulcers.

1	Cover	age requirements for skin substitute grafts/CTPs
2	To qua	alify as a skin substitute graft/CTP the product MUST be:
3	1.	A non-autologous human cellular or tissue product (e.g., dermal or epidermal,
4		cellular and acellular, homograft or allograft), OR non-human cellular and tissue
5		product (i.e., xenograft), OR biological product (synthetic or xenogeneic) applied
6		as a sheet, allowing scaffold for skin growth, intended to remain on the recipient
7		and grow in place or allow recipient's cells to grow into the implanted graft
8		material AND
9	2.	Supported by high-certainty evidence to demonstrate the product's safety,
10		effectiveness, and positive clinical outcomes in the function as a graft for DFUs
11		and/or VLUs. Substantial equivalence to predicate products does not allow
12		sufficient evidence to support similar cleared products.
13		
14	Note:	Liquid or gel preparations are not considered grafts. Their fluidity does not allow
15		lacement and stabilization of the product on the wound.
16		
17	The fo	llowing are considered reasonable and necessary (per episode of care):
18	1.	The maximum number of applications of a skin substitute graft/CTP within the
19		episode of skin replacement therapy (defined as 12 to 16 weeks from the first
20		application of a skin substitute graft/CTP) is 8 applications. The mean number of
21		skin substitute graft/CTP applications associated with wound healing is 4; however,
22		with documentation of progression of wound closure under the current treatment
23		plan and medical necessity for additional applications, up to 8 applications may be
24		allowed. Use of greater than 4 applications requires an attestation from the provider
25		showing that requirements specified here have been met and the additional
26		applications are medically necessary. In absence of this attestation, denial of the
27		additional applications will occur.
28	2.	The usual episode of care for skin substitute grafts/CTP is 12 weeks; however, some
29		wounds may take longer to heal therefore 16 weeks is allotted with documentation
30		that includes progression of wound closure under current treatment plan.
31	3.	The skin substitute graft/CTP must be used in an efficient manner utilizing the most
32		appropriate size product available at the time of treatment.
33		• Excessive wastage (discarded amount) should be avoided by utilization of
34		size appropriate packaging of the product consistent with the wound size.
35		The graft must be applied in a single layer without overlay of product or
36		adjacent skin in compliance with the correct label application techniques for
37		the skin substitute graft/CTP.
38	4.	Only skin substitute grafts/CTP with labeled indications for use over exposed
39		muscle, tendon, or bone will be considered reasonable and necessary for those
40		indications.

Limitations 1 The following are considered **NOT** reasonable and necessary: 2 1. Greater than 8 applications of a skin substitute graft/CTP within an episode of care 3 (up to 16 weeks). 4 2. Repeat applications of skin substitute grafts/CTP when a previous application was 5 unsuccessful. Unsuccessful treatment is defined as an increase in size or depth of 6 an ulcer, no measurable change from baseline, and no sign of improvement or 7 indication that improvement is likely (such as granulation, epithelialization, or 8 progress towards closure). 9 3. Application of skin substitute grafts/CTP in patients with inadequate control of 10 underlying conditions or exacerbating factors, or other contraindications (e.g., 11 active infection, progressive necrosis, active Charcot arthropathy of the ulcer 12 extremity, active vasculitis, or ischemia). 13 4. Use of surgical preparation services (e.g., debridement), with routine, simple, or 14 repeat skin replacement surgery with a skin substitute graft/CTP. 15 5. All liquid or gel skin substitute products/CTP for ulcer care. 16 6. Placement of skin substitute grafts/CTP on an infected, ischemic, or necrotic wound 17 bed. 18 19 20 For more information on applicable codes for specific skin substitute products/CTP please refer to Local Coverage Determination (LCD): Skin Substitute Grafts/Cellular and Tissue-21 Based Products for the Treatment of Diabetic Foot Ulcers and Venous Leg Ulcers 22 (L35041). 23

24

HCPCS codes covered	if selection criteria are met	
Q4101	Apligraf, per sq cm	
ICD-10 codes covered	if selection criteria are met	
E08.621	Diabetes mellitus due to underlying condition with foot ulcer	
E09.621	Drug or chemical induced diabetes mellitus with foot ulcer	
E10.621	Type 1 diabetes mellitus with foot ulcer	
E11.621	Type 2 diabetes mellitus with foot ulcer	
E13.621	Other specified diabetes mellitus with foot ulcer	
I83.001 - I83.029	Varicose veins of lower extremities with ulcer	
183.201 - 183.229	Varicose veins of lower extremities with ulcer and inflammation	
I87.311 - I87.319	Chronic venous hypertension (idiopathic) with ulcer	
187.331 - 187.339	Chronic venous hypertension (idiopathic) with ulcer and inflammation	

Dermagraft:			
HCPCS codes covered if	selection criteria are met		
Q4106	Dermagraft, per sq cm		
ICD-10 codes covered if s	selection criteria are met		
E08.621	Diabetes mellitus due to underlying condition with foot ulcer		
E09.621	Drug or chemical induced diabetes mellitus with foot ulcer		
E10.621	Type 1 diabetes mellitus with foot ulcer		
E11.621	Type 2 diabetes mellitus with foot ulcer		
E13.621	Other specified diabetes mellitus with foot ulcer		
Q81.2	Epidermolysis bullosa dystrophica		
Transcyte:			
No specific code			
ICD-10 codes covered if s	selection criteria are met		
T20.011A - T25.799S	Burns		
Orcel:			
No specific code			
HCPCS codes covered if	selection criteria are met		
Q4100	Skin substitute, not otherwise specified		
ICD-10 codes covered if s	selection criteria are met		
Q81.2	Epidermolysis bullosa dystrophica		
T20.011A - T25.799S	Burns		
Biobrane biosynthetic dr	essing:		
No specific code			
CPT® codes covered if se	election criteria are met		
15050, 15100 - 15261	Autograft/tissue cultured autograft		
ICD-10 codes covered if s	selection criteria are met		
T20.011A - T25.799S	Burns		

Integra Dermal Regenerat and Integra Meshed Bilay	tion Template, Integra Bilayer Matrix Wound Dressing, er Wound Matrix:		
HCPCS codes covered if s	election criteria are met		
C9363	Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter		
Q4104	Integra Bilayer Matrix Wound Dressing (BMWD), per sq sm		
Q4105	Integra Dermal Regeneration Template (DRT), or Integra Omnigraft Dermal Regeneration Matrix, per sq cm		
ICD-10 codes covered if se	election criteria are met		
T20.011A - T25.799S	Burns		
Artiss:			
HCPCS codes covered if s	election criteria are met		
C9250	Human plasma fibrin sealant, vapor-heated, solvent- detergent (Artiss), 2ml		
ICD-10 codes covered if se	election criteria are met		
T20.011A - T25.799S	Burns		
Oasis Wound Matrix:			
HCPCS codes covered if s	election criteria are met		
Q4102	Oasis Wound Matrix, per sq cm		
ICD-10 codes covered if se	election criteria are met		
E08.621	Diabetes mellitus due to underlying condition with foot ulcer		
E09.621	Drug or chemical induced diabetes mellitus with foot ulcer		
E10.621	Type I diabetes mellitus with foot ulcer		
E11.621	Type II diabetes mellitus with foot ulcer		
E13.621	Other specified diabetes mellitus with foot ulcer		
183.001 - 183.028	Varicose veins of lower extremities with ulcer		
I83.201 - I83.229	Varicose veins of lower extremities with ulcer and inflammation		
I87.311 - I83.319	Chronic venous hypertension with ulcer		
187.331 - 187.339	Chronic venous hypertension with ulcer and inflammation		
Graftjacket Regenerative	Tissue Matrix:		
HCPCS codes covered if s	election criteria are met		
Q4107	Graftjacket, per sq cm		

E08.621, E09.621, E10.621, E11.621, E13.621	08.621, E09.621, E10.621, Diabetes mellitus			
Epicel :				
No specific code				
CPT® codes covered if sele 15150 - 15157	Tissue cultured skin autograft, face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits			
ICD-10 codes covered if sel	ection criteria are met			
*T20.30XA - *T20.39XS, T20.711A - *T20.79XS	Burn and corrosion of third degree of face, head, and neck			
*T21.30XA - *T21.39XS, *T21.70XS - *T21.79XS	Burn and corrosion of third degree of trunk			
*T22.30XA - T22.399S, *T22.70XA - T22.799S	Burn and corrosion of third degree of shoulder and upper limb			
T23.301A - T23.399S, T23.701A - T23.799S	Burn and corrosion of third degree of wrist and hand			
T24.301A - T24.399S, T24.701A - T24.799S	Burn and corrosion of third degree of lower limb, except ankle and foot			
T25.311A - T25.399S, T25.711A - T25.7799S	Burn and corrosion of third degree of ankle and foot			
**T31.30 - T31.99, T32.30 - T32.99	Burn and corrosion 30 to 90 percent or more of body surface			
CPT® codes covered if sele	ection criteria are met			
***15271 - 15278	Application of skin substitute graft			
*Use additional external ca (X00-X19, X75-X77, X96-X	use code to identify the source, place, and intent of the bur (02, NO2)			

**Burn and corrosion codes inclusive of third degree burns only, as described within the scope of these codes.

*** Graft application codes must be associated with one of the grafts listed above.

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Surgical Preparation and Skin Replacement (CPT® codes 15002 – 15005) 1 1. Per the definitions and the guidelines in CPT® Code Book codes CPT® codes 2 15002/15005 are not appropriate codes to use when performing a non-surgical 3 application of a skin substitute. 4 2. CPT® codes 15002/15005 are only appropriately used in place of service inpatient 5 hospital, outpatient hospital or ambulatory surgical center with regional or general 6 anesthesia to resurface an area damaged by burns, traumatic injury, or surgery. An 7 operative report is required and must be available upon request. 8 9 CPT® codes 15002-15005 are to be used for the initial traumatic wound preparation 10 (removal of appreciable nonviable tissue) and cleaning to provide a viable wound surface 11 (primary intention healing) for placement of an autograft, flap, skin substitute graft or for 12 negative pressure wound therapy. Primary intention presumes that the performance of the 13 skin preparation and the application of the autograft, flap, skin substitute graft or for 14 negative pressure wound therapy is to heal the wound. 15 16 CPT® codes 15002-15005 are NOT to be used for the removal of nonviable tissue/debris 17 in chronic wounds left to heal by secondary intention. CPT® 11042-11047 and CPT® 18 97597-97598 are to be used for this. 19 20 CPT® codes 15002-15005 are selected based on the anatomic area and size of the 21 prepared/debrided defect. For multiple wounds, the choice of code is based on the 22 aggregate sum of the surface area of all similarly grouped wound types. 23 24 Codes 15002 - 15005 should NOT be reported for the removal of nonviable tissue/debris 25 in a chronic wound (e.g., venous, or diabetic) when the wound is left to heal by secondary 26 intention. Regarding CPT® codes 15002-15005: 27 Use when preparing a proper wound surface for the placement of a graft, flap, skin 28 replacement, skin substitute, or negative pressure therapy. 29 • Appreciable nonviable tissue is always removed. 30 A clean wound bed may be created by incisional release of a scar contracture, 31 resulting in a surface defect from separation of tissue. 32 The purpose of these codes is to prepare the wound to heal by primary intention or 33 • negative pressure wound therapy. 34 The patient's condition may require that final closure may be delayed. 35 • 36 Use CPT[®] codes 15271 - 15278 for the surgical preparation or creation of recipient site 37 for the tissue skin graft. Regarding CPT® codes 15271-15278: 38 Wound prep codes are separate from skin substitute graft application codes. 39 • The ankle is considered 'leg' in terms of skin substitute graft application. 40 • • Wound areas that skin substitute grafts will be applied are measured 41 42 AFTER prep/debridement.

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• Bill either the 'small' leg/ankle skin substitute graft codes or the 'large' 1 2 skin substitute graft codes (see description below). 3 • Bill either the 'small' foot/toe skin substitute graft codes or the 'large' skin substitute graft codes (see description below). 4 • It is acceptable to bill both the leg/ankle and the foot/toe skin substitute graft 5 application codes if you are treating both the leg/ankle and the foot/toe. 6 • Do not discount an 'add-on code'; do not apply a '-51' modifier. 7 8 'Small Wounds' - for wounds known to have an aggregate wound size up to a maximum 9 of 100 cm². The codes represent the first 25 cm² or 1% of body area in infants and children, 10 and additional 25 cm^2 or 1% of body area in infants and children, up to that maximum 100 11 cm^2 wound area. 12 13 14 'Large Wounds' - for wounds known to have an aggregate wound size beginning at 100 cm² or greater. The 'small wound' codes would not be used in these cases; instead, 15 surgeons would use the 'large wound' codes which begin with a wound area of 100 cm² or 16 greater. "Large wound" codes refer to: 1) the initial 100 cm² or 1% of body area in infants 17 and children, and 2) each additional 100 cm² or 1% of body area in infants and children. 18 19 20 **CPT® Codes and Descriptions CPT®** Code **CPT® Code Description** Surgical preparation or creation of recipient site by excision of open wounds, burn eschar, or scar (including subcutaneous tissues), or 15002 incisional release of scar contracture, trunk, arms, legs; first 100 sq cm or 1% of body area of infants and children Surgical preparation or creation of recipient site by excision of open wounds, burn eschar, or scar (including subcutaneous tissues), or incisional release of scar contracture, trunk, arms, legs; each 15003 additional 100 sq cm, or part thereof, or each additional 1% of body area of infants and children (List separately in addition to code for primary procedure) Surgical preparation or creation of recipient site by excision of open wounds, burn eschar, or scar (including subcutaneous tissues), or incisional release of scar contracture, face, scalp, eyelids, mouth, 15004 neck, ears, orbits, genitalia, hands, feet and/or multiple digits; first 100 sq cm or 1% of body area of infants and children Surgical preparation or creation of recipient site by excision of open 15005 wounds, burn eschar, or scar (including subcutaneous tissues), or

incisional release of scar contracture, face, scalp, evelids, mouth,

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CPT® Code	CPT® Code Description	
	neck, ears, orbits, genitalia, hands, feet and/or multiple digits; each additional 100 sq cm, or part thereof, or each additional 1% of body area of infants and children (List separately in addition to code for primary procedure)	
15271	Application of skin substitute graft to trunk, arms, legs, total wound surface area up to 100 sq cm; first 25 sq cm or less of wound surface area	
15272	Application of skin substitute graft to trunk, arms, legs, total wound surface area up to 100 sq cm; each additional 25 sq cm wound surface area, or part thereof (List separately in addition to code for primary procedure)	
15273	Application of skin substitute graft to trunk, arms, legs, total wound surface greater than or equal to 100 sq cm; first 100 sq cm wound surface area, or 1% of body area of infants and children	
15274	Application of skin substitute graft to trunk, arms, legs, total wound surface greater than or equal to 100 sq cm; each additional 100 sq cm wound surface area, or part thereof, or each additional 1% of body area of infants and children, or part thereof (List separately in addition to code for primary procedure)	
15275	Application of skin substitute graft to face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits, total wound surface area up to 100 sq cm; first 25 sq cm or less wound surface area	
15276	Application of skin substitute graft to face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits, total wound surface area up to 100 sq cm; each additional 25 sq cm wound surface area, or part thereof (List separately in addition to code for primary procedure)	
15277	Application of skin substitute graft to face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits, total wound surface area greater than or equal to 100 sq cm; first 100 sq cm wound surface area, or 1% of body area of infants and children	

CPT® Code	CPT® Code Description
15278	Application of skin substitute graft to face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits, total wound surface area greater than or equal to 100 sq cm; each additional 100 sq cm wound surface area, or part thereof, or each additional 1% of body area of infants and children, or part thereof (List separately in addition to code for primary procedure)

For preparation of wounds on the trunk, arms, and/or legs, report 15002 for the first 100 sq cm of site prep. For additional preparation (beyond 100 sq cm) in the same anatomic areas, report add-on 15003. Because 15003 is an add-on code, report it only in addition to 15002. Likewise, for preparation of wounds of the face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits, report 15004 for the first 100 sq cm of site prep. For additional preparation (beyond 100 sq cm) in the same anatomic areas, report add-on 15005—again, only in addition to 15004.

8

Surgical preparation may be reported only once per wound. If the wound is prepared, but
not grafted (e.g., grafting won't occur until the next day), minimal preparation of the wound
bed is included in the graft code, as is removing a previous graft.

12

Codes 15002-15005 apply specifically to describe the work of preparing a clean and viable wound surface for placement of an autograft, flap, skin substitute graft or for negative pressure wound therapy, according to CPT® guidelines. Surgical prep codes would not be reported for removal of nonviable tissue or debris in a chronic wound when it is left to heal by secondary intention. When a wound requires serial debridement, report active wound management (97597-97598) or debridement (11042-11047). If a wound requires negative pressure wound therapy, 15002-15005 are applicable in addition to 97605-97606.

20

21 **DESCRIPTION/BACKGROUND**

A wound by true definition is any disruption of the integrity of skin, mucous membrane, 22 or organ tissue (Kujath & Michelsen, 2008). Wounds can be caused by mechanical, 23 thermal, chemical, and radiogenic trauma. To be distinguished from these are those wounds 24 that have their origin due to underlying pathologies, such as diabetes mellitus, chronic 25 venous/arterial insufficiency, and immunological or dermatological diseases (Kujath & 26 Michelsen, 2008). A wound may be classified in many ways; by its etiology, anatomical 27 location, by whether it is acute or chronic, by method of closure, by its presenting 28 symptoms or by the appearance of the predominant tissue types in the wound bed (Enoch 29 et al., 2004). Some of the most common causes of chronic wounds are tissue loads over 30 bony prominences and lower extremity wounds secondary to neuropathy and venous 31 hypertension (Irion, 2010). Occasionally wounds are due to ischemia. It is critical that the 32 clinician be able to perform a good differential diagnosis between the types of wounds 33

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1 (arterial, venous hypertension, neuropathic, and/or from lymphatic disease) because the 2 management of each wound differs and may be contraindicated in the presence of ischemia.

3

4 Wound Types

The two major types of wounds are acute or chronic wounds. Acute wounds will heal in 5 orderly and timely reparative processes that result in sustained restoration of anatomic and 6 functional integrity, usually in 30 days or less (Lazarus et al., 1994). Chronic wounds, on 7 the other hand, are wounds that fail to complete the reparative process of healing in the 8 expected period, usually greater than 30 days, or proceeded through the healing phase 9 without establishing the expected functional result due to an interruption in the biological 10 11 or physiologic process of normal healing (ECRI, 2010). Chronic wounds generally do not achieve wound closure without some type of intervention. The common chronic cutaneous 12 wounds include venous stasis ulcers, arterial insufficiency ulcers, neuropathic ulcers, and 13 14 pressure ulcers (Bello and Phillips, 2000).

15

Venous stasis ulcers occur when there is an improper functioning of the venous valves, 16 usually in the lower extremities, causing a back flow and increased pressure in veins (Bello 17 and Phillips, 2000; Palfreyman et al., 2007). The body needs the pressure gradient between 18 arteries and veins in order for the heart to pump blood forward through the arteries and 19 20 veins. When there is an interruption in this pressure gradient and the arteries have a significantly lower pressure than the veins, which is known as venous hypertension, the 21 blood is not pumped as effectively and causes it to pool in the lower extremities (Brem et 22 al., 2004; Stanley et al, 2005). The standard of care for venous stasis ulcers is compression 23 therapy at 30 to 40 mm Hg (Bello and Phillips, 2000; Palfreyman et al., 2007). Treatment 24 regimens focus on increasing venous return and decreasing edema (Burns et al., 2007; 25 Palfreyman et al., 2007). 26

27

Arterial ulcers are caused by an insufficient arterial blood supply. Arterial ulcers occur 28 because there is inadequate perfusion of skin and subcutaneous tissue, resulting in tissue 29 ischemia and necrosis, usually due to a complete or partial blockage of the arteries (Bello 30 and Phillips, 2000; Holloway, 1996). Arterial insufficiency occurs as a result of peripheral 31 arterial disease (PAD) and causes decreased perfusion to the tissues distal to an arterial 32 33 plaque formation. Reestablishment of an adequate vascular supply is a key factor to support proper healing. Comprehensive medical management would include wound care to the 34 ulcer itself and management to include control of the common causes of arterial ulcers 35 (diabetes mellitus, control of hypertension, smoking cessation, proper nutrition, and 36 moderate exercise) (Bello and Phillips 2000; Guo and DiPietro, 2010). 37

38

Neuropathic ulcers form as a result of peripheral neuropathy, typically seen with diabetic patients but can be due to other metabolic disease process (renal failure), trauma, or surgery. Peripheral neuropathy affects the sensory nerves responsible for detecting sensations such as temperature or pain (American Diabetes Association (AMA), 1999).

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This loss of sensation causes local paresthesias, usually in the feet and/or lower extremities, 1 which can lead to microtrauma, breakdown of the overlying tissues, and eventually 2 ulceration, often seen over pressure points on the foot. Peripheral neuropathy can also 3 damage motor nerves causing minor muscle wasting resulting in muscle imbalances that 4 can cause foot deformities, which can lead to more prominent bony areas giving rise to 5 additional pressure points prone to ulceration (AMA, 1999; Krestel Editors, 2010; Lazarus 6 et al., 1994). In addition to basic wound care management, other medical management 7 includes maintaining optimal blood sugar levels, pressure relief at the wound site, surgical 8 debridement, control of infection, and arterial reconstruction. 9

10

A pressure ulcer is an injury to the skin and/or underlying tissue over a bony prominence 11 that occurs as a result of pressure in conjunction with or without shear or friction. Pressure 12 ulcers can also result from poorly fitting casts or appliances. They can occur in soft tissue 13 areas due to the pressure effects of a foreign object such as a medical device. Because 14 muscle and subcutaneous tissue are more susceptible to pressure induced injury than 15 dermis and epidermis, pressure ulcers are often worse than their initial presentation. 16 Pressure ulcers are assessed and staged at the bedside as a clinical description of the depth 17 of observable tissue destruction. 18

19

For the purpose of this clinical practice guideline, the staging of pressure ulcers can be classified according to the National Pressure Ulcer Advisory Panel as follows (Black et al., 2007):

23

Pressure Ulcer Stage	Description	
(Suspected) Deep Tissue Injury	• I from pressure and/or snear The area may be preceded by	
Stage I	Intact skin with non-blanchable redness of a localized area usually over a bony prominence. Darkly pigmented skin may not have visible blanching; its color may differ from the surrounding area.	
Stage II	Partial-thickness loss of dermis presenting as a shallow open ulcer with a red-pink wound bed, without slough. May also present as an intact or open/ruptured serum-filled blister.	
Stage III	Full-thickness tissue loss. Subcutaneous fat may be visible, but bone, tendon, or muscle are not exposed. Slough may be present but does not obscure the depth of tissue loss. May include undermining and tunneling.	

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Pressure Ulcer Stage	Description		
Stage IVFull-thickness tissue loss with exposed bone, tendon, or muscle. Slough or eschar may be present on some part the wound bed. Often includes undermining and tunner			
Unstageable Full-thickness tissue loss in which the base of the ulcer i covered by slough (yellow, tan, gray, green, or brown) and/or eschar (tan, brown, or black) in the wound bed.			

The National Pressure Ulcer Advisory Panel (2009) recommends debridement of devitalized tissue within the wound bed or edge of pressure ulcers when appropriate to the individual's condition and consistent with the overall goals of care.

5

6 Osteomyelitis

Osteomyelitis is inflammation of the bone caused by an infecting organism. Although bone is normally resistant to bacterial colonization, events such as trauma, surgery, presence of foreign bodies, or prostheses may disrupt bony integrity and lead to the onset of bone infection. Osteomyelitis can also result from hematogenous spread after bacteremia. When prosthetic joints are associated with infection, microorganisms typically grow in biofilm, which protects bacteria from antimicrobial treatment and the host immune response.

13

Acute osteomyelitis presents with acute inflammatory cells, edema, vascular congestion, and small-vessel thrombosis. In early disease, infection extends into the surrounding soft tissue, which compromises the vascular supply to the bone, as well as host response, surgery, and/or antibiotic therapy. Chronic osteomyelitis presents with pathologic findings of necrotic bone, formation of new bone, and polymorphonuclear leukocyte exudation, which is joined by large numbers of lymphocytes, histiocytes, and occasional plasma cells.

21 Surgery is indicated to treat osteomyelitis when the patient has not responded to specific antimicrobial treatment, if there is evidence of a persistent soft tissue abscess or 22 subperiosteal collection, or if concomitant joint infection is suspected. Debridement of 23 necrotic tissues, removal of foreign materials, and sometimes skin closure of chronic 24 unhealed wounds is necessary in some cases (Kishner et al., 2014). The Infectious Disease 25 Society of America (IDSA) guideline for the treatment of diabetic foot infections (Lipsky 26 27 et al., 2012) recommends surgical intervention ranging from minor (debridement) to major (resection, amputation) for diabetic foot infections such as osteomyelitis. 28

29

30 Wound Healing

Wound healing is traditionally divided into the following four phases: (1) exudative phase, (2) resorptive phase, (3) proliferative phase and (4) regenerative phase. Each of the traditional phases listed describe their biophysiological functions that occur during that phase that leads to the next phase (Kujath & Michelsen, 2008). In recent English language

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publications, wound healing is divided into the following four phases: hemostasis, inflammation, proliferation, and tissue remodeling or resolution (Guo and DiPietro, 2010; Kujath & Michelsen, 2008; Singer, 1999). There are many different medically accepted terms used for wound care that describe the phases of wound healing. For the purpose of this paper, wound healing will be referred to as a normal biological process in the human body that is achieved through four highly integrated and overlapping phases: hemostasis, inflammation, proliferation, and remodeling (Guo and DiPietro, 2010).

8

The primary goals of wound management are rapid wound closure and a functional, 9 mechanically stable and aesthetically acceptable scar (Kujath and Michelsen, 2008). 10 Wounds can heal either by primary intention or secondary intention depending upon 11 whether the wound may be closed with sutures or left to repair on its own, whereby 12 damaged tissue is restored by the formation of connective tissue and re-growth of 13 epithelium (Cooper, 2005). Cooper's definition of primary intention is when the edges of 14 the wound are approximated, and the individual layers of tissue are joined together either 15 by sutures, staples or tissue adhesives or a combination of all of these. Secondary intention 16 is when the wound sustains a degree of tissue loss where it appears that the wound closure 17 is impossible secondary to either the presence of infection and wound closure is undesirable 18 or wound edges are so far apart (Cooper, 2005). Primary wound healing is the 19 20 uncomplicated healing process that involves the non-infected, well-adapted wounds (Kujath & Michelsen, 2008). If the healing process is disturbed by local factors such as 21 infections, dehiscence, inadequate blood perfusion or systemic factors such as 22 immunocompromise, a situation of secondary wound healing develops (Cooper, 2005; 23 Kujath & Michelsen, 2008; Guo and DiPietro, 2010). 24

25

For the normal healing process to occur, the four phases of healing and their 26 biophysiological functions must occur in the proper sequence, at a specific time and 27 continue for a specific duration at an optimal intensity (Mathieu et al., 2006). There are 28 many factors that can affect wound healing which may interfere with one or more of the 29 healing phases, thus causing improper or impaired tissue repair and delays in wound 30 closure. Wounds that exhibit impaired healing, which can include delayed acute wounds 31 and/or chronic wounds, have failed to progress through the normal stages of healing. 32 33 Chronic wounds are examples of wounds that have a biological or physiological reason for not healing. It is the chronic wounds that frequently enter a state of pathological 34 inflammation due to postponed, incomplete, or uncoordinated healing process (Guo and 35 36 DiPietro, 2010).

37

38 Choice of Dressing

- A wound will require different management and treatment at various stages of healing. No dressing is suitable for all wounds; therefore, frequent assessment of the wound is required.
- 41 Considerations when choosing dressing products:
- 42

- Be able to control (remove) excess exudates. A moist wound environment is good, a wet environment is not beneficial
- Not stick to the wound, shed fibers or cause trauma to the wound or surrounding tissue on removal
 - Protect the wound from the outside environment bacterial barrier
 - Good adhesion to skin
- 7 Sterile
- Aid debridement if there is necrotic or sloughy tissue in the wound (caution with ischemic lesions)
- Keep the wound close to normal body temperature
- Conformable to body parts and doesn't interfere with body function
 - Be cost-effective
 - Diabetes choose dressings which allow frequent inspection
 - Non-flammable and non-toxic
- 14 15

13

1 2

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4

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6

Dry wound	Minimal exudate	Moderate exudate	Heavy exudate
Non adherent island dressing	Hydrogel	Calcium alginate	Hydrofibre
Hydrocolloid	Hydrocolloid	Hydrofibre	Foam
Films semi permeable	Silicone absorbent	Foams	Absorbent dressing
		Negative Pressure	Negative pressure wound therapy
		Hydrocolloid: paste/powder	Ostomy

16

17 EVIDENCE REVIEW

While there are numerous treatments that have been proposed as interventions to treat 18 19 chronic wounds, not all have been well-studied and there is not enough evidence to prove their safety and effectiveness. Some of the researched treatments that have some evidence 20 (but may not be confirmatory) to support their safety and effectiveness include ultrasound, 21 low level laser, electromagnetic (EM) therapy/diathermy, electrical stimulation (ES), 22 23 hyperbaric oxygen, surgical debridement, surgical revascularization of the affected area, myocutaneous skin flaps or grafting, use of various dressings (e.g., wet to dry, multilayer 24 compression bandages), negative pressure wound therapy (vacuum-assisted closure), and 25 the use of certain bioengineered skin substitutes. This paper will focus on those 26 interventions within the scope of practice of the wound care specialist. 27

28

Brolmann et al. (2012) completed a meta-analysis on the evidence for local and systemic
 wound care. Forty-four relevant reviews were included in this summary paper. Wounds

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included venous ulcers, acute wounds, pressure ulcers, diabetic ulcers, arterial ulcers, and
 miscellaneous chronic wounds. The authors summarized that strong evidence supports the
 effectiveness of therapeutic ultrasound, mattresses, cleansing methods, closure of surgical
 wounds, honey, antibiotic prophylaxis, compression, lidocaine-prilocaine cream, skin
 grafting, antiseptics, debridement, and hyperbaric oxygen therapy.

6

7 Electrical Stimulation (ES)

Electrical stimulation (ES) is one of several treatment modalities that have been studied for 8 the use of healing chronic wounds. Several randomized controlled trials have evaluated ES 9 with varying protocols using different currents and voltages for the healing of pressure 10 11 ulcers, venous stasis ulcers, arterial insufficiency ulcers, surgical wounds, and diabetic wounds (Houghton, 2003; Feedar et al. 1991; Fernandez et al. 2004). It is known that living 12 tissues possess electrical potentials that may play a role in the healing process. In early 13 studies by Wolcott et al. (1969), researchers showed that ischemic ulcers healed 14 significantly faster with the use of electrical stimulation. Researchers have studied the use 15 of ES with regards to the type of electrical current applied (low-intensity direct current, 16 low-intensity pulsed current, or high-voltage pulsed current) and the placement of 17 electrodes (in direct contact, close proximity, or to a skin wound), thereby creating an 18 electrical current that passes through the wound (Houghton, 2003; Feedar, 1991; 19 20 Fernandez, 2004; Ho, 2008; Recio et al., 2012).

21

Recio et al. (2012) studied the effectiveness of high-voltage electrical stimulation used to 22 manage stage III and IV pressure ulcers among adults with spinal cord injury (SCI). 23 Through retrospective studies the authors describe the care of adults with SCI with 24 recalcitrant pressure ulcers below the level of injury. Electrical stimulation was applied 25 directly into the wound bed: 60 minutes per session, 3-5 times per week; with an intensity 26 of 100 milliamperes and frequency of 100 pulses per second. Polarity was negative, 27 initially and was switched weekly. The amplitude and wave form were maintained 28 throughout each treatment session. The results showed that the long-standing (11-14 29 months) pressure ulcers were completely healed after 7 to 22 weeks of treatment with high-30 voltage ES. The study concluded that ES is effective for enhanced healing of Stage III-IV 31 ulcers otherwise unresponsive to standard wound care (Recio et al., 2012). 32

33

Houghton et al. (2003) studied the effect of high voltage pulsed current (HVPC) electrical 34 stimulation on healing chronic leg ulcers. The authors studied twenty-seven people with a 35 total of 42 chronic leg ulcers. The subjects were separated into subgroups according to 36 primary wound type (venous stasis, arterial insufficiency, diabetes) and then randomly 37 assigned to receive either HVPC (100 microseconds, 150V, 100Hz) or sham treatment for 38 39 45 minutes, 3 times weekly, for 4 weeks. Wound surface area and wound appearance were assessed during the initial evaluation, following 1- to 2- week period during which subjects 40 received only conventional wound therapy, after 4 weeks of sham or HVPC treatments, 41 and at 1 month post treatment. The results indicated that the use of HVPC to chronic leg 42

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ulcers reduced the wound surface area over the 4-week treatment period to approximately one half the initial wound sizes, which was over 2 times greater than that observed in wounds treated with the sham treatment. The authors concluded that HVPC administered 3 times a week is an effective treatment to accelerate wound closure of chronic lower extremity ulcers due to diabetes, or to arterial or venous insufficiency (Houghton et al., 2003).

8 Studies have not adequately evaluated the safety and effectiveness of unsupervised home 9 use of the electrical stimulation devices by a patient. Evaluation of the wound is an integral 10 part of wound management. It is recommended that when ES is used as an intervention to 11 treat chronic wounds, treatment should be conducted under the direct supervision of a 12 medical professional with the expertise in wound evaluation and management (CMS, 2004, 13 2003).

14

Barnes et al. (2014) conducted a review and meta-analysis of RCTs on electric stimulation vs. standard care for chronic ulcer healing. This systematic review also aimed to investigate the effect of different types of electrical stimulation on ulcer size reduction. Twenty-one studies were eligible for inclusion in the meta-analysis. Authors concluded that electrical stimulation appears to increase the rate of ulcer healing and may be superior to standard care for ulcer treatment.

21

Lala et al. (2015) conducted a systematic review and meta-analysis on the effects of 22 electrical stimulation therapy (EST) on healing pressure ulcers in individuals with spinal 23 cord injury (SCI). A meta-analysis with five studies demonstrated that EST significantly 24 decreased the ulcer size compared to standard wound care or sham EST. Another meta-25 analysis conducted with four studies showed that EST increased the risk of wound healing 26 by 1.55 times compared with standard wound care or sham EST. Because of the wide array 27 of outcome measures across studies, a single meta-analysis could not be conducted. 28 However, EST appears to be an effective adjunctive therapy to accelerate and increase 29 pressure ulcer closure in individuals with SCI. 30

31

Chen et al. (2020) evaluated the effectiveness of electric stimulation (ES) for diabetic foot 32 33 ulcer (DFU) treatment. Of the 145 randomized clinical trials initially identified, 7 studies (with a total of 274 patients) met the inclusion criteria. The percentage decrease in ulcer 34 area at 4 weeks was significantly greater in patients treated with ES and SWC than SWC 35 alone. The ulcer healing rate at 12 weeks was also significantly faster in the ES group. 36 Subgroup analysis showed comparable efficacies with different waveforms (monophasic 37 vs biphasic). Authors concluded that electrical stimulation appears to be an effective 38 39 adjunctive therapy for accelerating DFU healing.

- 40
- Avendaño-Coy et al. (2021) examined the effectiveness and safety of electrical
 microcurrent therapy (EMT) for improving wound healing and pain in people with acute

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or chronic wounds. Eight RCTs were included in the qualitative summary and seven in the 1 quantitative analysis (n = 337 participants). EMT plus standard wound care (SWC) 2 produced a greater decrease in wound surface and healing time that SWC alone, showing 3 moderate and low certainty in the evidence, respectively. However, no differences were 4 observed in the number of healed wounds, with very low quality of evidence. EMT 5 decreased perceived pain, but no differences in adverse effects were noted between groups. 6 Authors concluded that EMT is an effective, safe treatment for improving wound area, 7 healing time, and pain. Further clinical trials that include detailed intervention parameters 8 and protocols should be designed to lower the risk of bias. 9

10

11 Electromagnetic Therapy (ET)/Diathermy

Aziz et al. (2013) completed a Cochrane review on electromagnetic therapy for treating 12 venous leg ulcers to assess the effects of EMT on the healing of venous leg ulcers. Authors 13 concluded that there was no high-quality evidence that electromagnetic therapy increases 14 the rate of healing of venous leg ulcers, and further research is needed. Wang et al. (2024) 15 evaluated the effects of electromagnetic therapy (EMT) on the treatment of venous leg 16 ulcers (VLUs) by synthesizing and appraising available meta-analyses (MAs) and 17 systematic reviews (SRs). The search yielded five eligible studies. The reviews collectively 18 presented moderate methodological quality and a low risk of bias in several domains. 19 20 Reporting quality was high, albeit with inconsistencies in fulfilling certain PRISMA checklist items. The evidence quality, primarily downgraded due to small sample sizes, 21 was rated as moderate. While some studies suggest potential benefits of EMT in the 22 treatment of VLUs, the overall evidence is inconclusive due to methodological limitations 23 and limited sample sizes. This review underscores the need for future research with more 24 rigorous methodologies and larger cohorts to provide clearer insights into the efficacy of 25 EMT for VLUs. 26

27

28 Ultraviolet (UV) Light

Chen et al. (2014) sought to determine the effects of phototherapy on the healing of 29 pressure ulcers. Seven RCTs involving 403 participants were selected. All the trials were 30 at unclear risk of bias. Trials compared the use of phototherapy with standard care only (6 31 trials) or sham phototherapy (1 trial). Only one of the trials included a third arm in which 32 33 another type of phototherapy was applied. Overall, there was insufficient evidence to determine the relative effects of phototherapy for healing pressure ulcers. Variations in 34 studies did not allow for pooling of the studies to draw any conclusions as to whether 35 phototherapy is effective or not. Authors conclude that uncertainty exists as to the effects 36 of phototherapy in treating pressure ulcers. The quality of evidence is very low due to the 37 unclear risk of bias and small number of trials available for analysis. The possibility of 38 39 benefit or harm of this treatment cannot be ruled out. Further research is recommended.

- 40
- Inkaran et al. (2021) examined the effect of UV light on wound healing and infection in
 patients with skin ulcers or surgical incisions. Outcomes of interest included healing time,

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wound size and appearance, bacterial burden, and infection. Comparative and 1 noncomparative clinical studies were considered, including observational cohort, 2 retrospective, and randomized controlled studies. They addressed the research question: 3 "Does the use of UV light as an adjunct to conventional treatment help improve healing 4 and reduce infection in wounds?" The search yielded 30,986 articles, and screening 5 resulted in 11 studies that underwent final analysis. Of these (N = 27,833), seven (64%) 6 demonstrated an improvement in healing outcomes with adjunctive UV therapy, and the 7 results of four (36%) achieved statistical significance. Authors concluded there is limited 8 research on the utility of adjunctive UV therapy to improve wound healing outcomes in 9 humans. The majority of literature included in this review supported improved wound 10 healing outcomes with adjuvant UV therapy. Future well-designed randomized controlled 11 trials will be essential in further determining the benefit and utility of UV therapy in wound 12 healing. 13

14

15 Non-Contact Ultrasound

Olyaie et al. (2013) conducted a RCT to compare the effectiveness of standard treatment 16 and standard treatment plus either high-frequency ultrasound (HFU) or noncontact low-17 frequency ultrasound (NCLFU) on wound outcomes. Outcomes of both methods of 18 ultrasound therapy were better than standard care alone, and some differences between the 19 20 two ultrasound therapy groups were observed, but they were not statistically significant. Beheshti et al. (2014) compared high-frequency and MIST ultrasound therapy for the 21 healing of venous leg ulcers. All groups received the standard wound care. In the 22 ultrasound groups, HFU and MIST ultrasound therapy was administered to wounds 3 times 23 per week until the wound healed. Time of complete wound healing was recorded. Wound 24 size, pain, and edema were assessed at baseline and after 2 and 4 months. The authors 25 stated that this study showed the significant effectiveness of ultrasound therapy in wound 26 healing. Differences between the two ultrasound therapy groups were not statistically 27 significant. White et al. (2015) compared non-contact low-frequency ultrasound therapy to 28 the UK standard of care for venous leg ulcers. Both groups reported a reduction in pain 29 score. The authors suggest that outcome measures favored the non-contact low frequency 30 ultrasound therapy over standard of care, but the differences were not statistically 31 significant. A larger sample size with longer follow up would be prudent to confirm results. 32 33

In a single-site, evaluator-blinded RCT, Gibbons et al. (2015) completed a prospective, 34 randomized, controlled, multicenter trial comparing percent wound size reduction, 35 proportions healed, pain, and quality-of-life (QOL) outcomes in patients randomized to 36 standard care (SC) alone or SC and 40 kHz noncontact, low-frequency ultrasound (NLFU) 37 treatments 3 times per week for 4 weeks. All participants received protocol-defined SC 38 39 compression (30-40 mm Hg), dressings to promote a moist wound environment, and sharp debridement at the bedside for a minimum of 1 time per week. After 4 weeks of treatment, 40 average wound size reduction was $61.6\% \pm 28.9$ in the NLFU+SC compared to $45\% \pm 32.5$ 41 in the SC group (P = 0.02). Reductions in median (65.7% versus 44.4%, P = 0.02) and 42

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absolute wound area (9.0 cm² versus 4.1 cm², P = 0.003) as well as pain scores (from 3.0 1 to 0.6 versus 3.0 to 2.4, P = 0.01) were also significant. NLFU therapy with guideline-2 defined standard care should be considered for healing venous leg ulcers not responding to 3 SC alone. Rastogi et al. (2019) compared the efficacy of noncontact, low-frequency 4 airborne ultrasound (Glybetac) therapy with sham therapy added to standard treatment in 5 patients with neuropathic, clinically infected, or noninfected diabetic foot ulcer (DFU) 6 (wound size >2 cm2), Wagner grades 2 and 3. Patients received ultrasound or sham therapy 7 for 28 days dosed daily for first 6 days followed by twice a week for next 3 weeks along 8 with standard of care. The primary outcome was percentage of patients with at least >50%9 decrease in wound area at 4 week of intervention. Fifty-eight patients completed the study 10 11 protocol. A >50% reduction in wound area was observed in 97.1% and 73.1% subjects in ultrasound and sham groups, respectively. Wound contraction was faster in the first 2 12 weeks with ultrasound therapy, 5.3 cm2, compared with 3.0 cm2 with sham treatment. 13 Authors concluded that the airborne low-frequency ultrasound therapy improves and 14 hastens the healing of chronic neuropathic DFU when combined with standard wound care. 15

16

Kotronis and Vas (2021) evaluated the current evidence behind the NCLFU. Several 17 studies, especially those evaluating NCLFU technology, have demonstrated the potential 18 of ultrasound debridement to effectively remove devitalized tissue, control bioburden, 19 20 alleviate pain, and expedite healing. However, most of the studies are underpowered, involve heterogeneous ulcer types, and demonstrate significant methodological limitations 21 making comparison between studies difficult. Future clinical trials on ultrasound 22 debridement technology must address the design issues prevalent in current studies, and 23 report on clinically relevant endpoints before adoption into best-practice algorithms can be 24 recommended. 25

26

27 Chen et al. (2023) performed a meta-analysis to evaluate the effect of low-frequency ultrasound as an added treatment for chronic wounds. A systematic literature search up to 28 May 2022 was performed with 838 subjects with chronic wounds at the baseline of the 29 studies; 412 of them were using the low-frequency ultrasound (225 low-frequency high-30 intensity contact ultrasound for diabetic foot wound ulcers, and 187 low-frequency low-31 intensity non-contact ultrasound for a venous leg wound ulcers), and 426 were using 32 33 standard care (233 sharp debridement for diabetic foot wound ulcers and 193 sham treatments for venous leg wound ulcers). The low-frequency high-intensity contact 34 ultrasound for diabetic foot wound ulcers had significantly lower non-healed diabetic foot 35 wound ulcers at ≥ 3 months and a higher percentage of diabetic foot wound ulcers area 36 reduction compared with sharp debridement for diabetic foot wound ulcers. The low-37 frequency low-intensity non-contact ultrasound for a venous leg wound ulcers had a 38 39 significantly lower non-healed venous leg wound ulcers at ≥ 3 months and higher percentage venous leg wound ulcers area reduction compared with sham treatments for a 40 venous leg wound ulcers. The analysis of outcomes should be viewed with caution because 41

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1 of the low sample size of all the 17 studies in the meta-analysis and a low number of studies

- 2 in certain comparisons.
- 3

4 Ultrasound

A randomized controlled study of 305 subjects explored the efficacy of physical methods 5 for healing venous leg ulcers, including high-voltage electrical stimulation, ultrasound, and 6 low-level laser therapy, which was performed for 7 weeks (once a day, 6 days a week). 7 Results indicated high-voltage stimulation and ultrasound therapy are useful methods in 8 the conservative treatment of venous leg ulcers (Taradaj et al., 2012). Polak et al. (2014) 9 evaluated the effectiveness of ultrasound in the treatment of Stage II and Stage III pressure 10 11 ulcers in geriatric patients. Participants (age range of 71 to 95 years,) all with wounds that did not respond to previous treatment for at least 4 weeks, were randomly assigned to the 12 treatment group or control group. All patients received standard wound care (SWC); with 13 the treatment group also receiving ultrasound (1 MHz, 0.5 W/cm2, duty cycle of 20 %, 1 14 to 3 minutes/cm2; 1 session per day, 5 days a week). Patients were monitored for 6 weeks 15 or until wounds closed. Percent change in wound surface area (WSA), the weekly rate of 16 change in WSA, and the percentage of pressure ulcers that improved (i.e., decreased in size 17 by at least 50 % or closed) were used to compare differences. After 6 weeks of treatment, 18 the WSA of pressure ulcers decreased significantly in both groups with significantly 19 20 greater improvement in the treatment group (an average of 68.80 $\% \pm 37.23 \%$ compared with 37.24 % \pm 57.84 %; p = 0.047). The mean weekly change of WSA was greater in the 21 treatment group as well, but only for Stage II pressure ulcers than in the control group. The 22 authors concluded that the findings of this study showed US therapy can reduce the WSA 23 of pressure ulcers regardless of their shape, but further research is needed to establish how 24 ultrasound influences the healing of Stage III and Stage IV pressure ulcers. Tricco et al. 25 (2015) identified effective interventions to treat complex wounds through an overview of 26 systematic reviews. Overall, 99 systematic reviews were included; 54 were systematic 27 reviews with a meta-analysis (including data on over 54,000 patients) and 45 were 28 systematic reviews without a meta-analysis. Overall, 4% of included reviews were rated as 29 being of high quality (AMSTAR score greater than or equal to 8). Based on data from 30 systematic reviews including a meta-analysis with an AMSTAR score greater than or equal 31 to 8, promising interventions for complex wounds were identified. These included 32 33 bandages or stockings (multi-layer, high compression) and wound cleansing for venous leg ulcers; 4-layer bandages for mixed arterial/venous leg ulcers; biologics, ultrasound, and 34 hydrogel dressings for diabetic leg/foot ulcers; hydrocolloid dressings, electrotherapy, air-35 fluidized beds, and alternate foam mattresses for pressure ulcers; and silver dressings and 36 37 ultrasound for unspecified mixed complex wounds.

38

Chen et al. (2023) assessed the effect of ultrasound-supported wound debridement (USSD) in subjects with diabetic foot ulcer (DFU) in a meta-analysis. The selected studies contained 577 subjects with DFUs, 282 of them were using USSD, 204 were using standard care, and 91 were using a placebo. The USSD applied to DFU caused a significantly higher

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wound healing rate compared with the standard care with no heterogeneity and the placebo
with no heterogeneity. The USSD applied to DFUs caused a significantly higher wound
healing rate compared with the standard care and the placebo. Though cautions should be

- taken when interpreting these results given low sample sizes of included studies.
- 5

6 Low-Level Laser Therapy (LLLT)

Many researchers have proposed that low-level laser therapy (LLLT) may be an effective 7 treatment modality to promote wound healing and pain relief (Enwemeka, 2004; Hopkins, 8 2004; Posten, 2005). Samsun et al. (AHRQ, 2004) provided an overview of clinical and 9 methodological issues relevant to evaluating the evidence on interventions for wound 10 11 healing. The objective of this evidence report was to systematically review and synthesize the available evidence on the effectiveness of low-level laser treatment and vacuum-12 assisted closure for wound healing. Overall, the studies that met selection criteria for low-13 level laser were poor and do not permit definitive conclusions on whether low-light laser 14 increases the rate of healing for chronic wounds. The available data suggest that the 15 addition of laser therapy does not improve wound healing, as the vast majority of 16 comparisons in these studies do not report any group differences in the relevant outcomes. 17 With the majority of the studies, the low sample sizes and the lack of trends or patterns of 18 outcomes could be the reason for no definitive conclusions. Low light laser therapy has 19 20 potential to improve wound care, but there are limited reports of outcomes that have been demonstrated in well-controlled randomized trials (AHRO, 2004). Additionally, laser 21 parameters are not consistent from study to study and thus, results in difficulty in drawing 22 conclusions. 23

24

Enwemeka et al. (2004) used statistical meta-analysis to determine the overall treatment 25 effects of laser phototherapy (low-level laser) on tissue repair and pain relief. Thirty-four 26 articles on tissue repair and nine articles on pain control met inclusion criteria. Meta-27 analysis revealed a positive effect of laser phototherapy on tissue repair and pain control. 28 Further, analysis revealed the positive effects of various wavelengths of laser light on tissue 29 repair, with 632.8 nm having the highest treatment effect and 780 nm the least. The overall 30 treatment effect for pain control was positive as well. The authors concluded that laser 31 phototherapy is a highly effective therapeutic modality for tissue repair and pain relief 32 33 (Enwemeka et al., 2004). In another study by Enwemeka (2009), it was reported that inaccurate measurement and incorrect reporting dosages are major shortcomings of 34 phototherapy research. Enwemeka reported that there are as many as 30% of published 35 reports in the field lacking relevant information needed to determine a dosage or that 36 reported dosages that are not accurate. Further studies are needed to determine strategies 37 to improve dosages in the use of low-level laser for tissue repair and pain relief. 38

39

Posten et al. (2005) studied the mechanism and efficacy of low-level laser therapy (LLLT)
for wound healing. This group of researchers critically evaluated reported in vitro models
and in vivo animal and human studies, to assess the qualitative and quantitative sufficiency

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for the efficacy of LLLT in promoting wound healing. After the authors examined the 1 effects of LLLT on cell cultures in vitro, they concluded that some authors report an 2 increase in cell proliferation and collagen production using specific and somewhat arbitrary 3 laser settings with the helium neon (HeNe) and gallium arsenide (GaAs) lasers. Although 4 increases in cell proliferation and collagen production using specific laser settings was 5 reported, it could not be determined which properties (i.e., photothermal, photochemical, 6 or photomechanical) of the LLLT produced the positive effect (Posten et al., 2005). Some 7 studies using HeNe lasers reported improvements in surgical wound healing in a rodent 8 model; however, the results have not been duplicated in animals such as pigs, which have 9 skin that closely resembles that of humans. Studies that involved humans have beneficial 10 11 effects on superficial wound healing found in small case series and have not been replicated in larger studies (Posten et al., 2005). Although applications of high-energy (10-100W) 12 lasers are well established with significant supportive literature and widespread use, 13 conflicting studies in the literature have limited LLLT use in the United States to 14 investigational use only (Posten et al., 2005). 15

16

Another randomized, triple-blind, placebo-controlled design by Hopkins et al. (2004) 17 assessed the putative effects of LLLT on healing using an experimental model. Subjects 18 received LLLT from either a laser or a sham cluster head (8 J/cm2 for 2 minutes, 5 seconds) 19 20 to one of two randomly chosen wounds. Data were analyzed for wound contraction (area), color changes (chromatic red), and luminance. The results for group by wound by time 21 interaction showed at days 6, 8, and 10 follow-up testing revealed that the laser group had 22 smaller wounds (decreased area measurements) than the sham group for both the treated 23 and the untreated wounds. The authors concluded that LLLT resulted in the enhanced 24 wound healing as measured by wound contraction. The untreated wounds in subjects 25 treated with LLLT contracted more than the wounds in the sham group, thus LLLT may 26 produce an indirect healing effect on surrounding tissues. Data indicates that LLLT is an 27 effective modality to facilitate wound contraction of partial thickness wounds (Hopkins et 28 al., 2004). 29

30

A double-blinded RCT of 23 patients with diabetic foot ulcers who were randomly assigned 31 to LLLT or a sham control group. The treatment group received LLLT six times per week 32 33 for a minimum of two consecutive weeks, then laser therapy every other day up to complete healing of the ulcer for a maximum of 20 weeks. After 4 weeks of treatment, the 34 intervention group demonstrated significantly decreased ulcer size, but at 20 weeks, there 35 was no statistically significant difference in ulcer healing time between the two groups. 36 The authors recommended completion of additional studies with larger samples and longer 37 follow-up time (Kaviani et al., 2011). Another randomized controlled study of 34 patients 38 39 with venous leg ulcers demonstrated no significant differences in reduction of ulcer size between the laser treatment and control groups following a 9-week intervention period 40 (LeClere et al., 2010). A randomized controlled study of 305 subjects explored the efficacy 41 of physical methods for healing venous leg ulcers, including high-voltage electrical 42

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stimulation, ultrasound, and low-level laser therapy, which was performed for 7 weeks 1 (once a day, 6 days a week). Results indicated no significant effect or improvement in 2 healing with the use of laser therapy for venous ulcers. (Taradaj et al., 2012). Beckmann et 3 al. (2014) completed a systematic literature review of LLLT for wound healing of diabetic 4 ulcers. They concluded that although the majority of clinical studies show a potential 5 benefit of LLLT in wound healing of diabetic ulcers, there are several aspects in these 6 studies limiting final evidence about the actual outcomes. In summary, all studies give 7 enough evidence to continue research on laser therapy for diabetic ulcers, but clinical trials 8 using human models do not provide sufficient evidence to establish the usefulness of LLLT 9 as an effective tool in wound care regimes at present. Further well-designed research trials 10 are required to determine the true value of LLLT in routine wound care. 11

12

Zhou et al. (2021) aimed to synthesize and systematically review the best evidence to assess 13 the efficacy of low-level light therapy in improving healing of diabetic foot ulcers. Twelve 14 randomized controlled trials were included. Meta-analysis revealed that 30.90% of the 15 ulcer area was significantly reduced in the therapy group compared with the control group 16 with a very large effect. A 4.2 cm² reduction of the ulcer area was observed in the therapy 17 group compared with the control group with a very large effect. In addition, diabetic foot 18 ulcers in the therapy group were 4.65 times more likely to heal completely than those in 19 20 the control group. Authors conclude that low-level light therapy accelerates wound healing and reduces the size of diabetic foot ulcers. However, the review does not allow any 21 recommendation for the best treatment parameters required to achieve improved healing. 22 Future trials need to include a good design and large sample size in defining the optimal 23 treatment parameters for ulcers of different sizes. 24

25

Sutton et al. (2021) provided a comprehensive narrative review and critical appraisal of 26 research investigating photobiomodulation (PBM), formerly known as low level laser 27 therapy which includes lasers and light emitting diodes (LEDs), as a treatment to promote 28 diabetic foot and lower leg ulcer (DFU) healing for humans. A total of 13 studies, with a 29 total of 417 participants, were included in this review. The studies were critically appraised 30 using the PEDro scale, which revealed weaknesses in study designs such as small sample 31 sizes and problems with reproducibility with respect to the laser protocols. Characteristics 32 33 of PBM that improved wound healing were wavelengths of 630 nm-660 nm and infrared wavelengths of 850 or 890 nm, and radiant exposure levels of 3 J/cm2-7 J/cm2. PBM was 34 beneficial for superficial and deep DFUs. Controlled blood glucose levels and adherence 35 to best practices (i.e., pressure off-loading, optimized wound dressing changes, appropriate 36 debridement) could have been a factor in the beneficial outcomes. Authors concluded that 37 regardless of the laser characteristics chosen, in the majority of studies PBM as a treatment 38 39 for DFUs improved healing rate when compared with standard wound care alone. However, weaknesses across the studies indicate that further research is required. 40

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1 Zhang et al. (2024) evaluated the impact of red and infrared light on the healing of DFUs

2 and provided evidence-based recommendations for future clinical adjunctive treatments of

3 DFUs. A total of 28 studies, involving 1,471 patients, were included. The meta-analysis

4 showed that groups treated with red and infrared light had a significantly higher ulcer

- 5 healing rate, shorter ulcer healing time, increased peak blood flow velocity in the dorsalis
- 6 pedis artery, and reduced wound pain score. Authors concluded that the use of red and 7 infrared light as an adjunctive treatment for DFUs is more beneficial than conventional
- 8 wound care. However, due to limitations in the quality and sample size of the included
- 9 studies, further high-quality research is needed to validate these conclusions.
- 10

11 Negative Pressure Wound Therapy (NPWT)

Negative Pressure Wound Therapy (NPWT) is used to describe the treatment of a wound 12 with topical negative pressure including atmospheric pressure therapy or dressing, vacuum 13 sealing technique, foam suction dressing, vacuum compression, vacuum pack, sealed 14 surface wound suction or sealing aspirative therapy (National Institute for Health and 15 Clinical Excellence, 2005). The principles of the application of NPWT to a wound may aid 16 in the healing process due to the following mechanisms: 1) wound contraction, 2) 17 stimulation of granulation tissue formation, 3) continuous wound cleansing after adequate 18 primary surgical debridement, 4) continuous removal of exudates, and 5) reduction of 19 20 interstitial edema (AHQR, 2009; Willy et al., 2007). NPWT is primarily intended for chronic wounds that have not healed when treated with either standard care or other forms 21 of wound care (ECRI, 2009). The development of negative pressure techniques for wound 22 healing derives from two theories: removal of wound exudates while decreasing edema 23 and concentrations of inhibitory factors and increasing blood flow; and negative pressure 24 stretches and deforms the tissue and disturbs the extracellular matrix which induces 25 biochemical responses that promote wound healing (ERCI, 2009). 26

27

The Centers of Medicare and Medicaid Services partnered with the Agency for Health 28 Research and Quality (AHRQ) to commission a review of NPWT devices. AHRQ 29 contracted with the Institute Evidence-based Practice Center to perform the review 30 (AHRQ, 2009). The report specifically examined the use of NPWT for treatment of the 31 following wound types: diabetic foot ulcers, pressure ulcers, vascular ulcers (both venous 32 33 and arterial), burn wounds, surgical wounds (particularly infected sternal wounds) and trauma-induced wounds. This technology assessment report on NPWT found that the 34 systematic reviews of NPWT reveal several important points about the use of NPWT 35 modality. First, all the systematic reviews noted a lack of high-quality clinical evidence 36 supporting the advantages of NPWT compared to the other wound treatments. The lack of 37 high-quality evidence resulted in many of the systematic reviewers relying on low-quality 38 retrospective studies to judge the efficacy of NPWT technology. Secondly, the other 39 systematic reviews found no studies published that directly compared the different types 40 of NPWT devices or components. Direct comparison studies are needed to help determine 41 the importance of the dressing approaches (foam or gauze) that may provide the best 42

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potential for wound healing. Thirdly, other systemic reviews concluded that NPWT must 1 be evaluated according to wound type. Wound healing varies according to the type of 2 wound being treated and NPWT benefits described for one type of wound cannot be 3 transferred to other wound types (AHRQ, 2009). The overall assessment concluded that 4 the available evidence cannot be used to determine a significant therapeutic distinction of 5 a particular NPWT system (AHRQ, 2009). Due to lack of studies comparing one NPWT 6 system to another NPWT system, the severity of adverse events for one NPWT compared 7 to another could not be determined (AHRQ, 2009). 8

9

A multi-center randomized controlled study by Blume et al. (2008) evaluated the safety 10 and clinical efficacy of NPWT compared with advanced moist wound therapy (AMWT) 11 (predominately hydrogels and alginates) to treat foot ulcers in diabetic patients. Complete 12 ulcer closure was defined as skin closure (100% reepithelization) without drainage or 13 dressing requirements. Patients were randomly assigned to either NPWT or AMWT and 14 received standard off-loading as needed. The trial evaluated treatment until day 112 or 15 ulcer closure by any means. Patients whose wounds achieved ulcer closure were followed 16 at 3 and 9 months. The authors showed a greater proportion of the foot ulcers achieved 17 complete ulcer closure with NPWT than with AMWT within the 112-day active treatment 18 phase. The patients that received the NPWT experienced significantly fewer secondary 19 20 amputations. In assessing the overall safety, no significant difference between the groups was observed in treatment-related complications such as infection, cellulitis, and 21 osteomyelitis at 6 months. The authors of this study concluded that NPWT appears to be 22 as safe as and more efficacious than AMWT for the treatment of diabetic foot ulcers 23 (Blume et al., 2008). In 2015, a Cochrane review was completed by Dumville et al. on 24 NPWT for treating pressure ulcers in any care setting. Authors concluded that there is 25 currently no high quality RCT available regarding the effects of NPWT compared to 26 alternatives for the treatment of pressure ulcers. Also, they express that high uncertainty 27 remains about the potential benefits or harms or both of treatment using NPWT. An update 28 of the Cochrane review was completed in 2019. Despite the addition of 25 trials, results 29 were consistent with the earlier review, with the evidence judged to be of low or very low 30 certainty for all outcomes. Consequently, uncertainty remains about whether NPWT 31 compared with a standard dressing reduces or increases the incidence of important 32 33 outcomes such as mortality, dehiscence, seroma, or if it increases costs.

34

The US Food and Drug Administration (FDA) issued a Preliminary Public Health 35 Notification: Serious Complications Associated with NPWT Systems. The FDA issued the 36 alert to make individuals aware of deaths and serious complications, especially bleeding 37 and infection, associated with the use of NPWT systems, and to provide recommendations 38 39 to reduce the risk (FDA, 2009; FDA, 2011). Although complications are rare, if NPWT is not used properly by trained medical personnel, complications can occur. The FDA 40 recommends selecting patients for NPWT carefully, after reviewing the most recent device 41 labeling and instructions, and that the patient is monitored frequently in an appropriate care 42

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1 setting by trained practitioner. The patient's condition, including the wound status, wound

2 location, and co-morbidities must be considered and monitored prior and during NPWT

3 treatment. The FDA recommends numerous patient risk factors/characteristics need to be

- 4 considered before the use of NPWT. The FDA recommends that NPWT is contraindicated
- 5 for these wound types/conditions:
- 6 Necrotic tissue with eschar present
- 7 Untreated osteomyelitis
- 8 Non-enteric and unexplored fistulas
- 9 Malignancy in the wound
- 10 Exposed vasculature
- 11 Exposed nerves
 - Exposed anastomotic site
 - Exposed organs, such as eyes
- 13 14

12

The FDA issued an updated report (February 2011) on the original Preliminary Public 15 Health Notification: Serious Complications Associated with NPWT Systems, issued in 16 2009. The FDA received reports of an additional six deaths and 97 injuries, for a total of 17 12 deaths and 174 injury reports since 2007. The new recommendation was in regard to 18 the safety and effectiveness of NPWT systems in newborns, infants and children; safety 19 and effectiveness has not been established at this time and currently there are no NPWT 20 systems cleared for use in these pediatric populations. The FDA will continue to monitor 21 adverse events associated with NPWT systems and will make available any new 22 23 information that might affect their use (FDA, 2009; FDA, 2011).

24

25 A systematic review of interventions to enhance healing of chronic ulcers of the foot in patients with diabetes concluded that overall, the heterogeneity and poor methodology 26 made it difficult to draw conclusions (Game et al., 2012). Forty-three studies were selected 27 for full review. They identified 10 categories: sharp debridement and wound bed 28 29 preparation with larvae and hydrotherapy; wound bed preparation using antiseptics, applications and dressing products; resection of the chronic wound; hyperbaric oxygen 30 therapy (HBOT); compression or negative pressure therapy; products designed to correct 31 aspects of wound biochemistry and cell biology associated with impaired wound healing; 32 application of cells, including platelets and stem cells; bioengineered skin and skin grafts; 33 electrical, electromagnetic, lasers, shockwaves and ultrasound; other systemic therapies 34 35 which did not fit in the above categories. Thus, for this specific condition and type of wound, conclusions as to the best evidence of treatment interventions are not possible due 36 to lack of controlled studies and design issues (Game et al., 2012). 37

38

Seidel et al. (2020) evaluated effectiveness and safety of negative pressure wound therapy (NPWT) in patients with diabetic foot wounds in clinical practice. Three hundred sixtyeight patients were randomized, and 345 participants were included in the modified intention-to-treat (ITT) population. Adult patients suffering from a diabetic foot ulcer at

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least for 4 weeks and without contraindication for NPWT were allowed to be included. NPWT was compared with standard moist wound care (SMWC) according to local (Germany) standards and guidelines. Primary outcome was wound closure within 16 weeks. Secondary outcomes were wound-related and treatment-related adverse events (AEs), amputations, time until optimal wound bed preparation, wound size and wound tissue composition, pain, and quality of life (QoL) within 16 weeks, and recurrences and wound closure within 6 months.

8

Authors concluded that NPWT was not superior to SMWC in diabetic foot wounds in 9 German clinical practice. Overall, wound closure rate was low. Documentation deficits and 10 11 deviations from treatment guidelines negatively impacted the outcome wound closure. Norman et al. (2020) assessed the effects of NPWT for preventing surgical site infections 12 (SSI) in wounds healing through primary closure, and to assess the cost-effectiveness of 13 NPWT in wounds healing through primary closure. Trials were included if they allocated 14 participants to treatment randomly and compared NPWT with any other type of wound 15 dressing or compared one type of NPWT with another type of NPWT. In this third update, 16 15 new randomized controlled trials (RCTs) and three new economic studies were added, 17 resulting in a total of 44 RCTs (7,447 included participants) and five economic studies. 18 Studies evaluated NPWT in the context of a wide range of surgeries including orthopaedic, 19 20 obstetric, vascular, and general procedures. All studies compared NPWT with standard dressings. Most studies had unclear or high risk of bias for at least one key domain. Authors 21 concluded that people experiencing primary wound closure of their surgical wound and 22 treated prophylactically with NPWT following surgery probably experience fewer SSI than 23 people treated with standard dressings (moderate-certainty evidence). There is no clear 24 difference in number of deaths or wound dehiscence between people treated with NPWT 25 and standard dressings (low-certainty evidence). There are also no clear differences in 26 secondary outcomes where all evidence was low or very low certainty. Most evidence on 27 pain is very low-certainty, but there is probably no difference in pain between NPWT and 28 standard dressings after surgery for lower limb fracture (moderate-certainty evidence). 29

30

Zens et al. (2020) performed a systematic review of randomized controlled trials (RCTs) 31 comparing the patient-relevant benefits and harms of NPWT with standard wound therapy 32 33 (SWT) in patients with wounds healing by secondary intention. Forty-eight eligible studies of generally low quality with evaluable data for 4,315 patients and 30 eligible studies with 34 missing data for at least 1386 patients were identified. A meta-analysis of all wound healing 35 data showed a significant effect in favor of NPWT. There was neither proof (nor indication 36 nor hint) of greater benefit or harm of NPWT for other patient-relevant outcomes such as 37 mortality and adverse events. Authors concluded that low-quality data indicate a greater 38 39 benefit of NPWT versus SWT for wound closure in patients with wounds healing by secondary intention. The length of hospital stay is also shortened. The data show no 40 advantages or disadvantages of NPWT for other patient-relevant outcomes. Publication 41

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1 bias is an important problem in studies on NPWT, underlining that all clinical studies need

- 2 to be fully reported.
- 3

Pedrazi et al. (2021) completed a systematic review, including a total of 466 patients, which 4 shows that NPWT as the initial treatment for burned children and after skin grafting has 5 been shown to produce promising results. In the majority of studies, skin graft take rate is 6 close to 100%. This therapy is particularly beneficial in the pediatric population because 7 of less frequent dressing changes and early mobilization. Authors note that NPWT is not 8 in the subject of controlled clinical trials in pediatric; most publications are case reports or 9 retrospective reviews. The sporadic complications include bleeding, local infections, and 10 mechanical device issues. Prospective randomized studies are needed to provide validated 11 rules. Putri et al. (2022) reviewed the risks and benefits of NPWT in surgical wounds with 12 the underlying malignant disease compared with conventional wound care (CWC). The 13 first outcome was wound complications, divided into surgical site infection (SSI), seroma, 14 hematoma, and wound dehiscence. The secondary outcome was hospital readmission. 15 Thirteen observational studies with 1,923 patients and seven RCTs with 1,091 patients 16 were included. NPWT group showed significant decrease in the risk of SSI and seroma in 17 observational studies with P value <0.05, as well as RCTs but were not significant. Wound 18 dehiscence and hospital readmission showed lower risks in NPWT group but were not 19 20 significant. Hematoma showed no significant difference. Authors concluded that NPWT is not contraindicated in cancer surgical wounds and can be considered a beneficial palliative 21 treatment to promote wound healing. Gillespie et al. (2022) summarized the evidence on 22 the effectiveness of negative pressure wound therapy (NPWT) for preventing SSI and other 23 wound complications in obese women after CS. Ten RCTs with 5,583 patients were 24 included; studies were published between 2012 and 2021. Nine RCTs with 5,529 patients 25 were pooled for the outcome SSI. Meta-analysis results suggest a significant difference 26 favoring the NPWT group, indicating an absolute risk reduction of 1.8% among those 27 receiving NPWT compared with usual care. The risk of blistering in the NPWT group was 28 significantly higher. All studies had high risk of bias relative to blinding of 29 personnel/participants. Only 40% of studies reported blinding of outcome assessments and 30 50% had incomplete outcome data. Authors concluded that the decision to use NPWT 31 should be considered both in terms of its potential benefits and its limitations. 32

33

Shi et al. (2023) evaluated the effectiveness of NPWT for treating adult with pressure ulcers 34 in any care setting in a Cochrane Review. Authors included published and unpublished 35 randomized controlled trials (RCTs) comparing the effects of NPWT with alternative 36 treatments or different types of NPWT in the treatment of adults with pressure ulcers (stage 37 II or above). This review included eight RCTs with a total of 327 randomized participants. 38 39 Six of the eight included studies were deemed to be at a high risk of bias in one or more risk of bias domains, and evidence for all outcomes of interest was deemed to be of very 40 low certainty. Most studies had small sample sizes (range: 12 to 96, median: 37 41 participants). Five studies compared NPWT with dressings, but only one study reported 42

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usable primary outcome data (complete wound healing and adverse events). This study had 1 only 12 participants and there were very few events; only one participant was healed in the 2 study (risk ratio (RR) 3.00, very low-certainly evidence). There was no evidence of a 3 difference in the number of participants with adverse events in the NPWT group and the 4 dressing group, but the evidence for this outcome was also assessed as very low certainty. 5 Changes in ulcer size, pressure ulcer severity, cost, and pressure ulcer scale for healing 6 (PUSH) sores were also reported, but authors were unable to draw conclusions due to the 7 low certainly of the evidence. One study compared NPWT with a series of gel treatments, 8 but this study provided no usable data. Another study compared NPWT with 'moist wound 9 healing', which did not report primary outcome data. Changes in ulcer size and cost were 10 11 reported in this study, but evidence was assessed as being of very low certainty; One study compared NPWT combined with internet-plus home care with standard care, but no 12 primary outcome data were reported. Changes in ulcer size, pain, and dressing change 13 times were reported, but evidence was assessed as being of very low certainty. None of the 14 included studies reported time to complete healing, health-related quality of life, wound 15 infection, or wound recurrence. Authors concluded that the efficacy, safety, and 16 acceptability of NPWT in treating pressure ulcers compared to usual care are uncertain due 17 to the lack of key data on complete wound healing, adverse events, time to complete 18 healing, and cost-effectiveness. Compared with usual care, using NPWT may speed up the 19 20 reduction of pressure ulcer size and severity of pressure ulcer, reduce pain, and dressing change times. Still, trials were small, poorly described, had short follow-up times, and with 21 a high risk of bias; any conclusions drawn from the current evidence should be interpreted 22 with considerable caution. In the future, high-quality research with large sample sizes and 23 low risk of bias is still needed to further verify the efficacy, safety, and cost-effectiveness 24 of NPWT in the treatment of pressure ulcers. Future researchers need to recognize the 25 importance of complete and accurate reporting of clinically important outcomes such as 26 the complete healing rate, healing time, and adverse events. 27

28

Horn et al. (2023) examined the use of negative pressure wound therapy for the treatment 29 of venous leg ulcers (VLU). Authors report that NPWT is underrecognized as a useful 30 adjunct in the management of VLUs. The literature has shown NPWT to be beneficial by 31 primarily reducing wound area while promoting granulation tissue formation; thus, this 32 33 therapy is a valuable adjunct in preparing the wound for either a cellular and tissue-based therapy and, more notably, for Split-Thickness Skin Grafts (STSG). This is likely 34 especially true for large VLUs. Although what is considered large may be somewhat 35 arbitrary, it appears that the benefit of NPWT increases with wound size. Management of 36 fluid and drainage appears to be a secondary reason to use NPWT. Most clinicians who 37 treat VLUs with adjunctive NPWT use it in conjunction with multilayer compression. It is 38 39 well recognized that increasing venous return with multilayer compression is mandatory for good ulcer healing. Thus, in any setting other than the inpatient hospital setting, for 40 most clinicians adjunctive NPWT is best used in addition to compressive dressing when 41 treating VLUs. 42

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Onderková et al. (2023) aimed to systematically review NPWT effectiveness, safety, and 1 comparative efficacy for head and neck wound healing. Thirty-one studies from a 2 systematic literature search were identified and analyzed for wound healing response, 3 overall success rate, improvements compared to conventional wound care, and variation in 4 pressure settings, treatment lengths, and dressing change frequency. NPWT showed 5 enhanced outcomes across diverse head and neck wounds, particularly complex post-6 reconstructive wounds, and severe infections. Despite the predominantly case report/series 7 evidence and lack of standardized NPWT protocols, its benefits over conventional care 8 were clear. NPWT emerges as a promising approach for head and neck wound 9 management, potentially improving patient outcomes and reducing complications. More 10 11 randomized controlled trials are needed to solidify the evidence and standardize NPWT application protocols. 12

13

Chen et al. (2024) updated the 2019 IWGDF evidence-based guideline on wound healing 14 interventions to promote healing of foot ulcers in persons with diabetes. Each 15 recommendation is based on the evidence found in the systematic review and, using the 16 GRADE summary of judgement items, including desirable and undesirable effects, 17 certainty of evidence, patient values, resources required, cost effectiveness, equity, 18 feasibility, and acceptability, recommendations were formulated that were agreed by the 19 20 authors and reviewed by independent experts and stakeholders. Authors made a number of conditional supportive recommendations for the use of interventions to improve healing of 21 foot ulcers in people with diabetes. These include the use of sucrose octasulfate dressings, 22 the use of negative pressure wound therapies for post-operative wounds, the use of 23 placental-derived products, the use of the autologous leucocyte/platelet/fibrin patch, the 24 use of topical oxygen therapy, and the use of hyperbaric oxygen. Although in all cases it 25 was stressed that these should be used where best standard of care was not able to heal the 26 wound alone and where resources were available for the interventions. 27

28

Wu et al. (2024) investigated the efficacy and safety of extracorporeal shockwave therapy 29 (ESWT) for DFUs. A total of 10 RCTs with moderate methodological quality were 30 included for data analysis. The findings showed that ESWT was significantly associated 31 with significantly complete healed ulcers and lower rate of unchanged ulcers compared to 32 33 controls. Subgroup analysis further revealed that ESWT was better than both hyperbaric oxygen therapy (HOT) and the standard of care (SOC). Moreover, ESWT also significantly 34 improved the average transcutaneous partial oxygen pressure. However, the rate of ≥ 50 % 35 improved ulcers and treatment-emergent adverse events were not significantly different 36 37 between the ESWT and controls. Authors concluded that ESWT has shown promising efficacy and a favorable safety profile in the treatment of DFUs. 38

39

40 Systemic Hyperbaric Oxygen Therapy (HBOT)

41 Systemic hyperbaric oxygen therapy (HBOT) involves the inhalation of pure oxygen gas 42 while enclosed in a high-pressure chamber (defined as pressure greater than standard

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atmospheric pressure). The pressures used are usually between 1.4 to 3.0 atmospheres 1 absolute (atm abs or ATA). The therapy works by supersaturating the blood tissues with 2 oxygen via increased atmospheric pressure as well as increased oxygen concentrations. 3 Studies have demonstrated that this therapy increases the available oxygen to the body by 4 10 to 20 times normal levels. Treatment may be carried out in either a mono-place chamber 5 pressurized with pure oxygen or in a larger, multi-place chamber pressurized with 6 compressed air, in which case the individual receives pure oxygen by mask, head tent, or 7 endotracheal tube. The number and duration of treatment sessions and the atmospheric 8 pressure during treatment varies depending on the specific condition being treated, the 9 severity of the condition, and the procedures developed by individual hospitals and clinics. 10 11 These individual procedures vary widely and have made the evaluation of the efficacy of hyperbaric oxygen therapy difficult. However, the medical specialty society which 12 represents the physicians who specialize in this type of medical treatment, called the 13 Undersea and Hyperbaric Medical Society (UHMS), created treatment recommendations 14 for a wide variety of conditions for which HBOT has been proven to provide significant 15 benefits. 16

17

The position regarding systemic hyperbaric oxygen is based on guidelines published by the Undersea and Hyperbaric Medical Society (2008). These guidelines provide recommendations for indications where hyperbaric oxygen therapy has been demonstrated to provide clinical benefits, and where there is adequate data to provide guidance regarding treatment duration, frequency, and depth of pressurization.

23

24 Lalieu et al. (2021) completed a retrospective, single-center cohort study between 2013 and 2019. All patients with a venous leg ulcer (VLU) from an outpatient clinic providing HBOT 25 and wound care were included. The primary outcome measure was wound healing, 26 determined at discharge from the center. Other outcome measures were improvement in 27 patient related outcome measures (PROMs), as assessed by the EQ-5D-3L questionnaire 28 and including quality of life (QoL) and pain score. Fifty patients were included, 53% 29 female, with a mean age of $73.4 (\pm 12.2)$. Most wounds (83%) had existed longer than 3 30 months before starting treatment. Patients received an average of 43 (±20) sessions of 31 HBOT. After treatment, 37 patients (63%) achieved complete or near-complete wound 32 healing. Wound size decreased from a median of 14 cm^2 to 0.5 cm^2 , a median decrease of 33 7.5 cm^2 (94%). Patients mostly reported improvement for all health aspects on the 34 questionnaire. Pain score decreased from 5.7 (\pm 2.5) to 2.1 (\pm 2.2) and health score increased 35 from 57.2 (± 15.6) to 69.9 (± 18.9). Authors concluded that patients with non-healing VLUs 36 may benefit from HBOT to achieve complete or substantial wound healing. They 37 recommend a well-designed randomized clinical trial with several patients allowing 38 39 enough statistical power, and of a reasonable duration, to establish the potential of additional HBOT on hard-to-heal venous ulcers. 40

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It is critical that interventions used to enhance the healing of chronic foot ulcers in diabetes 1 are backed by high-quality evidence and cost-effectiveness. In previous years, the 2 systematic review accompanying guidelines published by the International Working Group 3 of the Diabetic Foot performed 4-yearly updates of previous searches, including trials of 4 prospective, cross-sectional, and case-control design. Due to a need to re-evaluate older 5 studies against newer standards of reporting and assessment of risk of bias, Chen et al. 6 (2024) performed a whole new search from conception but limiting studies to randomized 7 control trials only. The literature search identified 22,250 articles, of which 262 were 8 selected for full text review across 10 categories of interventions. Overall, the certainty of 9 evidence for a majority of wound healing interventions was low or very low, with moderate 10 11 evidence existing for two interventions (sucrose-octasulfate and leucocyte, platelet and fibrin patch) and low-quality evidence for a further four (hyperbaric oxygen, topical 12 oxygen, placental derived products and negative pressure wound therapy). The majority of 13 interventions had insufficient evidence. Overall, the evidence to support any other 14 intervention to enhance wound healing is lacking and further high-quality randomized 15 control trials are encouraged. 16

17

Lalieu et al. (2023) analyzed wound healing results of hyperbaric oxygen therapy (HBOT) 18 for a variety of different wound types. This retrospective cohort study included all patients 19 20 treated with HBOT and wound care at a single hyperbaric center between January 2017 and December 2020. The primary outcome was wound healing. Secondary outcome 21 measures were quality of life (OoL), number of sessions, adverse effects, and treatment 22 cost. Investigators also examined possible influencing factors, including age, sex, type and 23 24 duration of wound, socioeconomic status, smoking status, and presence of peripheral vascular disease. A total of 774 treatment series were recorded, with a median of 39 25 sessions per patient. In total, 472 wounds (61.0%) healed, 177 (22.9%) partially healed, 41 26 (5.3%) deteriorated, and 39 (5.0%) minor and 45 (5.8%) major amputations were 27 performed. Following HBOT, median wound surface area decreased from 4.4 cm 2 to 0.2 28 cm 2, and patient QoL improved from 60 to 75 on a 100-point scale. Frequently recorded 29 adverse effects were fatigue, hyperoxic myopia, and middle ear barotrauma. Attending 30 fewer than 30 sessions and having severe arterial disease were both associated with a 31 negative outcome. Authors concluded that adding HBOT to standard wound care increases 32 33 wound healing and QoL in selected wounds. Patients with severe arterial disease should be screened for potential benefits. Most reported adverse effects are mild and transient. 34

35

Kwee et al. (2024) evaluated the effectiveness of hyperbaric oxygen therapy in the management of severe lower limb soft tissue injuries. In total 7 studies met the inclusion criteria, involving 229 patients. The studies included 2 randomized clinical trials, 1 retrospective cohort study, 3 case series and 1 case report. The randomized placebocontrolled clinical trial showed a significant increase in wound healing and decrease in the need for additional surgical interventions in the patient group receiving hyperbaric oxygen therapy when compared to those undergoing sham therapy. The randomized non-placebo-

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controlled clinical trial revealed that early hyperbaric oxygen therapy reduces tissue 1 necrosis and the likelihood of long-term complications. The retrospective cohort study 2 indicated that hyperbaric oxygen therapy effectively reduces infection rates and the need 3 for additional surgical interventions. The case series and case report presented beneficial 4 results with regard to wound healing when hyperbaric oxygen therapy was added to the 5 treatment regimen. Authors concluded that hyperbaric oxygen therapy is generally 6 considered a safe therapeutic intervention and seems to have a beneficial effect on wound 7 healing in severe lower limb soft tissue injuries when implemented as an addition to 8 standard trauma care. 9

10

Ogbeide et al. (2024) summarized the current management principles and the development 11 of new approaches to care in a narrative review. Authors noted that the management of 12 DFUs has significantly advanced over time, with a clear trend toward a compact, patient-13 centered, multidisciplinary approach. Optimal glycemic control, infection control, pressure 14 redistribution, re-vascularization, wound care, and debridement remain key to preventing 15 and managing DFUs. Emerging trends like hyperbaric oxygen therapy, negative wound 16 pressure therapy, skin substitutes, and growth factor therapy are promising, and there is a 17 need for further randomized and observational studies. 18

19

20 Damineni et al. (2025) assessed the primary clinical evidence supporting hyperbaric oxygen therapy (HBOT) in the management of DFUs. Six studies with a total of 391 21 patients were included in the final analysis, after applying relevant inclusion and exclusion 22 criteria. The majority of the studies indicated reduced major amputation rates, improved 23 ulcer healing rates, and decreased ulcer size and depth with HBOT compared to standard 24 care (SC). Most studies indicate that HBOT leads to lower rates of major amputations, 25 better ulcer healing, and reduced ulcer dimensions than SC. However, one study found no 26 significant differences in amputation rates or long-term wound healing between groups. 27 While most studies showed a low risk of bias in certain areas, moderate-to-high bias in key 28 aspects necessitated careful interpretation. Future high-quality RCTs with stringent 29 blinding, standardized protocols, and defined patient selection criteria are crucial to 30 confirm the effectiveness of HBOT, improve guidelines, and establish its long-term 31 viability. Although this review suggests that HBOT may be valuable for DFUs, additional 32 33 rigorous research is needed to reduce bias, enhance methodological consistency, and improve the reliability of the findings for clinical implementation. 34

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36 Undersea and Hyperbaric Medical Society Guidelines

The Undersea and Hyperbaric Medical Society's (UHMS) 2008 Hyperbaric Oxygen Therapy Committee suggests utilization of systemic hyperbaric oxygen therapy pressurization or 'HBOT' guidelines as described below regarding wound care:

- 40
- Arterial Insufficiencies Treatment varies depending upon the severity of the condition
 and the type of chamber used. In large multi-place chambers, treatments delivered between

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 2.0 and 2.5 ATA of oxygen for 90-120 minutes once or twice daily is standard. In monoplace chambers, treatment at 2.0 ATA of oxygen for 90-120 minutes once or twice daily is standard. Once the patient is stabilized, once daily treatment is recommended. Details of

4 specific conditions are below:

- 5 a. Diabetic lower extremity wounds
 - Patient with type 1 or type 2 diabetes with lower extremity wound due to diabetes; and
 - Wegner grade III or higher wound severity; and
- 9 o Patient has failed an adequate course of standard wound therapy (defined as
 30 days of standard treatment including assessment and correction of
 vascular abnormalities, optimization of nutritional status and glucose
 control, debridement, moist wound dressing, off-loading, and treatment of
 infection; and
- 14 Re-evaluations at 30 days must show continued progress.
- b. Arterial insufficiency ulcers May benefit patients who have persistent hypoxia
 despite attempts at increasing blood flow or when wound failure continues despite
 maximum revascularization.
- c. Pressure ulcers Not recommended for the routine treatment of decubitus ulcers.
 May be necessary for support of skin flaps and grafts showing evidence of ischemic
 failure, when the ulcer develops in the field of previous irradiated area for pelvic or
 perineal malignancies, or when progressive necrotizing soft tissue infection or
 refractory osteomyelitis is present.
 - d. Venous stasis ulcers May be required to support skin grafting in patients with concomitant peripheral arterial occlusive disease and hypoxia not corrected by control of edema.
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Stoekenbroek et al. (2014) completed a systematic review of randomized clinical trials 27 (RCTs) to assess the additional value of hyperbaric oxygen therapy (HBOT) in promoting 28 the healing of diabetic foot ulcers and preventing amputations was performed. Eligible 29 studies reported the effectiveness of adjunctive HBOT with regard to wound healing, 30 amputations, and additional interventions. Seven of the 669 identified articles met the 31 inclusion criteria, comprising 376 patients. Authors concluded that current evidence shows 32 some evidence of the effectiveness of HBOT in improving the healing of diabetic leg ulcers 33 in patients with concomitant ischemia. Larger trials of higher quality are needed before 34 implementation of HBOT in routine clinical practice in patients with diabetic foot ulcers 35 can be justified. A Cochrane Review (2015) by Kranke et al. assessed the benefits and 36 37 harms of adjunctive HBOT for treating chronic ulcers of the lower limb. Randomized controlled trials (RCTs) comparing the effect on chronic wound healing of therapeutic 38 regimens which include HBOT with those that exclude HBOT (with or without sham 39 40 therapy). Twelve trials (577 participants) were included. In people with foot ulcers due to diabetes, HBOT significantly improved the ulcers healed in the short term but not the long 41 term and the trials had various flaws in design and/or reporting that means we are not 42

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confident in the results. More trials are needed to properly evaluate HBOT in people with 1 chronic wounds; these trials must be adequately powered and designed to minimize bias. 2 Kumar et al. (2020) evaluated the efficacy of hyperbaric oxygen therapy (HBOT) as an 3 adjuvant to standard therapy for treatment of diabetic foot ulcers. A total of 54 patients 4 with diabetic foot ulcer of Wagner grade II-IV were recruited in this prospective, 5 randomized, double blind study. Patients were randomized to receive HBOT along with 6 standard therapy (group H; n = 28) or standard therapy alone (group S; n = 26). Patients 7 were given 6 sessions per week for 6 weeks and followed up for 1 year. Outcomes were 8 measured in terms of healing, and need for amputation, grafting or debridement. The 9 diabetic ulcers in 78% patients in Group H completely healed without any surgical 10 intervention while no patient in group S healed without surgical intervention. 2 patients in 11 group H required distal amputation while in Group S, three patients underwent proximal 12 amputation. Authors concluded that hyperbaric oxygen therapy is a useful adjuvant to 13 standard therapy and is a better treatment modality if combined with standard treatment 14 rather than standard treatment alone for management of diabetic foot ulcers. 15

16

26

Dauwe et al. (2014) completed a systematic review on whether hyperbaric oxygen therapy 17 works in facilitating acute wound healing given that the majority of the literature supports 18 its use for chronic wounds. A total of eight studies were found to meet criteria for 19 20 evaluation of adjunctive hyperbaric oxygen therapy in the treatment of complicated acute wounds, flaps, and grafts. Authors concluded that when combined with standard wound 21 management principles, hyperbaric oxygen therapy can augment healing in complicated 22 acute wounds. However, it is not indicated in normal wound management. Further 23 investigation is required before it can be recommended as a mainstay in adjuvant wound 24 therapy. 25

27 Wound Dressings

Application of wound dressing continues to be the standard of care for wound treatment; 28 however, the literature is inconclusive as it relates to standardized topical preparations and 29 types of dressings. Palfreyman et al. (2007) completed a Cochrane review and meta-30 analysis on dressings for venous leg ulcers. Dressing wounds is standard care. However, 31 there are different types of dressings that may improve healing. The authors reviewed all 32 33 randomized controlled trials (RCTs) that evaluated dressings applied to venous leg ulcers. Two hundred and fifty-four studies were discovered but only 42 of these fulfilled inclusion 34 criteria. Findings suggest that hydrocolloids were no more effective than simple low 35 adherent dressings used beneath compression. No other comparisons could be stated due 36 to insufficient evidence. Overall, no particular class or type of dressing appeared to be 37 better from a healing perspective than any other. According to the authors, determining 38 39 which dressing to apply should be based on local costs and preference of patient and practitioner. 40

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Roehrs et al. (2023) evaluated the effects of hyaluronic acid (and its derivatives) on the 1 healing of chronic wounds. Authors included randomized controlled trials that compared 2 the effects of hyaluronic acid (as a dressing or topical agent) with other dressings on the 3 healing of pressure, venous, arterial, or mixed-etiology ulcers and foot ulcers in people 4 with diabetes. Twelve trials (13 articles) were included in a qualitative synthesis, and four 5 trials in a quantitative analysis were combined. Overall, the included trials involved 1108 6 participants (mean age 69.60 years) presenting 178 pressure ulcers, 54 diabetic foot ulcers, 7 and 896 leg ulcers. Sex was reported for 1022 participants (57.24% female). Pressure 8 ulcers: It is uncertain whether there is a difference in complete healing; change in ulcer 9 size; or adverse events (none reported) between platelet-rich growth factor (PRGF) + 10 hyaluronic acid and PRGF because the certainty of evidence is very low (1 trial, 65 11 participants). It is also uncertain whether there is a difference in complete healing between 12 lysine hyaluronate and sodium hyaluronate because the certainty of evidence is very low. 13 Foot ulcers in people with diabetes It is uncertain whether there is a difference in time to 14 complete healing between hyaluronic acid and lyophilized collagen because the certainty 15 of evidence is very low. It is uncertain whether there is a difference in complete ulcer 16 healing or change in ulcer size between hyaluronic acid and conventional dressings because 17 the certainty of evidence is very low. Leg ulcers: Authors are uncertain whether there is a 18 difference in complete wound healing, percentage of adverse events, pain, or change in 19 20 ulcer size between hyaluronic acid + hydrocolloid and hydrocolloid because the certainty of evidence is very low (1 study, 125 participants). It is uncertain whether there is a 21 difference in change in ulcer size between hyaluronic acid and hydrocolloid because the 22 certainty of evidence is very low. Authors are uncertain whether there is a difference in 23 complete wound healing between hyaluronic acid and paraffin gauze because the certainty 24 of evidence is very low. When compared with neutral vehicle, hyaluronic acid probably 25 improves complete ulcer healing (4 studies, 526 participants; moderate-certainty 26 evidence); may slightly increase the reduction in pain from baseline (3 studies, 337 27 participants); and may slightly increase change in ulcer size, measured as mean reduction 28 from baseline to 45 days (2 studies, 190 participants). It is uncertain if hyaluronic acid 29 alters incidence of infection when compared with neutral vehicle (3 studies, 425 30 participants). Authors are uncertain whether there is a difference in change in ulcer size 31 (cm2) between hyaluronic acid and dextranomer because the certainty of evidence is very 32 33 low 1 study, 50 participants). The authors downgraded the certainty of evidence due to risk of bias or imprecision, or both, for all of the above comparisons. No trial reported health-34 related quality of life or wound recurrence. Measurement of change in ulcer size was not 35 homogeneous among studies, and missing data precluded further analysis for some 36 comparisons. Authors concluded that there is currently insufficient evidence to determine 37 the effectiveness of hyaluronic acid dressings in the healing of pressure ulcers or foot ulcers 38 39 in people with diabetes. Authors found evidence that hyaluronic acid probably improves complete ulcer healing and may slightly decrease pain and increase change in ulcer size 40 when compared with neutral vehicle. Future research into the effects of hyaluronic acid in 41

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the healing of chronic wounds should consider higher sample size and blinding to minimize
bias and improve the quality of evidence.

3

4 PRACTITIONER SCOPE AND TRAINING

5 Practitioners should practice only in the areas in which they are competent based on their 6 education, training, and experience. Levels of education, experience, and proficiency may 7 vary among individual practitioners. It is ethically and legally incumbent on a practitioner 8 to determine where they have the knowledge and skills necessary to perform such services 9 and whether the services are within their scope of practice.

10

It is best practice for the practitioner to appropriately render services to a member only if they are trained, equally skilled, and adequately competent to deliver a service compared to others trained to perform the same procedure. If the service would be most competently delivered by another health care practitioner who has more skill and training, it would be best practice to refer the member to the more expert practitioner.

16

Best practice can be defined as a clinical, scientific, or professional technique, method, or process that is typically evidence-based and consensus driven and is recognized by a majority of professionals in a particular field as more effective at delivering a particular outcome than any other practice (Joint Commission International Accreditation Standards for Hospitals, 2020).

22

Depending on the practitioner's scope of practice, training, and experience, a member's condition and/or symptoms during examination or the course of treatment may indicate the need for referral to another practitioner or even emergency care. In such cases it is prudent for the practitioner to refer the member for appropriate co-management (e.g., to their primary care physician) or if immediate emergency care is warranted, to contact 911 as appropriate. See the *Managing Medical Emergencies (CPG 159 – S)* policy for information.

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31 **REFERENCES**

Agency for Health Care Research and Quality (AHRQ). Samson D, Lefevre, F and
 Aronson N, Assessment Wound-Healing Technologies: Low-Level laser and Vacuum Assisted Closure Evidence Report/Technology, Report 111. AHRQ Publication No.
 05-E005-2. Rockville MD, December 2004

36

Agency for Health Care Research and Quality (AHRQ). Negative pressure wound therapy
 devices. AHRQ Technology assessment report July 29, 2009. Rockville MD. Retrieved
 on April 29, 2025 from https://archive.ahrq.gov/research/findings/ta/negative pressure-wound-therapy/negative-pressure-wound-therapy.pdf

American Diabetes Association. Consensus Development Conference on diabetic foot 1 wound care. Diabetes Care. 1999;22:1354-1360 2 3 American Medical Association. (current year). Current Procedural Terminology (CPT) 4 Current year (rev. ed.). Chicago: AMA 5 6 American Medical Association (current year). HCPCS Level II. American Medical 7 Association 8 9 American Medical Association. (current year). ICD-10-CM. American Medical 10 Association 11 12 APTA Guide to Physical Therapist Practice 4.0. American Physical Therapy Association. 13 Published 2023. Accessed [April 29, 2025]. https://guide.apta.org 14 15 Armstrong DG, Galiano RD, Orgill DP, et al. Multi-centre prospective randomised 16 controlled clinical trial to evaluate a bioactive split thickness skin allograft vs standard 17 of care in the treatment of diabetic foot ulcers. International Wound Journal. 18 2022;19(4):932-944 19 20 Armstrong D, Orgill D, Galiano R, et al. A purified reconstituted bilayer matrix shows 21 improved outcomes in treatment of non-healing diabetic foot ulcers when compared to 22 the standard of care: Final results and analysis of a prospective, randomized, controlled, 23 multi-centre clinical trial. International Wound Journal. 2024;21(4):e14882 24 25 Avendaño-Coy J, López-Muñoz P, Serrano-Muñoz D, Comino-Suárez N, Avendaño-26 López C, Martin-Espinosa N. Electrical microcurrent stimulation therapy for wound 27 28 healing: A meta-analysis of randomized clinical trials [published online ahead of print, 2021 Dec 4]. J Tissue Viability. 2021;S0965-206X(21)00132-7 29 30 Aziz Z, Cullum N, Flemming K. Electromagnetic therapy for treating venous leg ulcers. 31 Cochrane Database of Systematic Reviews 2013, Issue 2. Art. No.: CD002933. DOI: 32 10.1002/14651858.CD002933.pub5 33 34 Banerjee J, Lasiter A, Nherera L. Systematic review of cellular, acellular and matrix-like 35 products (CAMPs) and indirect treatment comparison between cellular/acellular and 36 37 amniotic/non-amniotic grafts in the management of diabetic foot ulcers. Advances in Wound Care. 2024(ja) 38 39 40 Baranski S, Ayello EA Wound Care Essentials: Practice Principles 2nd Edition Philadelphia PA Lippincott Williams & Wilkins 2008 41

Page 59 of 75

1	Barnes R, Shahin Y, Gohil R, Chetter I. Electrical stimulation vs. standard care for chronic
2	ulcer healing: a systematic review and meta-analysis of randomised controlled trials.
3 4	Eur J Clin Invest. 2014 Apr;44(4) :429-40
5	Barret JP, Dziewulski P, Ramzy PI, et al. Biobrane versus 1% silver sulfadiazine in second-
6	degree pediatric burns. Plast Reconstr Surg. 2000;105(1):62-65
7	
8	Beckmann KH, Meyer-Hamme G, Schröder S. Low level laser therapy for the treatment of
9	diabetic foot ulcers: a critical survey. Evid Based Complement Alternat Med.
10	2014;2014:626127
11	
12	Beheshti A, Shafigh Y, Parsa H, Zangivand AA. Comparison of high-frequency and MIST
13	ultrasound therapy for the healing of venous leg ulcers. Adv Clin Exp Med.2014 Nov-
14	Dec;23(6):969-75
15	
16	Bello Y, Phillips T, Recent Advances in Wound Healing. JAMA. 2000;283(6):716-718
17	
18	Black, J., Baharestani, M., Cuddigan, J., Dorner, B., Edsberg, L., Langemo, D. National
19	Pressure Ulcer Advisory, P. (2007). National Pressure Ulcer Advisory Panel's updated
20	pressure ulcer staging system. Dermatol Nurs, 19(4), 343-349; quiz 350
21	
22	Bluestein D, Javaheri A. Pressure ulcers: prevention, evaluation, and management. Am
23	Fam Physician. 2008;78(10):1186-1194
24	
25	Blume PA, Walters J, Payne W, Ayala J, Lantis J. Comparison of negative pressure wound
26	therapy using vacuum-assisted closure with advanced moist wound therapy in the
27	treatment of diabetic foot ulcers: a multicenter randomized controlled trial. Diabetes
28	Care. 2008 Apr;31(4):631-6. Epub 2007 Dec 27
29	
30	Brem H, Kirsner RS, Falanga V. "Protocol for the successful treatment of venous ulcers".
31	(2004) Am. J. Surg. 188 (1A Suppl): 1–8
32	
33	Brigido SA, Boc SF, Lopez RC. Effective management of major lower extremity wounds
34	using an acellular regenerative tissue matrix: A pilot study. Orthopedics. 2004;27(1
35	Suppl):s145-s149
36	
37	Brölmann FE, Ubbink DT, Nelson EA, Munte K, van der Horst CM, Vermeulen H.
38	Evidence-based decisions for local and systemic wound care. Br J Surg. 2012
39 40	Sep;99(9):1172-83
40	Durna II - Diastruali SI Diastia aurgany lawar antromity. In: Townsond, Schister Touthask
41 42	Burns JL, Blackwell SJ. Plastic surgery lower extremity. In: Townsend: Sabiston Textbook of Surgery. 19 th ed. Philadelphia, PA: Saunders Elsevier, 2012. Ch. 64 and 65

Page 60 of 75

1	Cabeza de Vaca, F. G., Macias, A. E., Ramirez, W. A., Munoz, J. M., Alvarez, J. A.,
2	Mosqueda, J. L., Sifuentes-Osornio, J. (2010). Salvaging diabetic foot through
3	debridement, pressure alleviation, metabolic control, and antibiotics Cabeza de Vaca
4	et al. Salvaging diabetic foot. Wound Repair & Regeneration, 18(6), 567-571. Doi:
5	10.1111/j.1524-475X.2010.00621.x
6	
7	Carpenter S F, A, Bahadur D, Estapa A, Bahm J. Evaluating the Number of Cellular or
8	Tissue-Based Product Applications Required for Treating Diabetic Foot Ulcers and
9	Venous Leg Ulcers in Non-Hospital Outpatient Department Settings. WOUNDS.
10	2024;35(8)
11	
12	Centers for Medicare and Medicaid. Decision memo for electrostimulation for wounds
13	(CAG-00068R) In: Medicare Coverage Database. Baltimore, MD: December 2003
14	
15	Centers for Medicare and Medicaid. Local Coverage Determination for Debridement
16	Services (L33614). Retrieved on April 29, 2025 from https://www.cms.gov/medicare-
17	coverage-database/details/lcd-
18	details.aspx?lcdid=33614&ver=26&bc=CAAAAAAAAAAAA
19	
20	Centers for Medicare and Medicaid. Local Coverage Determination (LCD): Outpatient
21	Physical and Occupational Therapy Services (L33631). Retrieved on April 29, 2025
22	from https://www.cms.gov/medicare-coverage-database/details/lcd-
23	details.aspx?lcdid=33631&ver=51&bc=CAAAAAAAAAAAA
24	Contara for Madianra and Madianid Lagal Coverage Determination (LCD): Surgical
25 26	Centers for Medicare and Medicaid. Local Coverage Determination (LCD): Surgical Dressings (L33831). Retrieved on April 29, 2025 from https://www.cms.gov/medicare-
26 27	coverage-database/view/lcd.aspx?lcdid=33831&ver=40&bc=CAAAAAAAAAA
27 28	coverage-ualabase/view/icu.aspx?iculu=55851&ve1=40&bc=CAAAAAAAAAAAAA
28 29	Centers for Medicare and Medicaid Services. Local Coverage Determination (LCD):
30	Wound Care (L35125). Retrieved on April 29, 2025 from
31	https://www.cms.gov/medicare-coverage-database/details/lcd-
32	details.aspx?lcdid=35125&ver=76&bc=CAAAAAAAAAA
33	
34	Centers for Medicare and Medicaid. National Coverage Determination (NCD) for
35	Hyperbaric Oxygen Therapy (20.29) Retrieved on April 29, 2025 from
36	https://www.cms.gov/medicare-coverage-database/details/ncd-
37	details.aspx?NCDId=12&ncdver=4&DocID=20.29&SearchType=Advanced&bc=IA
38	AAABAAAAA&
39	
40	Centers for Medicare and Medicaid. National Coverage Determination (NCD) for
41	Treatment of Decubitus Ulcers (270.4). Retrieved on April 29, 2025 from
42	https://www.cms.gov/medicare-coverage-database/details/ncd-

Page 61 of 75

1	details.aspx?NCDId=47&ncdver=1&DocID=270.4&ncd_id=270.4&ncd_version=1&
2	basket=ncd%25253A270%25252E4%25253A1%25253ATreatment+of+Decubitus+
3	Ulcers&bc=gAAAAAgAAAAAAA%3D%3D
4	
5	Centers for Medicare and Medicaid Services. National Coverage Determination (NCD) for
6	Electrical Stimulation (ES) and Electromagnetic Therapy for the Treatment of Wounds
7	(270.1). Retrieved on April 29, 2025 from https://www.cms.gov/medicare-coverage-
8	database/details/ncd-
9	details.aspx?NCDId=131&ncdver=3&bc=AAAAgAAAAAAA&
10	
11	Centers for Medicare and Medicaid Services. Local Coverage Determination (LCD): Skin
12	Substitute Grafts/Cellular and Tissue-Based Products for the Treatment of Diabetic
13	Foot Ulcers and Venous Leg Ulcers (35041). Retrieved on April 28, 2025 from
14	https://www.cms.gov/medicare-coverage-
15	database/view/lcd.aspx?LCDId=35041#:~:text=This%20Local%20Coverage%20Det
16	ermination%20(LCD,failed%20established%20methods%20of%20healing
17	
18	Chen C, Hou WH, Chan ES, Yeh ML, Lo HL. Phototherapy for treating pressure ulcers.
19	Cochrane Database Syst Rev. 2014 Jul 11;(7):CD009224
20	
21	Chen H, Xiao T, Zhang L, Liu N, Liang X, Li T, Wang J, Peng Y, Liu Y, Xu J. Effect of
22	ultrasound-supported wound debridement in subjects with diabetic foot ulcers: A meta-
23	analysis. Int Wound J. 2023 Sep;20(7):2618-2625
24	
25	Chen H, Yu Z, Liu N, Huang J, Liang X, Liang X, Liang M, Li M, Ni J. The efficacy of
26	low-frequency ultrasound as an added treatment for chronic wounds: A meta-analysis.
27	Int Wound J. 2023 Feb;20(2):448-457
28	
29	Chen P, Vilorio NC, Dhatariya K, Jeffcoate W, Lobmann R, McIntosh C, Piaggesi A,
30	Steinberg J, Vas P, Viswanathan V, Wu S, Game F. Guidelines on interventions to
31	enhance healing of foot ulcers in people with diabetes (IWGDF 2023 update). Diabetes
32	Metab Res Rev. 2024 Mar;40(3):e3644
33	
34	Chen P, Vilorio NC, Dhatariya K, Jeffcoate W, Lobmann R, McIntosh C, Piaggesi A,
35	Steinberg J, Vas P, Viswanathan V, Wu S, Game F. Effectiveness of interventions to
36	enhance healing of chronic foot ulcers in diabetes: A systematic review. Diabetes
37	Metab Res Rev. 2024 Mar;40(3):e3786
38	
39	Chen AC, Lu Y, Hsieh CY, Chen YS, Chang KC, Chang DH. Advanced Biomaterials and
40	Topical Medications for Treating Diabetic Foot Ulcers: A Systematic Review and
41	Network Meta-Analysis. Adv Wound Care (New Rochelle). 2024 Feb;13(2):97-113

1 2	Chen HL, Chung JWY, Yan VCM, Wong TKS. Polylactic Acid-Based Biomaterials in Wound Healing: A Systematic Review. Adv Skin Wound Care. 2023 Sep 1;36(9):1-8
3	Chen V Dy D Ly C A mete enclusion exemined the effect of enclosed reconnected
4 5	Chen Y, Du P, Lv G. A meta-analysis examined the effect of oxidised regenerated cellulose/collagen dressing on the management of chronic skin wounds. Int Wound J.
6	2023 May;20(5):1544-1551
7	
8	Cooper P. A review of different wound types and their principles of management in Wound
9	Healing: A systematic approach to advanced wound healing and management. 2005
10	Cromwell Press UK
11	
12	Cullum N, Nelson EA, Fletcher A, Sheldon T. Compression for venous leg ulcers.
13	Cochrane Database Syst Rev 2001;(2):CD000265. ECRI Institute. Hotline
14	
15	Damineni U, Divity S, Gundapaneni SRC, Burri RG, Vadde T. Clinical Outcomes of
16	Hyperbaric Oxygen Therapy for Diabetic Foot Ulcers: A Systematic Review. Cureus.
17	2025;17(2):e78655. Published 2025 Feb 6. doi:10.7759/cureus.78655
18	
19	Damoiseaux, J. (2013). Bullous skin diseases: classical types of autoimmune diseases.
20	Scientifica (Cairo), 2013, 457982
21	
22	Dauwe PB, Pulikkottil BJ, Lavery L, Stuzin JM, Rohrich RJ. Does hyperbaric oxygen
23 24	therapy work in facilitating acute wound healing: a systematic review. Plast Reconstr Surg. 2014 Feb;133(2):208e-15e
25	Suig. 2011100,135(2).2000 150
26	David G. Armstrong DPO, Robert D. Galiano, Paul M. Glat, Jarrod P. Kaufman, Marissa
20 27	J. Carter, Lawrence A. DiDomenico, Charles M. Zelen, Paul M. Glat, Jarrod P.
28	Kaufman, Marissa J. Carter, Lawrence A. DiDomenico, Charles M. Zelen. Use of a
29	purified reconstituted bilayer matrix in the management of chronic diabetic foot ulcers
30	improves patient outcomes vs standard of care: Results of a prospective randomised
31	controlled multi-centre clinical trial. Int Wound J. 2022;19(5):1197-1209
32	
33	Debridement Procedures for Managing Diabetic Foot Ulcers: A Review of Clinical
34	Effectiveness, Cost-effectiveness, and Guidelines. (2014). Ottawa ON: 2014 Canadian
35	Agency for Drugs and Technologies in Health
36	
37	Dryden, M. S. (2010). Complicated skin and soft tissue infection. J Antimicrob Chemother,
38	65 Suppl 3, iii35-44
39	
40	Dumville JC, Deshpande S, O'Meara S, Speak K. Foam dressings for healing diabetic foot

41 ulcers. Cochrane Database of Systematic Reviews 2013, Issue 6. Art. No.: CD009111

1	Dumville JC, Hinchliffe RJ, Cullum N, Game F, Stubbs N, Sweeting M, Peinemann F.
2	Negative pressure wound therapy for treating foot wounds in people with diabetes
3	mellitus. Cochrane Database of Systematic Reviews 2013, Issue 10. Art. No.:
4	CD010318
5	
6	Dumville JC, Keogh SJ, Liu Z, Stubbs N, Walker RM, Fortnam M. Alginate dressings for
7	treating pressure ulcers. Cochrane Database Syst Rev. 2015 May 21;5:CD011277
8	
9	Dumville JC, O'Meara S, Deshpande S, Speak K. Alginate dressings for healing diabetic
10	foot ulcers. Cochrane Database of Systematic Reviews 2013, Issue 6. Art. No.: CD000110
11	CD009110
12	Dumville JC, O'Meara S, Deshpande S, Speak K. Hydrogel dressings for healing diabetic
13	foot ulcers. Cochrane Database of Systematic Reviews 2013, Issue 7. Art. No.:
14	CD009101
15 16	CD009101
17	Dumville JC, Webster J, Evans D, Land L. Negative pressure wound therapy for treating
17	pressure ulcers. Cochrane Database Syst Rev. 2015 May 21;5:CD011334
19	pressure dicers. Coefficie Daubase Byst Rev. 2015 May 21,5. CD011554
20	Dumville JC, Stubbs N, Keogh SJ, Walker RM, Liu Z. Hydrogel dressings for treating
20	pressure ulcers. Cochrane Database Syst Rev. 2015 Feb 17;2:CD011226
22	1
23	Ehrenreich M, Ruszczak Z. Tissue-engineered temporary wound coverings. Important
24	options for the clinician. Acta Dermatovenerol Alp Panonica Adriat. 2006;15(1):5-13
25	
26	Eleftheriadou I, Samakidou G, Tentolouris A, Papanas N, Tentolouris N.
27	Nonpharmacological Management of Diabetic Foot Ulcers: An Update. Int J Low
28	Extrem Wounds. 2021;20(3):188-197
29	
30	Eneroth M, van Houtum WH, The value of debridement and Vacuum-Assisted Closure
31	(V.A.C.) Therapy in diabetic foot ulcers. Diabetes Metab Res Rev, 2008;May-Jun;24(
32	Suppl 1):S76-80
33	
34	Enoch S and Price P (2004) Cellular, molecular and biochemical differences in the
35	pathophysiology of healing between acute wounds, chronic wounds and wounds in the
36	aged. Retrieved on April 29, 2025 from
37	http://www.worldwidewounds.com/2004/august/Enoch/Pathophysiology-Of-
38	Healing.html
39	
40	Enwemeka CS. Intricacies of dose in laser phototherapy for tissue repair and pain relief,
41	Photomed Laser Surg. 2009 Jun; 27(3):387-93

Page 64 of 75

1 2 3	Enwemeka CS, Parker JC, Dowdy DS, Harkness EE, Sanford LE, Woodruff LD. The efficacy of low-power lasers in tissue repair and pain control: a meta-analysis study. Photomed Laser Surg. 2004 Aug; 22(4):323-9
4 5	ERCI Institute. Electrical Stimulation and Electromagnetic Therapy (AHRQ). Plymouth
5 6	Meeting PA: ERCI Institute Health Technology Assessment Information Service, Dec
7	2010. Available at: http://www.ecri.org
8	
9	ERCI Institute. Negative Pressure Wound Therapy Devices (AHRQ). Rockville, MD:
10	ERCI Institute Health Technology Assessment Information Service, July 2009.
11	Available at: http://www.ecri.org
12	
13	Eskes A, Vermeulen H, Lucas C, Ubbink DT. Hyperbaric oxygen therapy for treating acute
14	surgical and traumatic wounds. Cochrane Database Syst Rev. 2013 Dec
15	16;12:CD008059
16	
17	Faglia, E., Clerici, G., Caminiti, M., Quarantiello, A., Gino, M., & Morabito, A. (2006).
18	The role of early surgical debridement and revascularization in patients with diabetes
19	and deep foot space abscess: retrospective review of 106 patients with diabetes. Journal
20	of Foot & Ankle Surgery, 45(4), 220-226
21	Easter IA Klath IC Container CD Change Demol III an II align Enhanced with
22	Feedar JA, Kloth LC, Gentzkow, GD. Chronic Dermal Ulcer Healing Enhanced with Monophasia Pulsad Elastrical Stimulation Phys. Ther 1001:71:630-640
23	Monophasic Pulsed Electrical Stimulation Phys Ther 1991;71:639-649
24 25	Fernandez-Chimeno M, Houghton P, Holey L. Electrical stimulation for chronic wounds
25 26	(Cochrane Review). Intervention Protocol. In: The Cochrane Library, Issue I, 2004.
20 27	Oxford: Update Software
28	Onlord. Optimie
29	Fisher, T. K., Scimeca, C. L., Bharara, M., Mills, J. L., Sr., & Armstrong, D. G. (2010). A
30	step-wise approach for surgical management of diabetic foot infections. J Vasc Surg,
31	52(3 Suppl), 72S-75S
32	
33	Fletcher A, Cullum N, Sheldon TA. A systematic review of compression treatment for
34	venous leg ulcers. BMJ 1997;315:576-80
35	
36	Flanagan, M. (1997) A practical framework for wound assessment 2: methods. British
37	Journal of Nursing: 6; 6, $8 - 11$
38	
39	Flanagan, M. (1994) Assessment Criteria. Nursing Times: 90; 35, 76 – 86

1 2 3 4	Foster K, Greenhalgh D, Gamelli R et al; FS 4IU VH S/D Clinical Study Group. Efficacy and safety of a fibrin sealant for adherence of autologous skin grafts to burn wounds: Results of a phase 3 clinical study. J Burn Care Res. 2008;29(2):293-303
5 6 7 8 9	Game FL, Hinchliffe RJ, Apelqvist J, Armstrong DG, Bakker K, Hartemann A, Löndahl M, Price PE, Jeffcoate WJ. A systematic review of interventions to enhance the healing of chronic ulcers of the foot in diabetes. Diabetes Metab Res Rev. 2012 Feb;28 Suppl 1:119-41
10 11 12 13	Gibbons GW, Orgill DP, Serena TE, et al. A prospective, randomized, controlled trial comparing the effects of noncontact, low-frequency ultrasound to standard care in healing venous leg ulcers. Ostomy Wound Manage. 2015;61(1):16-29
14 15 16 17	Gillespie BM, Thalib L, Ellwood D, et al. Effect of negative-pressure wound therapy on wound complications in obese women after caesarean birth: a systematic review and meta-analysis. BJOG. 2022;129(2):196-207
18 19 20 21	Greer N, Foman NA, MacDonald R, et al. Advanced wound care therapies for nonhealing diabetic, venous, and arterial ulcers: A systematic review. Ann Intern Med. 2013;159(8):532-542
21 22 23	Guo S, DiPietro LA. Factors Affecting Wound Healing, J Dent Res 2010;89(3):219-229
24 25 26 27	Guo X, Mu D, Gao F. Efficacy and safety of acellular dermal matrix in diabetic foot ulcer treatment: a systematic review and meta-analysis. International Journal of Surgery. 2017;40:1-7
28 29 30 31 32	Gurtner G, Garcia, AD, Bakewell, K, Alarcon, JB. A retrospective matched-cohort study of 3994 lower extremity wounds of multiple etiologies across 644 institutions comparing a bioactive human skin allograft, TheraSkin, plus standard of care, to standard of care alone. International Wound Journal. 2020;17(1):55-64
33 34 35 36	Harding K, Sumner M, Cardinal M. A prospective, multicentre, randomised controlled study of human fibroblast-derived dermal substitute (Dermagraft) in patients with venous leg ulcers. Int Wound J. 2013;10(2):132-137
37 38	Haycocks, S., & Chadwick, P. (2012). Debridement of diabetic foot wounds. <i>Nursing Standard</i> , 26(24), 51-51-52, 54, 56 passim
 39 40 41 42 	Ho C, Bogie K. Pressure Ulcers. In: Frontera W, Silver J, Rizzo, T: Essentials of Physical Medicine and Rehabilitation: musculoskeletal disorders, pain and rehabilitation. 2 nd ed. Philadelphia, PA: Saunders Elsevier, 2008. Ch. 140

Page 66 of 75

1 2	Holloway GA, Arterial ulcers: assessment and diagnosis. Ostomy Wound Manage. 1996 Apr;42(3):46-8, 50-1
3	
4	Hopf HW, Ueno C, Aslam R, Burnand K, Fife C, and Grant L. et al. Guidelines for the
5	treatment of arterial insufficiency ulcers. Wound Repair Regen. 2006 Nov-
6	Dec;14(6):693-710
7	
8	Hopkins JT, McLoda TA, Seegmiller JG, Baxter D, Low-Level Laser Therapy Facilitates
9	Superficial Wound healing in Humans: a triple-blind, sham-controlled study. J Athletic
10	Training 2004:339(3):223-229
11	
12	Horn C, Fierro A, Lantis Ii JC. Use of negative pressure wound therapy for the treatment
13	of venous leg ulcers. Wounds. 2023 Jun;35(6):117-125
14	
15	Houghton PE, Kincaid CB, Lovell M, Campbell KE, Keast DH, Woodbury G, and Harris
16	K. Effect of Electrical Stimulation on Chronic Leg Ulcer Size and Appearance. Phys
17	Ther,2003;83:17-28
18	
19	Irion G Comprehensive Wound Management 2nd Edition, Thorofare, NJ: Slack Inc., 2010
20	
21	Joint Commission International. (2020). Joint Commission International Accreditation
22	Standards for Hospitals (7th ed.): Joint Commission Resources
23	
24	Joseph E, Hamori CA, Bergman S, Roaf E, Swann NF, Anastasi GW. A prospective,
25	randomized trial of vacuum-assisted closure versus standard therapy of chronic non-
26	healing wounds. Wounds 2000;12(3):60-7
27	
28	Kane, D. (2014). Surgical Management of Pressure Ulcers. In M. D. D. R. Thomas & M.
29	D. G. A. Compton (Eds.), Pressure Ulcers in the Aging Population (Vol. 1, pp. 99-
30	126): Humana Press
31	
32	Kaviani A, et al. A randomized clinical trial on the effect of low-level laser therapy on
33	chronic diabetic foot wound healing: a preliminary report. Photomedicine and Laser
34	Surgery 2011;29(2):109-14
35	
36	Kloth LC, McCulloch JM, Feedar MA Wound Healing: Alternatives in Management 2nd
37	Edition, Philadelphia PA: FA Davis 1995
38	
39	Kneisel, A., & Hertl, M. (2011). Autoimmune bullous skin diseases. Part 1: Clinical
40	manifestations. J Dtsch Dermatol Ges, 9(10), 844-856; quiz 857. doi: 10.1111/j.1610-
41	0387.2011.07793.x

1 2 3	Kotronis G, Vas PRJ. Ultrasound Devices to Treat Chronic Wounds: The Current Level of Evidence. Int J Low Extrem Wounds. 2020 Dec;19(4):341-349. doi: 10.1177/1534734620946660
3 4	10.11///1554/54020940000
5 6	Kranke P, Bennett MH, Martyn-St James M, Schnabel A, Debus SE, Weibel S. Hyperbaric oxygen therapy for chronic wounds. Cochrane Database Syst Rev. 2015 Jun
7	24;6:CD004123
8	
9	Kujath P, Michelsen A. Wounds from Physiology to Wound Dressing. Dtsch Arztebl Int
10	2008; 105(13):239-48
11 12	Kumar A, Shukla U, Prabhakar T, Srivastava D. Hyperbaric oxygen therapy as an adjuvant
12	to standard therapy in the treatment of diabetic foot ulcers. J Anaesthesiol Clin
13	Pharmacol. 2020 Apr-Jun;36(2):213-218. doi: 10.4103/joacp.JOACP_94_19. Epub
15	2020 Jun 15
16	
17	Kumbhar S, Bhatia M. Advancements and best practices in diabetic foot Care: A
18	comprehensive review of global progress. Diabetes Res Clin Pract. 2024;217:111845.
19	doi:10.1016/j.diabres.2024.111845
20	
21	Kwee E, Borgdorff M, Schepers T, et al. Adjunctive hyperbaric oxygen therapy in the
22 23	management of severe lower limb soft tissue injuries: a systematic review. Eur J Trauma Emerg Surg. 2024;50(3):1093-1100. doi:10.1007/s00068-023-02426-2
24 25	Lala D, Spaulding SJ, Burke SM, Houghton PE. Electrical stimulation therapy for the
26 27	treatment of pressure ulcers in individuals with spinal cord injury: a systematic review and meta-analysis. Int Wound J. 2015 Apr 13
28	
29	Lalieu RC, Akkerman I, van Hulst RA. Hyperbaric Oxygen Therapy for Venous Leg
30	Ulcers: A 6 Year Retrospective Study of Results of a Single Center. Front Med
31	(Lausanne). 2021;8:671678. Published 2021 Jul 28
32	Lalieu RC, Bol Raap RD, Smit C, Dubois EFL, van Hulst RA. Hyperbaric Oxygen Therapy
33 34	for Nonhealing Wounds-A Long-term Retrospective Cohort Study. Adv Skin Wound
35	Care. 2023 Jun 1;36(6):304-310
36	
37	Langer A, Rogowski W. Systematic review of economic evaluations of human cell-derived
38	wound care products for the treatment of venous leg and diabetic foot ulcers. BMC
39	Health Serv Res. 2009;9:115
40	
41	Lantis J, Lullove, Eric J, Liden, Brock, McEneaney, Patrick, Raphael, Allen, Klein, Robert,
42	Winters, Christopher, Huynh, Ruby N. Final efficacy and cost analysis of a fish skin

Page 68 of 75

1 2 3 4	graft vs standard of care in the management of chronic diabetic foot ulcers: a prospective, multicenter, randomized controlled clinical trial. Wounds: a Compendium of Clinical Research and Practice. 2023;35(4):71-79
5 6 7 8	Lazarus GS, Cooper DM, Knighton DR, Margolis DJ, and Pecoraro RE et al. Definitions and Guidelines of Assessment of Wounds and Evaluation of Healing, Arch Dermatol. Apr 1994;130:489-493
9 10 11 12	Lebrun, E., Tomic-Canic, M., & Kirsner, R. S. (2010). The role of surgical debridement in healing of diabetic foot ulcers. <i>Wound Repair Regen</i> , <i>18</i> (5), 433-438. doi: 10.1111/j.1524-475X.2010.00619.x
13 14 15 16	Leclere FM, Puechguiral IR, Rotteleur G, Thomas P, Mordon SR. A prospective randomized study of 980 nm diode laser-assisted venous ulcer healing on 34 patients. Wound Repair and Regeneration 2010;18(6):580-5
10 17 18 19 20	Levine, S. M., Sinno, S., Levine, J. P., & Saadeh, P. B. (2013). Current thoughts for the prevention and treatment of pressure ulcers: using the evidence to determine fact or fiction. <i>Ann Surg</i> , 257(4), 603-608
21 22 23 24 25	Lipsky, B. A., Berendt, A. R., Cornia, P. B., Pile, J. C., Peters, E. J., Armstrong, D. G. Infectious Diseases Society of, A. (2012). 2012 Infectious Diseases Society of America clinical practice guideline for the diagnosis and treatment of diabetic foot infections. <i>Clin Infect Dis</i> , 54(12), e132-173
26 27 28 29 30	 Liu, C., Bayer, A., Cosgrove, S. E., Daum, R. S., Fridkin, S. K., Gorwitz, R. J., Chambers, H. F. (2011). Clinical Practice Guidelines by the Infectious Diseases Society of America for the Treatment of Methicillin-Resistant Staphylococcus Aureus Infections in Adults and Children. <i>Clinical Infectious Diseases</i>
31 32 33 34	Lucas C, Coenen CHM, De Haan RJ. The effect of low level laser therapy (LLLT) on stage III decubitus ulcers (pressure sores); A prospective randomised single blind, multicentre pilot study. Lasers Med Sci 2000; 15(2):94-100
35 36 37 38	Lucas C, van Gemert MJ, de Haan RJ. Efficacy of low-level laser therapy in the management of stage III decubitus ulcers: a prospective, observer-blinded multicentre randomised clinical trial. Lasers Med Sci 2003; 18(2):72-7
38 39 40	Lundeberg T, Malm M. Low-power HeNe laser treatment of venous leg ulcers. Ann Plast Surg 1991; 27(6):537-9

Page 69 of 75

1 2 3	Machado, N. O. (2011). Necrotizing fasciitis: The importance of early diagnosis, prompt surgical debridement and adjuvant therapy. <i>North Am J Med Sci, 3</i> , 107-118
3 4 5	Malm M, Lundeberg T. Effect of low power gallium arsenide laser on healing of venous ulcers. Scand J Plast Reconstr Surg Hand Surg 1991; 25(3):249-51
6	aloois. Sound \$ 1 hast Reconstr Surg Hund Surg 1991; 25(5):219 51
7	Mathieu D, Linke J-C, Wattel F (2006). Non-healing wounds. In: Handbook on hyperbaric
8	medicine, Mathieu DE, editor. Netherlands: Springer, pp.401-427
9	
10	Menke NB, Ward KR, Witten TM, Bonchev DG, Diegelmann RF (2007). Impaired wound
11	healing. Clin Dermatol 25:19-25
12	
13	Mishra A, Kushare A, Gupta MN, Ambre P. Advanced Dressings for Chronic Wound
14	Management. ACS Appl Bio Mater. 2024;7(5):2660-2676.
15	doi:10.1021/acsabm.4c00138
16	
17	Moore ZEH, Cowman S. Wound cleansing for pressure ulcers. Cochrane Database of
18	Systematic Reviews 2013, Issue 3. Art. No.: CD004983
19	
20	National Institute for Health and Clinical Excellence (NICE). Overview: Pressure ulcers:
21	Prevention and management: Guidance. (2014). Retrieved April 29, 2025 from
22	http://www.nice.org.uk/guidance/cg179
23	
24	National Pressure Ulcer Advisory Panel. 2014. Treatment of Pressure Ulcers. In:
25	Prevention and treatment of pressure ulcers: clinical practice guideline. Retrieved on
26	April 29, 2025 from https://npiap.com/page/2014Guidelines
27	
28	Norman G, Goh EL, Dumville JC, Shi C, Liu Z, Chiverton L, Stankiewicz M, Reid A.
29	Negative pressure wound therapy for surgical wounds healing by primary closure.
30	Cochrane Database Syst Rev. 2020 Jun 15;6(6):CD009261. doi:
31	10.1002/14651858.CD009261.pub6
32	
33	Nussbaum EL, Biemann I, Mustard B. Comparison of ultrasound/ultraviolet-C and laser
34	for treatment of pressure ulcers in patients with spinal cord injury. Phys Ther 1994;
35	74(9):812-23
36	
37	Ogbeide OA, Okeleke SI, Okorie JC, et al. Evolving Trends in the Management of Diabetic
38	Foot Ulcers: A Narrative Review. Cureus. 2024;16(7):e65095. Published 2024 Jul 22.
39	doi:10.7759/cureus.65095
40	
41	O'Meara S, Martyn-St James M. Alginate dressings for venous leg ulcers. Cochrane
42	Database of Systematic Reviews 2013, Issue 4. Art. No.: CD010182

Page 70 of 75

1 2 3	O'Meara S, Martyn-St James M. Foam dressings for venous leg ulcers. Cochrane Database of Systematic Reviews 2013, Issue 5. Art. No.: CD009907
4 5	O'Meara S, Cullum N, Nelson EA, Dumville JC. Compression for venous leg ulcers. Cochrane Database Syst Rev. 2012 Nov 14;11:CD000265
6	
7 8	Onderková A, Butler PEM, Kalavrezos N. The efficacy of negative-pressure wound therapy for head and neck wounds: A systematic review and update. Head Neck. 2023
9	Dec;45(12):3168-3179
10	
11	Ontario Health (Quality) . Skin Substitutes for Adults With Diabetic Foot Ulcers and
12	Venous Leg Ulcers: A Health Technology Assessment. Ont Health Technol Assess Ser.
13 14	2021;21(7):1-165. Published 2021 Jun 4
14	Olyaie M, Rad FS, Elahifar MA, Garkaz A, Mahsa G. High-frequency and noncontact low-
	frequency ultrasound therapy for venous leg ulcer treatment: a randomized, controlled
16	study. Ostomy Wound Manage. 2013 Aug;59(8):14-20
17	study. Ostonny wound Manage. 2015 Aug, 59(8).14-20
18	Quahas N. Dhilling TI I as ulases Curr Drohl Domatal 1005 7:100 142
19 20	Ouahes N., Phillips TJ Leg ulcers Curr Probl Dematol. 1995-7:109-142
20	Dedule WW Demonsthern & Cohen DC Dessen C Armstrong DC Componetius
21	Padula WV, Ramanathan S, Cohen BG, Rogan G, Armstrong DG. Comparative
22	Effectiveness of Placental Allografts in the Treatment of Diabetic Lower Extremity
23	Ulcers and Venous Leg Ulcers in US Medicare Beneficiaries: a retrospective
24	observational cohort study using real-world evidence. Advances in Wound Care. 2024
25	
26	Palfreyman SJ, Lochiel R, Michaels JA. A systematic review of compression therapy for
27	venous leg ulcers. Vasc Med 1998;3:301-13
28	Defension CL Malan DA Mishala IA Describes for supervised has also setting
29	Palfreyman SJ, Nelson EA, Michaels JA. Dressings for venous leg ulcers: systematic
30	review and meta-analysis. BMJ 2007 Aug 4:335(7613):244
31	Dedrami NE Meiken S. Le Scale C. Negative Pressure Wound Thereasy in Dedictric Dum
32	Pedrazzi NE, Naiken S, La Scala G. Negative Pressure Wound Therapy in Pediatric Burn Patiente: A Systematic Paylor Adv Wound Core (New Pachella), 2021;10(5):270
33	Patients: A Systematic Review. Adv Wound Care (New Rochelle). 2021;10(5):270-280
34	280
35	Discus C. Commercial I. Challen III. et al. Discussion and altimetric franches from the
36	Pham C, Greenwood J, Cleland H, et al. Bioengineered skin substitutes for the management
37	of burns: A systematic review. Burns. 2007;33(8):946-957
38	Deb Etznetziek M. & Luching Harking I. M. (2012) Dullaus Disesse of Disketes
39 40	Poh-Fitzpatrick, M., & Junkins-Hopkins, J. M. (2013). Bullous Disease of Diabetes Treatment & Management. <i>Drugs and Diseases</i> . Retrieved from
41	http://emedicine.medscape.com/article/1062235-treatment

Page 71 of 75

Polak A, Franek A, Blaszczak E, et al. A prospective, randomized, controlled, clinical
 study to evaluate the efficacy of high-frequency ultrasound in the treatment of Stage II
 and Stage III pressure ulcers in geriatric patients. Ostomy Wound Manage.
 2014;60(8):16-28

- Posten W, Wrone DA, Dover JS, Arndt KA, Silapunt S, Alam M. Low-level laser therapy
 for wound healing: mechanism and efficacy. Dermatol Surg. 2005 Mar;31(3):334-40
- Purdue GF, Hunt JL, Still JM Jr, et al. A multicenter clinical trial of a biosynthetic skin
 replacement, Dermagraft-TC, compared with cryopreserved human cadaver skin for
 temporary coverage of excised burn wounds. J Burn Care Rehabil. 1997;18(1 Pt 1):52 57
- Putri IL, Adzalika LB, Pramanasari R, Wungu CDK. Negative pressure wound therapy
 versus conventional wound care in cancer surgical wounds: A meta-analysis of
 observational studies and randomised controlled trials. Int Wound J.
 2022;10.1111/iwj.13756
- Rastogi A, Bhansali A, Ramachandran S. Efficacy and Safety of Low-Frequency,
 Noncontact Airborne Ultrasound Therapy (Glybetac) For Neuropathic Diabetic Foot
 Ulcers: A Randomized, Double-Blind, Sham-Control Study. Int J Low Extrem
 Wounds. 2019 Mar;18(1):81-88
- Recio AC, Felter CE, Schneider AC, McDonald JW, High-voltage electrical stimulation
 for the management of stage III and IV pressure ulcers among adults with spinal cord
 injury: demonstration of its utility for recalcitrant wounds below the level of injury. J
 Spinal Cord Med. 2012 Jan;35(1):58-63
- 28

32

36

8

13

18

23

- Ricco, J. B., Thanh Phong, L., Schneider, F., Illuminati, G., Belmonte, R., Valagier, A., &
 Regnault De La Mothe, G. (2013). The diabetic foot: a review. *J Cardiovasc Surg*(*Torino*), 54(6), 755-762
- Roehrs H, Stocco JG, Pott F, Blanc G, Meier MJ, Dias FA. Dressings and topical agents
 containing hyaluronic acid for chronic wound healing. Cochrane Database Syst Rev.
 2023 Jul 27;7(7):CD012215
- Roje, Z., Roje, Z., Matic, D., Librenjak, D., Dokuzovic, S., & Varvodic, J. (2011).
 Necrotizing fasciitis: literature review of contemporary strategies for diagnosing and
 management with three case reports: torso, abdominal wall, upper and lower limbs. *World J Emerg Surg*, 6(1), 46

1 2 3	Romanelli M, Dini V, Bertone MS. Randomized comparison of OASIS wound matrix versus moist wound dressing in the treatment of difficult-to-heal wounds of mixed arterial/venous etiology. Adv Skin Wound Care. 2010;23(1):34-38
4 5 6 7	Santoianni P, Monfrecola G, Martellotta D, et al. Inadequate effect of helium-neon laser on venous leg ulcers. Photodermatology 1984;1245-9
7 8 9	Schunk C, Reed K. Clinical Practice Guideline: Examination and Intervention for Rehabilitation, Gaithersburg MD Aspen Publishers, 2000
10 11 12 13 14 15 16	Seidel D, Storck M, Lawall H, Wozniak G, Mauckner P, Hochlenert D, Wetzel-Roth W, Sondern K, Hahn M, Rothenaicher G, Krönert T, Zink K, Neugebauer E. Negative pressure wound therapy compared with standard moist wound care on diabetic foot ulcers in real-life clinical practice: results of the German DiaFu-RCT. BMJ Open. 2020 Mar 24;10(3):e026345
17 18 19	Shahi, N., Bradley, S., Vowden, K., & Vowden, P. (2014). Diabetic bullae: a case series and a new model of surgical management. <i>J Wound Care</i> , <i>23</i> (6), 326, 328-330
20 21 22	Shi J, Gao Y, Tian J, Li J, Xu J, Mei F, Li Z. Negative pressure wound therapy for treating pressure ulcers. Cochrane Database Syst Rev. 2023 May 26;5(5):CD011334
22 23 24	Singer AJ, Clark RAF. Cutaneous wound healing. N Engl J Med. 1999;341:738-746
24 25 26 27 28 29 30	Snyder DL, Sullivan N, Schoelles KM. Skin substitutes for treating chronic wounds. Technology Assessment Report. Prepared by the ECRI Institute Evidence-based Practice Center (EPC) under contract to the Agency for Healthcare Research and Quality (AHRQ), Contract No. HHSA 290-2007-10063. Project ID: HCPR0610. Rockville, MD: AHRQ; December 18, 2012
31 32 33 34 35 36	Snyder DL, Sullivan N, Margolis DJ, Schoelles K. Skin substitutes for treating chronic wounds. Technology Assessment Program Project ID No. WNDT0818. (Prepared by the ECRI Institute-Penn Medicine Evidence-based Practice Center under Contract No. HHSA 290-2015-00005-I) Rockville, MD: Agency for Healthcare Research and Quality. February 2020
37 38	Stanley AC, Lounsbury KM, Corrow K, et al. "Pressure elevation slows the fibroblast response to wound healing". (2005). J. Vasc. Surg. 42 (3): 546–51
 39 40 41 42 	Stoekenbroek RM, Santema TB, Legemate DA, Ubbink DT, van den Brink A, Koelemay MJ. Hyperbaric oxygen for the treatment of diabetic foot ulcers: a systematic review. Eur J Vasc Endovasc Surg. 2014 Jun;47(6):647-55

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1 2 3	Sussman C, Bates-Jensen BM. Wound Care: A Collaborative Practice Manual for Health Professionals (4th ed.). Gaithersburg, Maryland: Aspen Publishers, 2011
4 5 6	Sussman C, Bates-Jensen BM Wound Care: A Collaborative Practice Manual for Physical Therapists and Nurses (2d ed.). Gaithersburg, Maryland: Aspen Publishers, 2001
0 7 8 9	Sutton E, Ganie S, Chan C, Kaur A, Nussbaum E. Photobiomodulation and diabetic foot and lower leg ulcer healing: A narrative synthesis. Foot (Edinb). 2021;48:101847
10 11 12	Swanson, T., Asimus, M., & McGuiness, B. (2014). Wound Management for the Advanced Practitioner: IP Communications Pty, Limited
12 13 14 15 16	Taradaj J, Franek A, Blaszczak E, Polak A, Chmielewska D, Krol P, Dolibog P. Using Physical Modalities in the Treatment of Venous Leg Ulcers: A 14-year Comparative Clinical Study. Wounds. 2012 Aug;24(8):215-26
17 18 19	Thomas, DR. Pressure ulcers. In: Rakel RE, Bope ET, editors Conn's Current Therapy, 1st ed. Philadelphia, PA: WB Saunders;2011.Section 13
20 21 22	Tricco AC, Antony J, Vafaei A, et al. Seeking effective interventions to treat complex wounds: An overview of systematic reviews. BMC Med. 2015;13:89
23 24 25 26 27 28 29	U.S. Food and Drug Administration (FDA). Guidance for Industry and FDA Staff - Class II Special Controls Guidance Document: Non-powered Suction Apparatus Device Intended for Negative Pressure Wound Therapy (NPWT). Updated June 2015. Retrieved on April 29, 2025 from: https://www.fda.gov/medicaldevices/deviceregulationandguidance/guidancedocument s/ucm233275.htm
30 31 32	Vecin NM, Kirsner RS. Skin substitutes as treatment for chronic wounds: current and future directions. Frontiers in Medicine. 2023;10
33 34 35 36	Wang G, Zheng J, Wu H, Wu Y. Effects of electromagnetic therapy in treating patients with venous leg ulcers: An overview of systematic reviews. Int Wound J. 2024 Apr;21(4):e14852
37 38 39 40	Webster J, Liu Z, Norman G, Dumville JC, Chiverton L, Scuffham P, Stankiewicz M, Chaboyer WP. Negative pressure wound therapy for surgical wounds healing by primary closure. Cochrane Database Syst Rev. 2019 Mar 26;3:CD009261
40 41 42	Wheeler PC, Hardwicke HM, Rowley BA. Accelerated healing of skin ulcer by electrotherapy: preliminary clinical results, South Med J. 1969 Jul;62 (7):795-801

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1 2 3 4	White J, Ivins N, Wilkes A, Carolan-Rees G, Harding KG. Non-contact low-frequency ultrasound therapy compared with UK standard of care for venous leg ulcers: a single-centre, assessor-blinded, randomised controlled trial. Int Wound J. 2015 Jan 25
4 5 6	White-Chu EF, Conner-Kerr TA. Overview of guidelines for the prevention and treatment of venous leg ulcers: a US perspective. J Multidiscip Healthc. 2014;11(7):111-7
7	
8 9	Willy C, Voelker HU, Engelhardt M. Literature on the subject of vacuum therapy: review and update 2006. Eur J Trauma Emerg Surg 2007 Feb;33(1):33-9
10	
11	Wolcott LE, Wheeler PC, Hardwicke HM, Rowley BA. Accelerated healing of skin ulcers
12 13	by electrotherapy. South Med J. 1969;62(7): 795-801
13	Wu F, Qi Z, Pan B, Tao R. Extracorporeal shock wave therapy (ESWT) favors healing of
15	diabetic foot ulcers: A systematic review and meta-analysis. Diabetes Res Clin Pract.
16	2024;217:111843. doi:10.1016/j.diabres.2024.111843
17	, , , , , , , , , , , , , , , , , , ,
18	Wu S, Carter M, Cole W, et al. Best practice for wound repair and regeneration use of
19	cellular, acellular and matrix-like products (CAMPs). J Wound Care.
20	2023;32(Sup4b):S1-s31
21	
22	Zarei N, Hassanzadeh-Tabrizi SA. Alginate/hyaluronic acid-based systems as a new
23	generation of wound dressings: A review. Int J Biol Macromol. 2023 Dec 31;253(Pt
24	6):127249
25	
26	Zens Y, Barth M, Bucher HC, Dreck K, Felsch M, Groß W, Jaschinski T, Kölsch H, Kromp
27	M, Overesch I, Sauerland S, Gregor S. Negative pressure wound therapy in patients
28	with wounds healing by secondary intention: a systematic review and meta-analysis of
29	randomised controlled trials. Syst Rev. 2020 Oct 10;9(1):238. doi: 10.1186/s13643-
30	020-01476-6
31	
32	Zhang J, Zhao Y, Zhao X, Zhang J, Jing L. Efficacy and safety of red and infrared light in
33	the adjunctive treatment on diabetic foot ulcers: A systematic review and meta-
34	analysis. Complement Ther Clin Pract. 2024;57:101906.
35	doi:10.1016/j.ctcp.2024.101906
36	
37	Zhou Y, Chia HWA, Tang HWK, Lim SYJ, Toh WY, Lim XL, Cheng LJ, Lau Y. Efficacy
38	of low-level light therapy for improving healing of diabetic foot ulcers: A systematic
39	review and meta-analysis of randomized controlled trials. Wound Repair Regen. 2021
40	Jan;29(1):34-44

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