Acceleration of Diabetic Wound Healing with PHD2- and miR-210-Targeting Oligonucleotides

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In diabetes-associated chronic wounds, the normal response to hypoxia is impaired and many cellular processes involved in wound healing are hindered. Central to the hypoxia response is hypoxia-inducible factor-1α (HIF-1α), which activates multiple factors that enhance wound healing by promoting cellular motility and proliferation, new vessel formation, and re-epithelialization. Prolyl hydroxylase domain-containing protein 2 (PHD2) regulates HIF-1α activity by targeting it for degradation under normoxia. HIF-1α also upregulates microRNA miR-210, which in turn regulates proteins involved in cell cycle control, DNA repair, and mitochondrial respiration in ways that are antagonistic to wound repair. We have identified a highly potent short synthetic hairpin RNA (sshRNA) that inhibits expression of PHD2 and an antisense oligonucleotide (antimiR) that inhibits miR-210. Both oligonucleotides were chemically modified for improved biostability and to mitigate potential immunostimulatory effects. Using the sshRNA to silence PHD2 transcripts stabilizes HIF-1α and, in combination with the antimiR targeting miR-210, increases proliferation and migration of keratinocytes in vitro. To assess activity and delivery in an impaired wound healing model in diabetic mice, PHD2-targeting sshRNAs and miR-210 antimiRs both alone and in combination were formulated for local delivery to wounds using layer-by-layer (LbL) technology. LbL nanofabrication was applied to incorporate sshRNA into a thin polymer coating on a Tegaderm mesh. This coating gradually degrades under physiological conditions, releasing sshRNA and antimiR for sustained cellular uptake. Formulated treatments were applied directly to splinted full-thickness excisional wounds in db/db mice. Cellular uptake was confirmed using fluorescent sshRNA. Wounds treated with a single application of PHD2 sshRNA or antimiR-210 closed 4 days faster than untreated wounds, and wounds treated with both oligonucleotides closed on average 4.75 days faster. Markers for neovascularization and cell proliferation (CD31 and Ki67, respectively) were increased in the wound area following treatment, and vascular endothelial growth factor (VEGF) was increased in sshRNA-treated wounds. Our results suggest that silencing of PHD2 and miR-210 either together or separately by localized delivery of sshRNAs and antimiRs is a promising approach for the treatment of chronic wounds, with the potential for rapid clinical translation.

Keywords: HIF-1α, miR-210, sshRNA, wound healing, PHD2, antimiR, diabetes
Introduction

Diabetes affects ~25 million adults in the United States, with these numbers expected to double by the year 2050. Approximately one-quarter of these patients develop at least one foot ulcer in their lifetime, increasing the risk of lower limb amputations. The recurrence rate of diabetic ulcers is 66% and with subsequent ulcerations the amputation rate is 12%. Nonhealing diabetic wounds are thus a tremendous challenge for patients and caregivers alike and cost the U.S. healthcare system $9–13 billion annually. Microvascular impairments and dysfunctional neovascularization are the leading cause of pressure ulcers and chronic wounds in diabetic patients. These defects have been linked to a diminished cellular hypoxia response, of which the master regulator is the transcriptional factor, hypoxia-inducible factor-1α (HIF-1α).

HIF-1α levels are negatively regulated by prolyl hydroxylase domain-containing protein 2 (PHD2), a crucial cellular oxygen sensor that hydroxylates two specific proline residues in HIF-1α, tagging it for rapid degradation via the proteasome pathway in normoxia. However, in the hypoxic conditions of normal acute wounds, hydroxylation and degradation of HIF-1α are reduced, resulting in increased translocation of HIF-1α to the nucleus where it transactivates factors involved in vasculogenesis, angiogenesis, re-epithelialization, and cell survival that promote wound healing. These include vascular endothelial growth factor (VEGF), erythropoietin (EPO), platelet-derived growth factor (PDGF), fibroblast growth factor (FGF), transforming growth factor beta (TGF-β), stromal cell-derived factor 1 (SDF-1), and heat shock protein 1 (HSP-1) (Fig. 1). In diabetic patients, induction of these factors is hampered due to high-glucose-induced modification of p300, a cofactor of HIF-1α transactivation. Stabilization of HIF-1α may be achieved through inhibition of PHD2 expression, and such intervention can improve wound closure in diabetic mice (Fig. 1).

Hypoxia also induces the expression of certain microRNAs (miRNAs), called hypoxamirs, some of which are involved in wound healing. One such miRNA, miR-210, is upregulated by HIF-1α and targets E2F3, a cell cycle regulator that facilitates the G1/S transition. This activity is thought to be responsible for the finding that miR-210 attenuates keratinocyte proliferation as well as re-epithelialization in ischemic wound healing. The stabilization of HIF-1α through pharmacological inhibition of PHD2 alone would be expected to upregulate miR-210, and have negative consequences for wound healing.

We therefore hypothesize that stabilization of HIF-1α via PHD2 inhibition might be most effective if miR-210 is also inhibited. Because of the ease of combining oligonucleotides in a delivery cocktail, we decided to use an RNA-based approach to accelerate the wound healing process by inhibiting each of these hypoxia response targets (Fig. 1). To inhibit expression of PHD2, we used a novel class of short synthetic hairpin RNAs (sshRNAs); such RNAs have previously been shown to be highly potent inhibitors of hepatitis C virus in mouse models. sshRNAs induce cleavage of target RNAs by a novel Dicer-independent RNA interference (RNAi) pathway that is similar to the noncanonical miR-451 processing pathway. To inhibit miR-210, an antisense oligonucleotide (anti-miR) approach targeting miR-210 was used.

Delivery of oligonucleotides to tissues other than the liver remains a significant hurdle to the development of effective nucleic acid-based therapeutics. Because of the nature of wound sites, we have opted for a localized rather than systemic approach to delivery of the RNAi and antisense oligonucleotides. Using a novel layer-by-layer (LbL) technology, we formulated the oligonucleotides into a thin film polymer coating atop a nylon woven mesh. This
formulation provides slow release of the encapsulated oligonucleotides into the wound bed over the course of 7–10 days. Previous studies have shown that siRNAs delivered by this method are taken up by cells.29,30

In this study, we report the identification of an sshRNA (SG404) and an antimiR (SG608), both chemically modified, that potently inhibit PHD2 and miR-210, respectively, and promote keratinocyte growth and mobility in vitro. When these RNAs were formulated into LbL-coated dressings and applied to full-thickness excisional wounds in diabetic mice, time to wound closure was significantly reduced and increased expression was seen in markers of neovascularization and proliferation in the wound area. These effects were observed with each inhibitor alone and with enhanced effect when they were combined in a single LbL formulation.

Materials and Methods

In vitro target knockdown assays

Cells (primary keratinocytes [NHEK], HaCaT, or NIH-3T3) were seeded at 23,000 cells/well in 96-well or at 30,000 cells/well in 48-well plates 1 day before transfection. Triplicate transfections of PHD2 sshRNAs and miR-210 antimiRs at various concentrations along with nonspecific control (NSC) sshRNAs and antimiRs were performed using HiPerFect (Qiagen) or RNAiMAX (Invitrogen, Carlsbad, CA) following the manufacturer’s instructions. Total RNA was extracted using the RNeasy 96 or miRNeasy kits (Qiagen) and the biosensor assay to measure the activity of antimiR targeting miR-210 was performed as described in Supplementary Methods [Supplementary Data are available online at www.liebertpub.com/tea for details]. mRNA and miRNA levels were quantified by real-time quantitative polymerase chain reaction (RT-qPCR) on a 7500 Fast RT-PCR system incorporating the following oligonucleotides: (1) PHD2-specific sshRNA SG404, (2) miR-210 antimiR SG608, (3) SG404 combined with SG608, and (4) NSC RNA (SG221c). Oligonucleotide loading within the thin film coating is approximated to be 25 μg/cm². The RNA-coated dressing was applied directly to the wound bed.

Proinflammatory cytokine detection in vitro

Assessment of proinflammatory cytokine levels on transfection of PHD2 sshRNAs and miR-210 antimiRs into MRC-5 lung fibroblast cells was performed exactly as described in Dallas et al.20

Formulation of oligonucleotides for in vivo delivery

sshRNA and antimiR oligonucleotides were formulated into a thin film coating assembled by LbL deposition onto the surface of a woven nylon wound dressing (Tegaderm™; 3M, St. Paul, MN) by LayerBio, Inc. (See the Results section for a general description of the method.) Four different wound dressing samples were generated for in vivo testing incorporating the following oligonucleotides: (1) PHD2-specific sshRNA SG404, (2) miR-210 antimiR SG608, (3) SG404 combined with SG608, and (4) NSC RNA (SG221c). Oligonucleotide loading within the thin film coating is approximated to be 25 μg/cm². The RNA-coated dressing was applied directly to the wound bed.

Animals

All mice were housed in the Stanford University Veterinary Service Center in accordance with the National Institutes of Health (NIH) and institution-approved animal care guidelines. All procedures were approved by the Stanford Administrative Panel on Laboratory Animal Care.

Wound model

Fourteen-week-old male C57BL/6 db/db mice (BKS.Cg-Dock7m +/- Leprdb/J) were purchased from Jackson Laboratory (Bar Harbor, ME). A stented excisional wound healing model was used as previously described.33 Briefly, after induction of anesthesia, all hair on the dorsum of the mice was removed using a shaver and depilatory cream. Two excisional wounds were then generated on the dorsum of each mouse using a 6 mm disposable biopsy punch (Integra). To prevent contraction, each wound was splinted open using a silicone ring (Grace BioLabs) with outer and inner diameters of 16 and 10 mm, respectively, attached with cyanoacrylate glue (Krazy Glue, West Jefferson, OH) and 6-0 black nylon sutures (Ethicon).

Administration of LbL-formulated oligonucleotides

One day after wounding, oligonucleotide formulated dressings were cut with a 6 mm disposable biopsy punch and
placed into the wound bed. Treatments were secured in place by an occlusive adherent dressing (Tegaderm; 3M). As the wound healed and closed, each patch was carefully removed and trimmed to fit the wound area.

Wound analysis
Dressings were removed and digital photographs were taken of each wound on the day of surgery and every other day until closure. The analysis of the wound area was performed by measuring the wound area normalized to the inner diameter of the silicone ring. The ratio of the wound area for all time points to initial wound area on the day of surgery was measured and calculated by a blinded observer using ImageJ (NIH, Bethesda, MD).

Histological analysis
Wounds were harvested with a 2 mm ring of unwounded skin from euthanized mice. The wounds were then bisected into hemispherical pieces. The pieces of skin tissue were fixed overnight in 4% paraformaldehyde followed by a serial dehydration in ethanol and paraffin embedding. Five-micron-thick paraffin sections were prepared from wound tissue harvested on days 4, 7, and on wound closure after treatment with LbL-oligonucleotide formulations. Day 4 sections were stained for CD31 (Abcam, Cambridge, MA) and Ki67 (Abcam). Day 7 sections were stained for von Willebrand factor (vWF; antibody No. 7356; EMD Millipore, St. Louis, MO). Healed wounds were stained with hematoxylin and eosin (H&E) and trichrome. Nuclei were stained with DAPI (EMD Millipore). Quantification was performed using ImageJ.

Protein analysis
Wound tissue was harvested after euthanasia on days 2 and 4 post-treatment application similarly to the histology sections except that the remaining hemispherical sections were bisected and designated for protein or RNA analysis. Samples for protein analysis were snap-frozen in microcentrifuge tubes and stored at −80°C until testing. The tissue samples were then homogenized (KINEMATICA polytron; Thermo Fisher Scientific, Waltham, MA) and protein was extracted using RIPA buffer (Sigma-Aldrich, St. Louis, MO) and Halt protease inhibitor cocktail (Thermo Fisher Scientific, Waltham, MA). The isolated soluble protein extract was then analyzed for VEGF protein levels (ELISA Quantikine kits; R&D Systems, Inc., Minneapolis, MN).

Statistical analysis
Statistical analysis was performed using an unpaired Student’s t-test. Values are presented as mean ± standard error of the mean (SEM). p-values <0.05 were considered statistically significant.

Results
Identification of potent sshRNA inhibitors of human and mouse PHD2
We iteratively designed and screened for potency sshRNA inhibitors of both human and mouse versions of PHD2 based on target site prediction algorithms and previously reported siRNA sequences8–10,34 and then performed sequence walking around effective target sites. All sshRNAs were designed to have left-hand loop orientation, 19-bp stems, and a 2-nt loop, a structural design that was previously shown to be effective both in vitro22,23,35 and in vivo.20,21 Each sshRNA was screened for the ability to inhibit PHD2 expression in cell culture, resulting in the identification of SG302 (human) and SG402 (mouse) as the most potent inhibitors of PHD2 expression. Dose/response curves (Fig. 2A, B) show highly potent knockdown of PHD2 mRNA (IC50 = 3.5 pM for SG302, 29 pM for SG402).

Chemical stabilization of sshRNAs and antimiRs
Chemical modification of RNA oligonucleotides serves both to enhance biostability by increasing resistance to degradation by serum nucleases and to mitigate potential undesirable immune stimulatory effects. Chemically modified versions of SG302 and SG402 (SG302m1 and SG404, respectively) containing 2'-OMe modifications at select positions in the sense (or passenger) strand sequence and loop nucleotides as described in Dallas et al.20 and Ge et al.35 were synthesized and their ability to inhibit PHD2 mRNA expression was examined in cell culture. The inhibitory dose curves are shown in Figure 2A and B alongside their unmodified counterparts, confirming that these modified oligonucleotides retain high potency for their respective targets ([SG302m1] IC50, 14 pM; [SG404] IC50, 44 pM). A modified 5'-Rapid Amplification of cDNA (complementary DNA) Ends (5'-RACE) assay was performed to confirm that PHD2 mRNA was cleaved at the predicted site of RNAi-mediated cleavage (Supplementary Fig. S1).

Identification of a potent antisense inhibitor (antimiR) of miRNA-210
Under normoxic conditions, miR-210, a key miRNA upregulated by HIF-1α, has been shown to negatively impact wound healing by inhibiting keratinocyte growth and proliferation.18 By identifying an effective antimiR to miR-210, we aimed to abrogate the negative effects of miR-210 upregulation on stabilization of HIF-1α. AntimiRs were screened for potency in cell culture using a luciferase reporter biosensor (pSG247) assay. In this assay, rLuc is derepressed to an extent that correlates with antimiR efficacy. We tested a variety of chemical modification patterns, including various combinations of 2'-OMe, locked nucleic acid, and phosphorothioates (data not shown), with and without N,N-diethyl-4-(4-nitronaphthalen-1-ylazo)-phenylamine (“ZEN”) moieties at both the 5′- and 3′-ends. The ZEN modification increases the Tm for target binding and increases stability toward serum and cytoplasmic endonucleases and exonucleases.36 One antimiR (SG608), comprising all 2'-OMe-substituted residues with ZEN-modified ends, proved to be particularly potent in cell culture. SG608-transfected cells exhibited derepression of rLuc expression by up to threefold compared with cells lacking antimiR (Fig. 2C).

Simultaneous and specific inhibition of PHD2 and miR-210
Having demonstrated the specific inhibition of miR-210 and PHD2 targets with each inhibitor alone, we also tested...
them in combination in HaCaT cells (Fig. 2D) and monitored for effects on both PHD2 and miR-210 48 h after transfection by RT-qPCR. When PHD2 sshRNAs were transfected alone, PHD2 mRNA levels were reduced by 85%, while miR-210 levels were increased twofold, as expected. In contrast, when anti-miR-210 was transfected alone, PHD2 mRNA levels were unaffected, while miR-210 levels were strongly reduced. Finally, in cells cotransfected with both PHD2-sshRNA and antimiR-210, the PHD2 and miR-210 levels were reduced to the same extent as with each inhibitor alone.

Inhibition of PHD2-sshRNA results in increased HIF-1α expression

The efficacy of PHD2-sshRNA in upregulating HIF-1α was assessed by using a luciferase reporter assay. Forty-eight hours after cotransfection of the reporter plasmids and either PHD2-specific sshRNA (SG302) or NSC sshRNA (SG221c), we confirmed that SG302 was able to upregulate HIF-1α (Fig. 3A). We further validated this result by Western blot, where we observed a 1.8-fold increase in HIF-1α levels relative to controls (Supplementary Fig. S2A).

Demonstration of enhanced keratinocyte growth with PHD2 and miR-210 inhibitors

To measure cell migration, we performed a scratch assay with HaCaT cells as described in Kioka et al.37 Figure 3B and Supplementary Figure S2B show that keratinocytes narrowed the width of the cell-free region much more rapidly when transfected with the active anti-miR-210 and PHD2 sshRNAs than with PHD2 sshRNA alone or NSCs (p = 0.007, 0.001, and 0.003 for 24, 48, and 72 h time points, respectively).

Chemically modified oligonucleotides do not induce proinflammatory cytokines, are not cytotoxic, and have improved biostability

Unmodified RNAs are capable of inducing the undesired expression of proinflammatory cytokines, but this off-target effect can be avoided by incorporating 2′-OMe nucleotides.35,39 We evaluated the cytokine-inducing ability of sshRNAs and antimiRs with and without chemical modifications in human fetal lung fibroblast (MRC-5) cells, which
are known to be sensitive to immunostimulatory oligonucleotides and can be activated by cytosolic RNA through PHD2-dependent luciferase reporter. (B) PHD2 and miR-210 inhibitors increase keratinocyte migration in scratch assay. Percent scratch closure is shown at 24, 48, and 72 h after human HaCaT keratinocytes were transfected with the indicated inhibitors at 10 nM final concentration. SG302m is chemically modified PHD2 sshRNA, PHD2-NSC is a scrambled NSC sshRNA, SG302m+SG608 is cotransfected PHD2 sshRNA and N,N-diethyl-4-(4-nitronaphthalen-1-ylazo)-phenylamine (ZEN)-modified miR-210 antimiR-210, combination NSC is cotransfection with NSC sshRNA and NSC antimiR.

**FIG. 3.** Verification of functional activities induced by inhibition of PHD2 and miR-210. (A) Induction of HIF-1α by human PHD2-targeting sshRNA SG302 by transactivation of a PHD2-dependent luciferase reporter. (B) PHD2 and miR-210 inhibitors increase keratinocyte migration in scratch assay. Percent scratch closure is shown at 24, 48, and 72 h after human HaCaT keratinocytes were transfected with the indicated inhibitors at 10 nM final concentration. SG302m is chemically modified PHD2 sshRNA, PHD2-NSC is a scrambled NSC sshRNA, SG302m+SG608 is cotransfected PHD2 sshRNA and N,N-diethyl-4-(4-nitronaphthalen-1-ylazo)-phenylamine (ZEN)-modified miR-210 antimiR-210, combination NSC is cotransfection with NSC sshRNA and NSC antimiR.

**TABLE 1. IMMUNOSTIMULATORY PROPERTIES OF SHORT SYNTHETIC HAIRPIN RNAs AND ANTIMI�RS**

<table>
<thead>
<tr>
<th>SSHRNA</th>
<th>TNF-α</th>
<th>IL-6</th>
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<tbody>
<tr>
<td>No inhibitor</td>
<td>1.00 ± 0.02</td>
<td>1.02 ± 0.14</td>
</tr>
<tr>
<td>Poly I:C</td>
<td>263.03 ± 6.41</td>
<td>1427.51 ± 31.32</td>
</tr>
<tr>
<td>SG302</td>
<td>0.49 ± 0.02</td>
<td>0.68 ± 0.13</td>
</tr>
<tr>
<td>SG302m</td>
<td>2.25 ± 0.43</td>
<td>1.01 ± 0.10</td>
</tr>
<tr>
<td>AntimiR</td>
<td>No inhibitor</td>
<td>1.00 ± 0.25</td>
</tr>
<tr>
<td>Poly I:C</td>
<td>6494.43 ± 434.01</td>
<td>528.38 ± 24.93</td>
</tr>
<tr>
<td>SG601</td>
<td>0.95 ± 0.50</td>
<td>1.21 ± 0.40</td>
</tr>
<tr>
<td>SG608</td>
<td>1.32 ± 0.39</td>
<td>0.97 ± 0.00</td>
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sshRNA, short synthetic hairpin RNA.

found in serum. sshRNAs without chemical modification are susceptible to degradation in 10% human serum despite being largely double-stranded. We found that stability increased on 2'-OMe modification at alternate nucleotides of the sense strand and each nucleotide in the UU loop (SG404 vs. SG402, Supplementary Fig. S4). SG608, which is a fully modified single-stranded oligonucleotide, was shown to be extremely stable in 10% human serum, with t1/2 > 96 h.

**Oligonucleotide formulation for in vivo delivery**

SG404 and SG608 were formulated using LbL technology (LayerBio, Inc.). LbL is a technique through which a thin film is generated based on the alternating adsorptions of two or more polymers or other species on the basis of complementary interactions (charge, hydrogen bonding). For this study, negatively charged nucleic acids (SG404, SG608, and/or control oligonucleotides) were coassembled via LbL with positively charged polymers into a thin multilayer coating onto the surface of a nylon woven dressing (Tegaderm) following previously demonstrated methods. Briefly, the layers are applied by alternately immersing the substrate into reservoirs containing solutions of positively and negatively charged polymers, respectively, rinsing after each immersion. The RNA-coated dressing is applied directly to the wound bed and biodegrades over 7–10 days with a sustained release of ssRNA-polyacation polyplexes.

To confirm effective delivery of these RNAs in vivo, a fluorescent-tagged version of PHD2 sshRNA (SG405) formulated by LbL was applied to full-thickness excisional wounds in db/db mice (Supplementary Methods). The wounds were imaged 2 and 6 days after application (Fig. 4A) and fluorescence intensity was quantified (Fig. 4B). The results showed uptake of SG405-sshRNA into cells in the dermal layer of the wound area at both time points.

**LbL delivery of SG404 and SG608 improves wound healing**

To assess the activity of the therapeutic oligonucleotides, LbL-formulated SG404, SG608, or a combination of SG404/SG608 was applied to full-thickness excisional wounds (n = 8) in diabetic mice 1 day postwounding and compared with both an untreated control group and group treated with an LbL-formulated NSC sshRNA. As shown in Figure 4C–E, SG404- and SG608-treated wounds closed 4 days faster than untreated control wounds: on average, 17.25 days after wounding versus 21.25 days [p (SG404 vs. control) = 0.002 and p (SG608 vs. control) = 0.0006]. Wounds treated with both SG404 and SG608 simultaneously closed 4.75 days faster (SEM 0.32) than untreated control. At earlier time points, SG404, SG608, and the combination show a similar rate of closure (51%, 55%, 56%, respectively, vs. 40% control at day 10), but after 2 weeks, the combination treatment results in accelerated wound closure over the individual treatments (86%, 84%, 95% vs. 68% control at day 16) (Fig. 4C). LbL-formulated NSC-treated wounds closed 1 day faster than untreated control wounds, suggesting that the dressing itself may partially contribute to the accelerated wound closure. Nevertheless, the specific treatments SG404, SG608, and combined SG404/SG608 still closed significantly faster than LbL-NSC (p = 0.01, 0.006, and 0.0003, respectively).
Trichrome and H&E staining of remodeled wound beds demonstrated increased collagen deposition and appendage remodeling, particularly in the SG608 treatment. We observed a more defined dermal layer and increased collagen in the combination treatment (Supplementary Fig. S5). In all of the specific treatment groups, there was an increase in the average number of appendages per wound versus controls: SG404 (27.6 ± 4.3), SG608 (38.4 ± 18.2), SG404+SG608 (40.6 ± 13.6) versus untreated (16.8 ± 4.3) and NSC (16.4 ± 6.4).

**PHD2-sshRNA and antimiR-210 treatments enhance neovascularization and cellular proliferation**

We then looked at how each treatment, both alone and in combination, affected neovascularization with CD31 staining. Histological samples obtained at day 4 after the start of treatment showed enhanced neovascularization, with the combination treatment showing the greatest enhancement, over controls (Fig. 5A, red staining). Samples were also stained for Ki-67, a marker for proliferation. Consistent with accelerated closure of wounds receiving oligonucleotide treatment, there was an increase in Ki-67 staining (Fig. 5B, red staining). A significant increase in neovascularization was also observed at day 7 by vWF staining (Supplementary Fig. S6) for SG404-treated wounds over NSC and untreated control wounds in a separate experiment in which only the effect of SG404 was examined.

**Molecular analysis**

Analysis of RNA levels in total RNA isolated from wound tissue harvested day 4 after start of treatment showed a reduction in miR-210 levels only in treatment groups in which SG608 was applied, which also confirms delivery of the antimiR to the cells in the wound area (Fig. 5C).

Analysis of VEGF and SDF-1 levels in wound tissue harvested on day 2 showed a significant (p < 0.05) increase in the group treated with SG404 alone (Supplementary Fig. S7), as expected, based on the induction of VEGF by HIF-1α (Fig. 1). Tissue harvested on day 4 (Fig. 5D) showed a similar result for SG404 alone. However, the two groups treated with SG608 (alone or combined with SG404) showed no change in VEGF levels over controls, consistent with the angiogenesis-promoting effects of miR-210, which at this time point apparently counter the VEGF-inducing effects of SG404.

**Discussion**

Chronic diabetic wounds, particularly pressure ulcers, affect 2.5 million people in the United States and cost the healthcare system an estimated $11.6 billion annually. While interventions exist, current treatments are only moderately effective. Critical cellular and molecular pathways responsible for normal wound healing, including the
hypoxia response regulated by HIF-1α, are impaired in diabetes.\textsuperscript{50–60} Diabetes-related defects have lower levels of signaling that impair neovascularization, fibroblast function, and wound healing.\textsuperscript{12,50,51,61–68} The effectiveness of introducing growth factors has been limited, in part, by the presence of high levels of proteases in the wound environment. Unlike proteins, RNAs can be chemically stabilized against degradative enzymes without compromising their functional activity. Thus, they represent a promising class of targeted therapeutic agents for this challenging condition.

In this study, we examined the use of chemically modified RNAs as therapeutic agents for treating chronic diabetic wounds by targeting the HIF-1α wound healing pathway in db/db mice. In particular, SG404, a chemically modified sshRNA targeting PHD2, and SG608, a modified anti-miR-210, were each shown to significantly accelerate wound closure in full-thickness punch wounds, alone and in combination. A key element in the treatment was the use of LbL technology to deliver high local concentrations of therapeutic RNA.

Several lines of evidence presented here support the modes of action as outlined in Figure 1, with SG404 inhibiting PHD2 and stabilizing HIF-1α, and SG608 inhibiting miR-210 and abrogating its negative effects on keratinocyte growth and mobility. First, irrelevant sshRNAs of the same structural design as SG404 have little or no effect (Figs. 3A and 4C, E). Second, SG404 was shown to knock down PHD2 and upregulate levels of HIF-1α in keratinocytes, consistent with the role of PHD2 in destabilizing HIF-1α (Fig. 3B and Supplementary Fig. S2B). Similarly, SG608 was shown to derepress targets of miR-210 in vitro (Fig. 2C) and knock down levels of miR-210 in the wound margins in vivo (Fig. 5C). Third, factors downstream of HIF-1α that are involved in wound healing (VEGF, SDF-1) are upregulated in the wound margins treated with SG404 alone (Fig. 5D and Supplementary Fig. S7). Fourth, transfection of SG404 and SG608 into HaCaT cells results in increased growth and mobilization in scratch assays (Fig. 3B and Supplementary Fig. S2B). Finally, staining for vWF and CD31 shows increased angiogenesis at days 4 and 7 and staining for Ki-67 shows increased proliferation at day 4 post-treatment initiation (Fig. 5 and Supplementary Fig. S6).

The effectiveness of inhibiting miR-210 alone suggests that this miRNA may play a significant role in chronic wounds. miR-210 is upregulated on binding of HIF-1α to a hypoxia responsive element (HRE) in the promoter of miR-210,\textsuperscript{69} and hence its levels are elevated when HIF-1α is stabilized. Biswas et al.\textsuperscript{18} reported that levels of miR-210 are elevated in ischemic cutaneous wounds and are associated with impairment of keratinocyte growth and re-epithelialization. In addition, miR-210 is implicated in a number of other processes related to wound healing, including angiogenesis, mitochondrial metabolism, and oxidative stress.\textsuperscript{2} While its role in promoting angiogenesis is consistent with a positive role in wound healing, the other processes it promotes tend to be detrimental. This is particularly true in the case of keratinocyte proliferation, which miR-210 inhibits by targeting E2F3 and thereby blocking entry into S phase and initiation of DNA replication. The suppression by miR-210 of mitochondrial respiration, while allowing cells to survive longer under hypoxic conditions, reduces available adenosine triphosphate (ATP), which is required in abundance for wound healing.

**FIG. 5.** Neovascularization and cell proliferation are improved on inhibition of PHD2 and miR-210 in diabetic murine wounds. (A) Images taken on day 4 of treatment show improved neovascularization at the wound periphery. The combination treatment showed greatest improvement compared to untreated and NSCs. Red: CD-31, blue: DAPI. (B) Increased cell proliferation was observed in the SG404 and SG608 groups. Red: Ki-67, blue: DAPI. (C) Verification of specific knockdown of miR-210 by SG608 in tissue isolated at day 4. (D) Enhancement of VEGF protein levels (ELISA) with PHD2 knockdown (SG404) at day 4. ELISA, enzyme-linked immunosorbert assay; VEGF, vascular endothelial growth factor.
Our results support the concept that inhibition of miR-210 and hence reduction of the above detrimental influences of miR-210 on wound healing more than compensate for the proangiogenic effects of this miRNA. Indeed, moderate impairment of angiogenesis alone does not necessarily retard normal wound healing. The observation that anti-miR-210 appears to counter the upregulation of VEGF seen on knockdown of PHD2 at day 4 is interesting in light of the increased extent of neovascularization seen at this time point (Fig. 5D). It may be that other proangiogenic influences make up for the reduction in VEGF, or that VEGF acts at an earlier time point (Supplementary Fig. S7).

The method of RNA delivery used in this study, an LbL coating of RNA on Tegaderm mesh, has important advantages as a practical treatment approach. Electrostatic complexation and the solid-phase nature of the LbL film limit the exposure of nucleic acids to environmental conditions, affording long-term storage stability that is important for stockpiling in clinics, including in resource-challenged settings. Importantly, LbL formulations allow for topical application of therapeutic oligonucleotides, providing high local concentrations spread over several days with very low systemic exposure.

While both RNAi and antisense oligonucleotides have been reported previously for the ability to accelerate wound healing individually, to our knowledge this is the first study to combine both modalities. The efficacy illustrates the potential of combining therapeutic RNAs targeting both mRNA and miRNA in a single formulation with potential superiority to either modality on its own.

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Disclosure Statement


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Supplementary Data

Supplementary Methods

Preparation of short synthetic hairpin RNAs and antimiR oligonucleotides

Short synthetic hairpin RNAs (sshRNAs) were chemically synthesized and high-performance liquid chromatography purified either by Integrated DNA Technologies (Coralville, IA) or TriLink Biotechnologies (San Diego, CA). sshRNAs were resuspended and annealed to monomeric hairpin conformations, and analyzed by denaturing polyacrylamide gel electrophoresis as described in Dallas et al.1 AntimiR oligonucleotides were resuspended in RNase-free H2O. The sequences of sshRNA SG404 and antimiR SG608 are as follows. SG404: 5'- CUGAUUG GCUCUGAGUUUCUUGAACUCAAGCCCAAUUCAG-3'; SG608: 5' - UCAGCCGCUGUCACACGCA-3'.

Cell culture

Human kidney 293FT (Invitrogen, Carlsbad, CA) and human keratinocyte cell line HaCaT (Robyn Hickerson, TransDerm) were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Cambrex) with 10% fetal bovine serum (Hyclone), 2 mM l-glutamine, and 1 mM sodium pyruvate. Human primary keratinocytes (Normal Human Epidermal Keratinocytes; ATCC) were cultured in Dermal Cell Basal Medium (ATCC) supplemented with Keratinocyte Growth Kit (ATCC). Mouse fibroblast cell line NIH-3T3 (ATCC) was cultured in MEM and 10% fetal calf serum. Human kidney 293FT (Invitrogen, Carlsbad, CA) and Human primary keratinocytes were cultured in DMEM (Cambrex) with 10% fetal bovine serum (Hyclone), 2 mM l-glutamine, and 1 mM sodium pyruvate. Human fetal lung fibroblast line MRC-5 were cultured in Dulbecco's modified Eagle medium (DMEM; Cambrex) with 10% fetal calf serum.

Real-time quantitative polymerase chain reaction of messenger RNA and microRNA

Complementary DNA (cDNA) for messenger RNA (mRNA) analysis was synthesized from 10 μL of total RNA samples using the High-Capacity cDNA Kit (Applied Biosystems). cDNA for microRNA (miRNA) quantification was reverse transcribed using the TaqMan-MicroRNA Reverse Transcription Kit (Applied Biosystems). Real-time quantitative polymerase chain reaction (RT-qPCR) amplification was performed using 5×HOT FIREPol Probe qPCR Mix Plus (ROX; Solis BioDyne), and the appropriate TaqMan probes (Applied Biosystems); human prolyl hydroxylase domain-containing protein 2 (PHD2; Hs00254392_m1), human GAPDH (Hs99999905_m1), mouse PHD2 (Mm00459770_m1), mouse GAPDH (Mm00475085_m1), miR-210 (00512), and sno-234 (001234).

Western blots

One day before transfection, 293FT cells were seeded at 90,000 cells/well in 48-well plates. sshRNAs (final concentrations 1 or 10 nM) along with 300 ng carrier plasmid pUC19 were transfected with Lipo2K. Cells treated with 500 μM CoCl2 (final concentration) served as a positive control. Forty-eight hours post-transfection, the cells were lysed in 100 μL of 1× LDS lysis buffer (Novex) supplemented with 100 mM DTT and incubated at 95°C for 10 min. Twenty microliters of lysate was loaded on NuPage 4–12% Bis Tris gel (Invitrogen) and resolved at 60 V for 20 min, followed by 1.5 h at 100 V. Resolved proteins were transferred to a nitrocellulose membrane at 30 V for 3 h. The membrane was blocked with 5% nonfat dry milk in TBST overnight at room temperature. The membrane was then stained with hypoxia-inducible factor-1α (HIF-1α) antibody (H1alpha67; Novus Biologicals), followed by staining with goat anti-mouse IgG-AP (Santa Cruz Biotechnologies). Proteins were visualized using an NBT/BCIP system (Promega) and quantified using ImageJ software.

Construction of miR-210 biosensor plasmid (pSG247)

The miR-210 biosensor reporter (pSG247) was prepared by subcloning double-stranded oligonucleotides containing four tandem complementary miR-210-3p binding sites (5’-TACGCCGCTGTCACACGCAG-3’), each separated by a short unrelated 8 bp sequence, as XhoI-Not1-sticky fragments into the 5’-untranslated region of the Renilla luciferase (rLuc) gene of the psiCHECK-2 dual luciferase reporter vector (Promega). This vector also contains constitutively expressed firefly luciferase (fLuc) for transfection normalization.

Supplemental biosensor assay

Triplicate transfections were performed in HaCaT cells with 0.2 ng of pSG247, 50 ng of carrier plasmid pUC19, and miR-210-specific antimiRs (0.3–10 nM) or nonspecific control (NSC) antimiRs over the same concentration range using Lipo2K (Invitrogen). Twenty-four hours post-transfection, CoCl2 was added to a final concentration of 500 μM to stimulate expression of miR-210. At 48 h, cells were lysed in 150 μL of passive lysis buffer (Promega) and 10 μL of lysate was used to measure fLuc and rLuc levels with the Dual-Glo Luciferase Assay (Promega) on a TR 717 Microplate Lumimeter (Applied Biosystems). miR-210 antimiR activity was measured by the derepression of rLuc expression relative to cells not transfected with antimiRs, normalizing to fLuc.

Serum stability assay and electrophoresis

sshRNAs (3.35 μg) were incubated with 10% human serum (Sigma, St Louis, MO) in phosphate-buffered saline at 37°C. An aliquot was taken out at different time points, immediately mixed with 2× gel loading buffer (Ambion, Austin, TX), and stored at -80°C. Gel electrophoresis was performed under denaturing conditions (12% polyacrylamide, 20% formamide, and 8 M urea) and stained with SYBR Gold (Invitrogen).

Cytotoxicity assay

One day before transfection, HaCaT cells were seeded at 15,000 cells/well in a 96-well plate. Transfections were performed in triplicate, using HiPerFect (Qiagen). sshRNAs and antimiRs were transfected at 10, 30, and 100 nM alongside mock transfection controls. One micromolar (in dimethyl
sulfoxide (DMSO)) staurosporine was used as a positive control. Cell viability was measured 48 h post-transfection using the Vybrant MTT Cell Proliferation Assay Kit (Invitrogen) following the manufacturer’s protocol. Absorbance at 570 nm was measured on a Thermomax microplate reader (Molecular Devices), and percent viability was calculated relative to wild-type cells.

**Mapping of RNA target cleavage sites by 5'-rapid amplification of cDNA ends analysis**

NIH-3T3 cells were transfected as described in the main text with SG404 and NSC sshRNA (SG221c). Untransfected cells were an additional negative control. Total RNA was extracted 18 h post-transfection. The mRNA was then subjected to 5'-RACE (rapid amplification of cDNA ends) analysis by first ligating an adapter of sequence 5'-CGAC UGGAGCAGGACACUGACAUUGGACUGAAGGAG UAGAAA-3' to mRNAs at their 5' ends. Ligated RNAs were reverse transcribed using the PHD2-specific primer 5'-CTGGCAACATCATCGCAGGAG-3' and amplified by PCR using primers 5'-ACTGAAGGAGTAGAAAGCCCA-3' and 5'-AGCAGCCAAGAGCAGTCACA-3'. PCR was started with 1 cycle of 94°C for 2 min, 5 cycles of 94°C for 30 s, and 70°C for 30 s, followed by 25 cycles of 94°C for 30 s and 60°C for 30 s and 70°C for 30 s. The PCR products were analyzed on a 2% agarose gel and the band with the predicted length (251 bp) of the cleavage product was excised and sequenced (Retrogen).

**Fluorescence uptake of PHD2-sshRNA**

SG405, an sshRNA of identical sequence and modification pattern as SG404 but additionally containing an AlexaFluor-594 label, was formulated by layer-by-layer (LbL-SG405) and applied to full-thickness excisional wounds on db/db mice. An additional untreated control group was included for comparison. In all groups, n = 6. On day 7 post-treatment application, animals were sacrificed and sections were prepared as described in the main text. Ten histology slides were prepared per wound. Immunofluorescence was monitored with von Willebrand factor (vWF) antibody (No. 7356; EMD Millipore, St. Louis, MO). Three images were captured per slide. Quantification was performed by blinded analysis using two independent observers, measuring fluorescence intensity by ImageJ.

**RNA analysis**

Tissue sections designated for RNA analysis were immediately immersed in RNAlater solution (Invitrogen), incubated overnight at 4°C, and then stored at −80°C until testing. Lysing Matrix “D” beads (1.2 g) (MP Biomedicals, Solon, OH) and 1 mL of QIAZOL (Qiagen) were added and the samples were processed with a FastPrep-24 homogenizer (MP Biomedicals) by four 60-s cycles (6 m·s⁻¹) followed by cooling on ice. Cellular debris was removed by centrifugation at 12,000 g for 2 min. The homogenate was extracted with chloroform and the aqueous layer was used to isolate total RNA using an RNeasy Plus Universal Mini Kit (Qiagen). RT-qPCR for quantification of miRNA was performed as described above.

**Supplementary Reference**

SUPPLEMENTARY FIG. S1. 5’-RACE analysis shows predicted RNA interference (RNAi)-mediated cleavage product of PHD2 mRNA. Two percent agarose/TBE gel showing 5’-RACE products from amplification of total RNA isolated from NIH-3T3 cells transfected with SG404, NSC sshRNA, or wt cells. The predicted band (251 bp) is only detected in SG404-transfected cells and was confirmed by sequencing. cDNA, complementary DNA; mRNA, messenger RNA; NSC, nonspecific control; PHD2, prolyl hydroxylase domain-containing protein 2; RACE, rapid amplification of cDNA ends; sshRNA, short synthetic hairpin RNA; wt, wild type.
SUPPLEMENTARY FIG. S2. (A) Western blot analysis of induction of HIF-1α by PHD2-specific sshRNA SG302 compared with cells not transfected with inhibitor (no inhibitor) or cells transfected with NSC sshRNA. Cells treated with CoCl₂ serve as a positive control analyzed by Western blot. Lamin (loading control). (B) Representative images of the scratch wounds at 0, 24, and 48 h for combination SG302m PHD2-sshRNA+SG608 miR-210 antimiR versus NSC sshRNA and antimiR. HIF-1α, hypoxia-inducible factor-1α.
SUPPLEMENTARY FIG. S3. MTT cell toxicity assay to measure cell viability 48 h after transfection of HaCaT cells with sshRNAs (A) and antimiR-210 (B) at the indicated final concentrations. In (A), percent cell viability is normalized to mock transfection with HiPerFect reagent (HPF only). In (B), percent cell viability is normalized to mock transfection with HiPerFect (no inhibitor). Staurosporine (1 μM) is a positive control for cell toxicity. SG302, PHD2-targeting sshRNA; SG608, N,N-diethyl-4-(4-nitronaphthalen-1-ylazo)-phenyl amine (ZEN)-modified antimiR-210.
SUPPLEMENTARY FIG. S4. Stability of oligonucleotides in human serum. Oligonucleotides were incubated with 10% human serum for various times in PBS at 37°C. At each time point, an aliquot was removed, mixed with 2× gel loading buffer, and immediately stored at −80°C. (A) Analysis of unmodified sshRNA SG402 and 2′-OMe-stabilized sshRNA SG404 by 12% denaturing PAGE (12% polyacrylamide, 20% formamide, and 8 M urea) and staining with SYBR Gold (Invitrogen, Carlsbad, CA). (B) Analysis of SG608 antimiR by 12% denaturing PAGE and staining with methylene blue. PAGE, polyacrylamide gel electrophoresis; PBS, phosphate-buffered saline.
SUPPLEMENTARY FIG. S5. Representative images of (A) H&E and (B) trichrome staining of fully healed wounds from all treatment groups. Images on the left are histology sections magnified at 5×. Images on the right are magnifications of the corresponding boxes from the 5× images (scale bar = 50 μm). The area corresponding to the healed wound lies in between the two vertical lines in the 5× images. H&E, hematoxylin and eosin.
SUPPLEMENTARY FIG. S5.  (Continued).
SUPPLEMENTARY FIG. S6. SG404 treatment leads to a sequence-specific increase in neovascularization. (A) Representative images of vWF staining at day 7. Green, vWF staining; blue, DAPI. Arrows denote in-focus examples of blood vessels. (B) Quantification of vWF fluorescence. *p < 0.05. vWF, von Willebrand factor.
SUPPLEMENTARY FIG. S7. Downstream factors that enhance wound healing are induced by day 2 after treatment with LbL-SG404. ELISA measuring protein levels for SDF-1 and VEGF comparing untreated, NSC, and LbL-SG404 ($n = 5$ for all groups) from tissue harvested on day 2 after treatment application. *$p < 0.05$ versus NSC. ELISA, enzyme-linked immunosorbent assay; LbL, layer-by-layer; SDF-1, stromal cell-derived factor 1; VEGF, vascular endothelial growth factor.