Potent laminin-inspired antioxidant regenerative dressing accelerates wound healing in diabetes

Yunxiao Zhu, Zdravka Cankova, Marta Iwanaszko, Sheridan Lichtor, Milan Mrksich, and Guillermo A. Amee

The successful treatment of chronic dermal wounds, such as diabetic foot ulcers (DFU), depends on the development of safe, effective, and affordable regenerative tools that the surgeon can rely on to promote wound closure. Although promising, strategies that involve cell-based therapies and the local release of exogenous growth factors are costly, require very long development times, and result in modest improvements in patient outcome. We describe the development of an antioxidant shape-conforming regenerative wound dressing that uses the laminin-derived dodecapeptide A5G81 as a potent tethered cell adhesion, proliferation, and haptotaxis-inducing ligand to locally promote wound closure. A5G81 immobilized within a thermoresponsive citrate-based hydrogel facilitates integrin-mediated spreading, migration, and proliferation of dermal and epidermal cells, resulting in faster tissue regeneration in diabetic wounds. This peptide-hydrogel system represents a paradigm shift in dermocoductive and dermoinductive strategies for treating DFU without the need for soluble biological or pharmacological factors.

D

Published online June 11, 2018.

PNAS license.

1To whom correspondence may be addressed. Email: milan.mrksich@northwestern.edu or g.amee@northwestern.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1804262115/-/DCSupplemental.

Published online June 11, 2018.


February 2018.
significantly improved the adhesion, spreading, proliferation, and migration of human dermal and epidermal cells in the PPCN network in an integrin-independent manner. This work reports a tethered laminin-derived peptide that is capable of enhancing all of these cellular processes, which are critical to improving wound healing. Upon validation in a large animal model, PPCN-ASG81 may have a faster path to market as a tool to promote healing in diabetic foot ulcers.

Results

The Complete ASG81 Dodecapeptide Sequence Is Necessary to Promote the Adhesion and Haptokinesis of Human Epithelial Keratinocytes. The cost-effective fabrication and scale-up of a peptide-based regenerative dressing requires the identification of the minimum peptide length and amino acid sequence that is necessary to support the adhesion of dermal fibroblasts and epithelial keratinocytes. To achieve these tasks, we performed amino acid truncation and alanine substitution of individual residues of ASG81 using thiol-terminated peptides that were conjugated to self-assembled monolayers (SAMs) of maleimide-terminated dioctadecyl disulfide on gold-coated glass coverslips (all peptides at 1 mol% surface density) (Fig. 1) (18). Both of these modifications resulted in a significant loss of cell adhesion to the immobilized peptide, confirming that the complete ASG81 dodecapeptide and original sequence are necessary for cell adhesion to ASG81 using thiol-terminated peptides that were conjugated to SAMs presenting alanine-substituted versions of ASG81 (Fig. S1). In addition to cell adhesion, the migration of adherent human epithelial keratinocytes (HEKas) on SAMs that display the various peptides, also at the same surface density, was evaluated using live cell time-lapse imaging. Individual cell tracking revealed that the immobilized ASG81 peptide significantly increased the average migration distance of HEKas (Fig. 2A) and effected a twofold increase in their average migration rate (Fig. 2B and SI Appendix, Fig. S2). In addition, the adherent cells were significantly more mobile on the ASG81-presenting surface, demonstrating this peptide’s superior haptokinetic properties (Fig. 2 and SI Appendix, Fig. S2). Consistent with the adhesion study, we found that very few cells adhered to surfaces displaying the inactive peptides, and the migration of these cells was very limited (Fig. 2 and SI Appendix, Fig. S1).

To further determine what other cellular processes are triggered by the cell–peptide interaction, we performed a global gene expression analysis. We compared the transcriptional activities of HEKas cultured on the ASG81-presenting monolayers to those grown on the RGD-presenting monolayers, both at the same surface peptide density of 1% that has been shown to reduce nonspecific interactions (19). Consistent expression readouts were obtained from triplicate experiments within each group, confirming the reliability of the peptide-SAM surface functionalization method. To generate an integrated view of the effects of ASG81 on HEKas, we performed pathway analysis using GOrilla gene ontology enrichment analysis (20). This analysis provides gene clustering information based on known functional interactions such as protein–protein interactions of gene regulation interactions and demonstrates the interconnectedness between genes. Among all of the cellular networks, three cell surface receptors and cell proliferation related networks were identified with a P value lower than 0.05 and were merged together to generate an integrated picture of relevant relationships among the pathways (Fig. 2C and D). Data analysis shows that, during the 12-h incubation time, significant gene expression changes occur in genes involved in cytokine-mediated signaling pathway (gene ontology [GO]: 0019221), cell surface receptor signaling pathway (GO: 0007166), and the negative regulation of growth pathway (GO: 45926). The first two pathways, which include keratinocyte development markers such as KRT7 and KRT19, were mostly up-regulated in cells grown on ASG81 surfaces. These results are consistent with the elevated cell matrix interactions and cellular activity levels triggered by ASG81. Genes associated with negative regulation of cell growth were found to be up-regulated in cells grown on surfaces displaying RGD and not on surfaces displaying ASG81. Additionally, migration-associated genes including ICAM1, OAS2, and IFT2 were up-regulated in the ASG81 group (21, 22). These results are consistent with the increased cell proliferation and migration rates observed in the ASG81 group.

The Biological Functions of ASG81 Can Be Harnessed Within a 3D Thermoresponsive Antioxidant Macromolecular Network. We synthesized an ASG81-presenting thermoresponsive and antioxidant macromolecular network (ASG81-PPCN) by conjugating the cysteine-terminated ASG81 peptide to maleimide-functionalized PPCN (Fig. 3A). With a similar strategy, we also synthesized RGD-presenting PPCN (RGD-PPCN) and inactive peptide-presenting PPCN (inactive-PPCN) as per MALDI data (Fig. 3B and SI Appendix, Fig. S3). The final concentration of the peptide motif in the peptide-PPCN was verified to be 1 mM for all groups throughout the study (SI Appendix, Fig. S4). Tethering the peptide to the macromolecule lowered PPCN’s intrinsic lower critical solution temperature (LCST) by a few degrees (24 °C for ASG81-PPCN vs. 27 °C for PPCN) (SI Appendix, Figs. S5 and S6). The LCST of the peptide-macromolecule conjugate allows the material to be applied as a liquid to the wound and transition to a gel due to the tissue’s increased temperature (Fig. 3C). This property also allows gentle removal of the hydrogel by rinsing the wound with cooled saline during dressing changes, reducing additional trauma to regenerating tissue. Oxidative stress has been shown to exacerbate inflammation and delay the healing of diabetic foot ulcers (23). Therefore, we...
and displayed fibroblast morphology. In contrast, cells entrapped in PPCN or inactive-PPCN remained rounded, suggesting minimal interactions with the matrix (Fig. 4A).

These findings confirm that the observed cell–matrix interactions are facilitated through specific ligand–receptor interactions between the cells and the tethered A5G81 and RGD peptides. It is significant that we observed an increased density of cells entrapped in the A5G81-PPCN hydrogels. Cell cycle analysis of cells entrapped in the various matrices at day 5 showed that a higher percentage of cells in A5G81-PPCN are in the DNA synthesis phase (Fig. 4B and SI Appendix, Table S1). This enhanced DNA synthesis activity within the cell population continued to increase from 21% at day 5 to 27% at day 10. A similar pattern was also observed in the RGD-PPCN group, with an increase from 13% at day 5 to 21% at day 10. The proliferation activity of cells entrapped in PPCN or inactive-PPCN was significantly lower than those measured in A5G81-PPCN and RGD-PPCN (Fig. 4B). Consistently, the total DNA quantification results show that A5G81-PPCN supported a higher cell proliferation rate over the course of 10 d than the other groups, confirming that A5G81 is able to promote the proliferation of HDFs in 3D (Fig. 4C). To further understand the mechanism for the enhanced proliferation of HDFs in A5G81-PPCN, we investigated the binding interactions between tethered A5G81 and the integrin receptors α3β1 and α6β1, which we confirmed are expressed by the cells using anti-α3 and anti-α6 antibodies (Fig. 4 D and E). Blocking only α3 or α6 leads to a partial decrease in cell proliferation, whereas blocking both α3 and α6 reduced cell proliferation to values that were comparable to those observed in the other hydrogels. Antibody blocking did not have an impact on cell proliferation in PPCN, inactive-PPCN, and RGD-PPCN. These results suggest that the increased cell proliferation within assessed whether the intrinsic antioxidant properties of PPCN were affected by the tethering of A5G81. The results of a lipid peroxidation inhibition assay show that conjugation of the peptides to PPCN resulted in protection against oxidation that was 125% and 160% that of PPCN’s for RGD-PPCN and A5G81-PPCN, respectively (Fig. 3D). Further study of the intact peptides revealed that this enhanced antioxidant property of the peptide-functionalized PPCN is likely due to the intrinsic antioxidant property of the peptide itself (SI Appendix, Fig. S7), as certain amino acids are known to have antioxidant activities due to the reactivity of their side chains (24). Therefore, the thermo-responsive and antioxidant properties of PPCN, in combination with the properties of the tethered laminin-derived peptide A5G81, are expected to provide an optimal microenvironment that promotes tissue regeneration.

All of the hydrogel formulations supported the long-term viability of human dermal fibroblasts (HDFs) (Fig. 4A). Given that a scaffold for wound healing should facilitate the spreading, migration, and proliferation of surrounding cells into the scaffold, we used cytoplasmic calcine staining, confocal microscopy, and flow cytometry to evaluate cell spreading and proliferation of HDFs within the hydrogels. HDFs seeded in A5G81-PPCN and RGD-PPCN began to spread and interact with the surrounding matrix at day 5 postseeding, and, by day 10, cells were fully elongated...
A5G81-PPCN is mediated by the specific integrin-peptide interactions between the HDFs and tethered A5G81.

**A5G81-PPCN Significantly Enhances Closure Rate, Reepithelialization, and Granulation Tissue Formation in Wounds of Diabetic Animals.**

We hypothesized that the observed enhanced in vitro cell proliferation and migration effects in A5G81-PPCN will translate to an accelerated wound healing response in vivo. To test this hypothesis, we investigated wound closure in a splinted excisional wound diabetic mouse model (25). This preclinical animal model is widely accepted as a first step to study wound healing therapies that could potentially benefit diabetic patients. All animals received two splinted dorsal wounds. One wound was treated with A5G81-PPCN, and the contralateral wound was treated with either RGD-PPCN, inactive-PPCN, or PPCN. Wounds treated with A5G81-PPCN healed at a significantly faster rate than wounds treated with the PPCN, inactive-PPCN, and RGD-PPCN, with 45% closure achieved at day 10 relative to 20% for inactive-PPCN and PPCN, and 31% for RGD-PPCN (Fig. 5 A–C). A5G81-PPCN also achieved 79% healing by day 15 and complete closure by day 22 postwounding (Fig. 5D and SI Appendix, Fig. S10). Wound reepithelialization and granulation tissue formation was assessed histologically through measurements of the epithelial gap and granulation tissue thickness (Fig. 6 A–C and SI Appendix, Fig. S10). Histology of the regenered wounded tissue at day 30 revealed close to complete reepithelialization in A5G81-PPCN–treated wounds, whereas PPCN, inactive-PPCN, and RGD-PPCN–treated wounds had remaining gaps of 2.15 ± 0.82 mm, 2.85 ± 1.08 mm, and 0.74 ± 0.64 mm, respectively (Fig. 6C and SI Appendix, Fig. S9). A5G81-PPCN–treated wounds had a multilayered epithelium structure that closely resembled healthy epidermis of the intact skin (26). The epithelial layer was significantly thinner in the RGD-PPCN–treated wounds, and discontinuities were observed in wounds treated with PPCN and inactive-PPCN (Fig. 6D and SI Appendix, Fig. S10B). A5G81-PPCN–treated wounds were also populated by a significant amount of α3 positive cells, which further supports the role of A5G81 in promoting cell infiltration and proliferation during the healing process (Fig. 6D). Staining for the macrophage cell marker F4/80 shows few macrophages in all groups, with the least amount within the A5G81-PPCN–treated tissue, suggesting a more subdued inflammatory status consistent with a more advanced stage of the healing process (Fig. 6D). To assess whether a significant difference in reepithelialization was also observed at an earlier time point, the wounded tissue of animals that received A5G81-PPCN and inactive-PPCN was assessed via histology at day 10 postwounding (SI Appendix, Fig. S11 A–C). Wounds treated with A5G81-PPCN have an average gap size of 0.4 ± 0.3 mm compared with 2.4 ± 0.6 mm for the wounds treated with inactive-PPCN (P < 0.001; SI Appendix, Fig. S11C). All wounds treated with A5G81-PPCN had granulation tissue that was more prominent relative to the contralateral wound (Fig. 6B and SI Appendix, Fig. S11C).

To benchmark A5G81-PPCN against another clinically used wound dressing for diabetic wound care management, we performed a side-by-side comparison study where one wound was treated with A5G81-PPCN and the contralateral wound was treated with the Promogran Prisma wound dressing (Systagenix), which is based on collagen and cellulose impregnated with silver. A5G81-PPCN outperformed Prisma starting on day 6 postwounding, with 50% of the wound area closed by day 15 and 90% closed by day 25. In contrast, wounds treated with Prisma exhibited 10% and 75% wound closure by days 15 and 25 postwounding, respectively (SI Appendix, Fig. S12 A–C). To mimic the presence of fluid in the wound, a condition often found in human wounds and not murine models, additional experiments...
were conducted with preswollen wet Prisma. A5G81-PPCN also outperformed the wet Prisma matrix (SI Appendix, Fig. S12 D–F).

Discussion
In order for chronic diabetic wounds to heal, skin cells must migrate and proliferate in a harsh microenvironment. We demonstrate that A5G81 is a potent adhesion ligand that is capable of facilitating both of these processes in an integrin-dependent manner that mimics laminin (11, 13). We also show that A5G81 covalently linked to a thermoresponsive antioxidant macromolecule is able to accelerate healing in splinted excisional wounds in diabetic mice. This animal model of hyperglycemia results in the characteristic diabetic complications observed in diabetic wound healing, including reduced chemokine and growth factor release, impaired angiogenesis, prolonged inflammation, and increased oxidative stress (25, 27). Another advantage of this model is the capability to mimic human wound healing processes by using a splint around the wound to prevent wound contraction, which is characteristic of wound healing in rodents, as they are loose-skin animals (28, 29). Therefore, the wound is allowed to heal mostly through tissue regeneration, resembling the healing process of humans (SI Appendix, Fig. S8) (30). Although various peptides have been successfully used to enhance wound healing in diabetic mice, most of these treatments involve topical application as a solution, requiring multiple repeat applications (31–33). Furthermore, a majority of those studies also did not take into account skin contraction during the healing process. Therefore, it is unclear how treatments evaluated in rodent nonsplinted wound healing models affect the regeneration component of the healing process, which is the relevant process to wound healing in humans (31). For example, Van Slyke et al. (34) reported 70% closure in nonsplinted wounds within 7 d postwounding with the use of a soluble peptide derived from angiopoietin. There was no control or accounting for how the treatment affects skin contraction, making it difficult to assess the potential benefit of their peptide for treating human wounds. Xiao et al. (35) recently reported the use of a tethered angiopoietin-derived peptide to heal nonsplinted wounds in diabetic mice and reported a 60% wound closure at day 14 with low-peptide treatment and 75% with high-peptide treatment. The peptide was tethered to a chitosan/collagen gel and added to the wound. Their histology data at days 14 and 21 postwounding show both contraction and regeneration as mechanisms for wound closure. The granulation tissue, as per their histology data, was also significantly less developed than what we observed in our study, reinforcing the importance of extracellular matrix proteins in promoting physiological wound healing. As for laminin-derived peptides, several dermal wound healing studies have reported that these peptides can play a positive role in accelerating wound closure (12, 33). Reported mechanisms include the promotion of angiogenesis and the support of keratinocyte migration (33, 36, 37). A brief summary of dermal wound healing studies that used laminin-derived peptides is included in SI Appendix to provide additional context to our findings (SI Appendix, Table S2).

To our knowledge, A5G81 is the only laminin-derived peptide to date that exhibits a potent and synergistic effect on migration and proliferation for both keratinocytes and dermal fibroblasts. Furthermore, at the same molar concentration of tethered peptide, the effects of A5G81 were superior to those of RGD, a peptide commonly used to promote cell adhesion. The accelerated wound healing observed in wounds treated with A5G81-PPCN compares favorably to wound-closure rates reported in this mouse model using growth factors and stem cell strategies. We previously reported 75% wound closure at day 15 and complete closure at 24 d postwounding when splinted wounds were treated with the slow release of low-dose stromal cell-derived factor 1α (38). Galiano et al. (39) reported an accelerated healing time of 17 d with the high dose of topical application of vascular endothelial growth factor. Two other research groups that investigated the treatment of wounds with allogeneic mesenchymal stem cells reported wound closures of 45% and 56% at 28 d postwounding (40, 41). Although promising, therapies that use cells, the local release of drugs, or the local release of macromolecules to promote wound healing will face higher regulatory challenges due to safety concerns, extended product development times, and higher costs.

Hydrogel dressings composed exclusively of peptides or extracellular matrix fragments have also shown promising results in treating dermal wounds in diabetic mice and can have an easier path to market (42, 43). However, the reported hydrogels do not have the shape-conforming and intrinsic antioxidant properties exhibited by the peptide-PPCN scaffold reported herein. Xiao et al. (35) and others have demonstrated that inhibiting oxidative stress in the wound can improve the wound healing response. Therefore, we probed healed wounds for markers of oxidative DNA damage. Wounds treated with A5G81-PPCN had significant attenuation of DNA oxidative damage relative to all other treatments (SI Appendix, Fig. S13). The decreased oxidative tissue damage may be due to the higher antioxidant nature of A5G81-PPCN or to a more advanced stage of healing in A5G81-PPCN–treated wounds. Therefore, this biomaterial-based approach to treat wound healing in diabetes may work through the simultaneous positive impact on the adhesion, proliferation, migration, and oxidative stress status of dermal cells. The strongest evidence for the treatment of diabetic ulcers using peptide hydrogels comes from a clinical trial using Argidene Gel—a peptide hydrogel matrix composed of RGD and sodium hyaluronate (44). The use of the RGD
hydrogel resulted in 4 times the number of patients with complete wound healing compared with the placebo group (44). Based on our results, we expect that the ASG81-PPCN dressing will provide a greater benefit to the patient.

Conclusion

We have identified and characterized a laminin-derived peptide with unique receptor-mediated and antioxidant properties that are beneficial to the wound healing process. Conjugating this peptide to PPCN resulted in a shape-conforming material that can be used as a regenerative dressing to enhance wound healing. Notably, ASG81 outperformed the fibronectin–derived cell adhesion peptide RGD in its ability to induce cell motility and proliferation in vitro, and the mechanism was shown to involve the integrin subunits α5 and α6. Future studies will investigate ASG81-PPCN in a large animal model for wound healing.

Materials and Methods

Peptide Conjugation and Characterization.

Conjugation was achieved using the bifunctional linker N-p-maleimidopropionic acid hydrazide (BMPH) (Thermo Fisher) The completion of the reaction between the peptide and the BMPH-PPCN was confirmed by quenching free thiol groups in the reaction medium over time using Ellman's reagent. The successful conjugation was confirmed using MALDI-TOF (Fig. 3B and SI Appendix, Fig. S3). The generation of primary amines within PPCN due to the conjugation was quantified with the trinitrobenzenesulfonic assay and used to calculate the final concentration of peptide tethered to PPCN using a standard curve for each peptide. The 1 mol% peptide on all SAM surfaces was verified by the peak intensity ratio between the peptide-maleimide-PEG-alkanethiols and the PEG-end-capped alkanethiols via MALDI-TOF mass spectrometry (18).

Diabetic Wound Healing Model.

All the experimental procedures involving animals were approved by the Institutional Animal Care and Use Committee of Northwestern University. The in vivo performance of the hydrogels was evaluated using an split-squared exudation wound model in type II diabetic mice [B6.Ks(Duart-LeprFloy)]. The Jackson Laboratory as previously described (25, 38). Doughnut-shaped acrylate splints were attached to the left and right dorsal sides of the mouse, and a 6-mm circular, full-thickness wound was made in the center of each splinted area. Forty microliters of PPCN solution was applied to each wound bed and covered with TegaDerm.

Statistical Analysis.

GraphPad Prism 6.0c was used for two-way ANOVA tests to measure differences for experiments with multiple data sets. A Tukey test was performed between groups with significant differences to correct for the multiple pair-wise comparisons. A value of P ≤ 0.05 was considered to be statistically significant.

Details for the materials and methods used for our protocols are provided in SI Appendix.