

NexoBrid®

Disruptive Therapy for Burn Care FDA and EMA approved

- Approved in 40+ countries; 15,000+ patients treated to date
- Poised to replace standard of care for eschar removal in severe burns
- Minimizes need for surgery and significantly reduces blood loss
- Topical application at bedside
- Preserves viable tissue and improves patient outcomes (scar quality and function)
- c-GMP sterile manufacturing facility to support global demand



EscharEx®

Bioactive Enzymatic Debridement Drug Candidate for Chronic Wound

- Rapid, effective, safe debridement for two indications: Venous Leg Ulcers (VLU) and Diabetic Foot Ulcers (DFU)
- Easy to use topical application for all patient settings
- Debrides chronic ulcers within 4-8 applications
- Promotes granulation tissue and reduces bacteria & biofilm
- Demonstrated superiority over SANTYL®
- Targets a \$2.5B market
- De-risked program: based on 3 successful Phase II trials
 - Ongoing Phase III trial for VLU
 - Planned trial for DFU
- Research collaborations with Mölnlycke, Solventum, Coloplast, Convatec and MIMEDX



Global Collaborations



Validated Enzymatic Technology Platform

Solid Balance Sheet

\$39M cash (as of Mar 31, 2025)

\$20M revenues in 2024

3:1 demand to production capacity

\$115M+ BARDA funding (to date)

\$15M+ DoD funding (to date)