Fidgetin-Like 2 siRNA Enhances the Wound Healing Capability of a Surfactant Polymer Dressing

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Microtubules (MTs) are intracellular polymers that provide structure to the cell, serve as railways for intracellular transport, and regulate many cellular activities, including cell migration. The dynamicity and function of the MT cytoskeleton are determined in large part by its regulatory proteins, including the recently discovered MT severing enzyme Fidgetin-like 2 (FL2). Downregulation of FL2 expression with small interfering RNA (siRNA) results in a more than twofold increase in cell migration rate in vitro as well as translates into improved wound-healing outcomes in vivo mouse models. Here we utilized a commercially available surfactant polymer dressing (SPD) as a vehicle to deliver FL2 siRNA. To this end we incorporated collagen microparticles containing FL2 siRNA into SPD (SPD-FL2-siRNA) for direct application to the injury site. Topical application of SPD-FL2 siRNA to murine models of full-thickness excision wounds and full-thickness burn wounds resulted in significant improvements in the rate and quality of wound healing, as measured clinically and histologically, compared with controls. Wound healing occurred more rapidly and with high fidelity, resulting in properly organized collagen substructure. Taken together, these findings indicate that the incorporation of FL2 siRNA into existing treatment options is a promising avenue to improve wound outcomes.

Keywords: Fidgetin-like 2, siRNA, wound healing, re-epithelialization, regeneration, microtubules

INTRODUCTION

Wound healing is a complex physiological process that, when not effectively completed, can lead to serious complications such as chronic wounds, infections, and ultimately amputations.1 Current approaches in acute wound healing are generally limited to passive dressings, skin-like matrices, and growth factor–based approaches that address only the initial, inflammatory phase of the healing process.2–5 One area undergoing significant development is the use of poloxamer-based hydrogels called surfactant polymer dressings
These physiologically amenable polymers can repair damaged cell membranes, effectively encapsulate small interfering RNA (siRNA), and increase hydration of the wound. Furthermore, these SPDs are clinically useful as they remain at the site of application and restrict drug delivery to the desired areas. Identification and incorporation of a therapeutically useful siRNA target into an SPD could be used to further improve wound outcomes.

Our previous work that involved screening of biological agents for cell migration phenotypes identified a novel regulator of the microtubule (MT) cytoskeleton, termed Fidgetin-like 2 (FL2), as an important negative regulator of cell migration. siRNA-mediated depletion of FL2 results in a significant increase in the velocity and directionality of keratinocyte and fibroblast motility in vitro. Moreover, we found that in vivo depletion of FL2 strongly promoted wound closure and repair. Of note, FL2 depletion enhanced healing by inducing a targeted, transient disruption of cortical MT within the leading edge of fibroblasts and keratinocytes. This localized alteration bypasses the manipulation of larger signaling cascades, which have been shown to induce numerous unintended side effects. Having identified FL2 as a promising therapeutic target we began development of an SPD containing FL2 siRNA.

As poloxamer technology has matured, commercially available SPDs are more widely used. We identified one biocompatible and nonionic SPD containing poloxamer 188 as a candidate for siRNA incorporation. This SPD has previously been shown to improve wound-healing outcomes (improved wound closure rates, reduction of inflammation, pain, and odor) in nonhealing wounds and is now in a U.S.-based clinical trial to determine its efficacy in the treatment of wounds of mixed etiologies. Looking to utilize the complimentary healing mechanisms of FL2 siRNA and SPD, we synthesized collagen microparticles containing FL2 siRNA and incorporated these into the SPD for topical application in two in vivo wound-healing models.

Here, we demonstrate that the incorporation of FL2 siRNA collagen microparticles into SPD (SPD-FL2-siRNA) significantly enhances healing outcomes compared with controls (SPD without microparticles and SPD-Control-siRNA), as measured clinically and histologically in two murine wound-healing models, namely full-thickness excision and full-thickness burn. Following SPD-FL2-siRNA treatment, there was a decrease in inflammation and an increase in revascularization at the wound site, while cell proliferation was unaffected. Wound healing occurred more rapidly and with high fidelity, resulting in properly organized collagen substructure. Taken together, these findings indicate that the incorporation of FL2 siRNA into existing treatment options is a promising avenue to improve wound management.

**CLINICAL PROBLEM ADDRESSED**

Slow healing and open wounds are a serious medical problem faced by patients worldwide. Without proper re-epithelialization, these wounds are at risk of infection, increasing the risk of amputation or mortality. Slow-healing wounds also bring the risk of scarring and skin deformation. A therapeutic that can more quickly close wounds would directly address these issues.

**MATERIALS AND METHODS**

**Surfactant polymer dressing**

A nonionic SPD (PluroGel®, Medline Industries, Inc.) was used for siRNA delivery.

**siRNA**

Sequences targeting mouse FL2 messenger RNA (mRNA) (SASI_Mm02_00354635; Sigma-Aldrich) or a nontargeting negative control siRNA (SIC001; Sigma-Aldrich) were used.

**Collagen microparticle and SPD incorporation**

Ten grams of sodium bis(2-ethylhexyl) sulfosuccinate (AOT; Sigma-Aldrich) was dissolved in 34 mL n-hexane, and 2 mL 5% collagen-I dissolved in acetic acid was added. The resulting micro-emulsion was stirred for 45 min until it became clear. This solution was then evaporated to remove hexane. The residue was washed and then suspended in nuclease-free water and lyophilized. Then 100 mg of lyophilized powder was then treated with 1,000 μL of 25 μM siRNA solution and re-lyophilized. This material was then suspended in 1.25 mL SPD, at 4°C for 2 h, and then lyophilized. The lyophilized powder was then added with 1.25 mL nuclease-free water and 1.25 mL SPD.

**In vivo studies**

Animal experiments were performed according to the guidelines published by the Institute of Laboratory Animal Resources of National Research Council, and animal care for this study was approved by the Institutional Animal Care and Use Committee of the Albert Einstein College of Medicine.

**Punch biopsy excision**

Before procedures, mice were anesthetized with a ketamine–xylazine cocktail. Female Balb/c mice...
(6–8 weeks; National Cancer Institute, Frederick, MD) were shaved and then wounded uniformly on their dorsa using a 5 mm punch biopsy tool. To counteract wound contraction typically seen in mice, we used an established silicone splinting method to encourage the formation of granulation tissue and re-epithelialization. Splints were checked multiple times a day during the study and replaced as needed. Wounds were treated with 10 μL of SPD, SPD-Control-siRNA, or SPD-FL2-siRNA on days 0, 2, and 4.

**Burn protocol**

Before procedures, mice were anesthetized with a ketamine–xylazine cocktail. Female Balb/c mice (6–8 weeks; National Cancer Institute, Frederick, MD) were shaved and then burned uniformly on their dorsa using a 5 mm brass probe heated to 100°C. Mice were provided buprenorphine as needed following injury. Wounds were treated with 10 μL of SPD, SPD-Control-siRNA, or SPD-FL2-siRNA on days 0, 2, and 4.

**Wound measurements**

Wounds were measured by investigators blinded to the treatment groups using surgical calipers. Measurements were made in four planes (N-S, W-E, NE-SW, NW-SE), tabulated, and compared with day 0. Histologically, wound edges were defined by the distance between the first hair follicle encountered at each end of the wound.

**Histochemistry**

Wound sites were harvested, fixed overnight in HistoChoice, bisected through the center of the wound, and embedded in paraffin blocks. Seven-micrometer sections were cut and slides were deparaffinized, washed in phosphate buffered saline (PBS)-0.01% Triton X100, unmasked with antigen retrieval buffer tris(hydroxymethyl)aminomethane/ethylene diaminetetraacetic acid pH 9.0, and blocked with 5% normal goat serum in PBS. Sections were stained overnight with their respective antibody and visualized using a 3,3’-diaminobenzidine (DAB) kit (Vector Labs).

**Microscopy**

Whole wound sections were imaged with a P250 High Capacity Slide Scanner (3D Histech) using brightfield with a 20× air objective, numerical aperture 0.8, and complementary metal–oxide semiconductor (CMOS) color camera (VCC-FC60FR19CL; CIS Corp.). Measurements of wound sizes were made using the CaseViewer analysis software (3D Histech). The size of wounds was determined by measuring the maximum length between wound edges. Quantifications of proliferating cell nuclear antigen (PCNA), cluster of differentiation 45 (CD45), and platelet endothelial cell adhesion molecule-1 (PECAM1) were conducted as described elsewhere. Herovici analysis was conducted following a previously described approach.

**Statistical analysis**

Data are expressed as mean ± standard error of the mean. Differences between treatment groups were calculated using unpaired Student’s Mann–Whitney nonparametric t-tests, or two-way analysis of variances. Significance was set at p < 0.05.

**Commercial antibodies and stains**

Commercially produced antibodies used in immunohistochemistry included PECAM-1 (1:50; ab28364; Abcam), CD45 (1:200; ab10558; Abcam), PCNA (1:1,000; ab152112; Abcam), and Keratin 14 (1:500; 906004; BioLegend). Alternatively, sections were stained with hematoxylin (#3530-32; Ricca Chemical Company) and eosin (#17372-87-1; Acros Organics) or Herovici (#KHER; American MasterTech) following the manufacturer’s suggested protocol.

**Quantitative PCR**

Skin of SPD, SPD-Control-siRNA, and SPD-FL2-siRNA–treated mice was excised at set time points after wounding. The Next Advance Bullet Blender was used to pulverize the tissue, and RNA was extracted with Trizol reagent (15596-026; Invitrogen). Complementary DNA was synthesized the same day using the SuperScript IV First-strand synthesis system (18091050; Invitrogen). Quantitative PCR (qPCR) was performed using Life Technologies Universal Master Mix II (4440040) with FIGNL2 primers (Mm.PT.58.21940655; IDT Technologies). ActB (actin) served as the reference gene (Mm.PT.58.33540333; IDT Technologies). Resulting data were quantified using the comparative 2–ΔΔCt method. The average of control skin wounds was normalized to 1 and used for relative quantification.

**RESULTS**

**SPD-FL2-siRNA knocks down FL2 messenger RNA abundance**

Delivery of FL2-targeting siRNA to the cells within the wound zone was achieved by direct topical application of our therapeutic surfactant polymer containing siRNA microparticles (Fig. 1A). Reduction of FL2 levels was confirmed by qPCR analysis on injured skin 3 days post initial treatment (Fig. 1B). Wounds that received FL2 siRNA had a 51% reduction in FL2 mRNA levels com-
pared with their control counterpart (SPD with collagen microparticles containing nontargeting siRNA). These data indicate that FL2 siRNA was effectively being delivered and was causing degradation of FL2 mRNA transcripts at the site of application.

**SPD-FL2-siRNA expedites excisional wound re-epithelialization**

Following full-thickness excision, significant improvements in wound size were first noted in SPD-FL2-siRNA–treated wounds 3 days post wounding (Fig. 2A, B). On day 4, open wound areas of SPD alone and SPD-Control-siRNA–treated mice (21.8% and 23.7% open, respectively) were nearly twice the size of SPD-FL2-siRNA–treated wounds (11.04% open; n = 22 wounds per group; Fig. 2C). Treatments continued through day 6 at which point wounds became too small to measure.

Histological analysis of wounds confirmed a significant reduction in wound size following SPD-FL2-siRNA treatment; SPD-FL2-siRNA wounds were reduced to about half the length of controls.

**Figure 1.** FL2 levels are reduced in injured skin treated with SPD-FL2-siRNA. (A) Schematic of microparticle siRNA incorporation into SPD. (B) Mice were burned and then treated at days 0 and 2 with the designated agent. Burns were harvested on day 3 and RNA isolated for subsequent qPCR analysis. SPD-FL2-siRNA treatment significantly reduced FL2 mRNA relative to SPD-Control-siRNA treatment. n≥6; nonparametric Mann–Whitney test; *p = 0.026; error bars = SEM. FL2, Fidgetin-like 2; mRNA, messenger RNA; qPCR, quantitative PCR; SEM, standard error of the mean; siRNA, small interfering RNA; SPD, surfactant polymer dressing. Color images are available online.

**Figure 2.** SPD-FL2-siRNA treatment stimulates healing in a mouse full-thickness excision model. (A) Representative time course of healing following the application of surfactant polymer treatment in BALB/c mice. (B, C) Graphs quantifying remaining wound area over time and at day 4. Red bars are SPD treatment; blue bars, SPD-Control-siRNA; green bars, SPD-FL2-siRNA; n = 22 wounds per group; Student’s t-test; ***p≤0.0008; error bars = SEM. Significance confirmed at individual time points and with a two-way ANOVA. ANOVA, analysis of variance. Color images are available online.
(average wound length of 2.5 ± 0.2 mm for SPD, 2.4 ± 0.3 mm for SPD-Control-siRNA, and 1.2 ± 0.3 mm for SPD-FL2-siRNA treatments, respectively; Fig. 3A, B). Furthermore, we noted that in a few cases (2 out of 7), SPD-FL2-siRNA–treated wounds had hair follicles present within the wound zone, as confirmed by keratin 14 staining (Fig. 3A inset). No such structures were observed in controls. These clinical and histological assessments indicate that SPD-FL2-siRNA was more effectively stimulating re-epithelialization and regeneration following full-thickness excision compared with controls.

**SPD-FL2-siRNA improves burn wound healing**

The effect of SPD-FL2-siRNA on wound healing was further tested in a full-thickness burn model. Burns were measured by two investigators blinded to the treatment groups using surgical calipers every day and tracked until closure (Fig. 4A, B). The wound sites enlarged in size for 4 days post injury, increasing up to 215% the size of the original burn area in control groups. SPD-FL2-siRNA–treated burns, however, did not enlarge as dramatically, reaching a maximum of 160% the size of the original burn area. The reduced burn expansion was followed by significantly improved healing from day 4 to 9 compared with SPD and SPD-Control-siRNA–treated burns (Fig. 4B, C). Due to scabbing, burns became too difficult to measure using surgical calipers following day 9.

Histological analysis of wound tissue 14 days post injury showed that SPD-FL2-siRNA–treated burns underwent a significant reduction in burn wound area (SPD and SPD-Control-siRNA were 35.5% and 39.3% larger, respectively; data not shown). All SPD-FL2-siRNA–treated burns had closed completely at day 14, whereas 25% and 30% of SPD and SPD-Control-siRNA–treated wounds, respectively, remained unhealed (data not shown). Furthermore, SPD-FL2-siRNA–treated burns had a significant reduction in epidermal thickness compared with controls, more closely resembling normal skin (Fig. 4D). These clinical and histological assessments indicate that SPD-FL2-siRNA promotes re-epithelialization and regeneration of wound tissue following full-thickness burns.

**SPD-FL2-siRNA improves regeneration of the skin**

We next characterized the quality of regenerated tissue through immunohistochemical staining.

First we looked to define FL2’s role in cell proliferation as an increase in cell proliferation could potentially account for the shortened re-epithelialization time. PCNA staining of burn wound sections revealed that there was no significant difference in the percentage of PCNA-positive cells between the three treatment groups (Fig. 5A, B). These data indicate that FL2 knockdown does not affect cell proliferation during healing and suggests that cell migration is the
likely driver of hastened re-epithelialization compared with controls.

Managing inflammation is an essential process for effective wound healing. To assess the impact of FL2 knockdown on inflammatory cell presence at day 14, sections were stained for CD45, a marker of leukocytes and other immune cells. CD45 staining was present in all three treatment groups. However, we observed that SPD-FL2-siRNA–treated wounds contained fewer distinct clusters of CD45-positive cells within the burn zone than SPD and SPD-Control-siRNA groups (Fig. 5A, C). After day 7, reduction of CD45-positive cells was indicative of a return to baseline conditions.21

Revascularization of the burn zone was quantified by PECAM1 staining. Significantly more vessels per field were present in SPD-FL2-siRNA–treated wounds (99.5 ± 4.3 vessels per field) than SPD and SPD-Control-siRNA treatments (73.5 ± 6.8 and 62.7 ± 7.9 vessels per field, respectively; Fig. 5A, D). This data suggest that FL2 knockdown may enhance angiogenesis in wounded skin to provide oxygen and nutrients to the wound site for use in the rapidly regenerating wound zone.

**SPD-FL2-siRNA affects collagen III to collagen I ratios in burn scars**

In the early stages of wound healing, myofibroblasts deposit collagen III in the wound zone, increasing the ratio of collagen III to collagen I. Collagen III accounts for up to 50% of total collagen in the early stages of wound healing, up from levels in uninjured skin by about 20%.24 As the scar matures, the ratio of collagen III to collagen I decreases to approximate normal levels.

To determine whether FL2 knockdown influences the ratio of collagen III to collagen I during burn healing, burns were analyzed using Herovici stains. Herovici is routinely used to quantify the ratio of collagen I (which stains red) and III (which stains blue) through multispectral imaging. Our results indicate that there were increased levels of mature collagen (reduction in collagen III to collagen I ratio) in SPD-FL2-siRNA–treated burn tissues, which further supports our hypothesis that FL2 knockdown accelerates wound healing.

**DISCUSSION**

Treating wounds with a SPD provides significant hydration to the wound site and increases
antimicrobial efficacy, improving healing outcomes. The data shown here support our hypothesis that the incorporation of FL2 siRNA, a therapeutic with a complimentary healing mechanism, into an SPD further improves wound-healing outcomes. Further development of alternative SPD formulations and optimization of dosing regimens is an area ripe for future investigation.25

Improving current therapeutic options with FL2-targeting siRNA

Several wound-healing models, including dermal (excisional and burns), ocular (abrasions and burns), and 3D co-cultures of keratinocytes and fibroblasts, have underscored the importance of rapid re-epithelialization as a key determinant of effective healing outcomes.26–32 The clinical and histological characterization of treated wounds presented

Figure 5. FL2 knockdown enhances revascularization and reduces inflammation without affecting cell proliferation. (A) Immunohistochemical staining of skin 14 days post burn and treatment using PCNA, PECAM1, or CD45 antibodies, respectively. Central regions of the burns were imaged at 20 × or 40 × magnification and the number of staining events quantified. (B) PCNA staining showed no significant differences between treatment groups; n ≥ 4 wounds per condition, ≥3 fields per wound; Student’s t-test; n.s. = not significant. (C) Significant differences were observed between controls and SPD-FL2-siRNA treatment groups following PECAM1 staining; n ≥ 5; Student’s t-test; **p = 0.0092. (D) Significant differences were observed between controls and SPD-FL2-siRNA treatment groups following CD45 staining; n ≥ 10; Student’s t-test; p = **0.0027; ***p = 0.0003; error bars = SEM. CD45, cluster of differentiation 45; PCNA, proliferating cell nuclear antigen; PECAM1, platelet endothelial cell adhesion molecule-1. Color images are available online.
above indicate that SPD-FL2-siRNA stimulates more efficient re-epithelialization and progresses wounds into subsequent stages of healing (resolution of inflammation) and regeneration (collagen maturation, angiogenesis) more quickly than controls.

The extent of inflammatory response following injury is a first indicator of potential wound outcome. It is noteworthy that SPD-FL2-siRNA-treated wounds show an expedited resolution of inflammation at earlier time points (Fig. 4B) and reduced CD45$^+$ staining at later points (Fig. 5A); there is now evidence that more effective re-epithelialization is correlated with shortened periods of inflammation and that reduced CD45$^+$ levels 7 days post injury is indicative of further progressed wounds. Next, as re-epithelialization continues, leading edge cells generate a “healing” scaffold into the wound zone. This structure creates a path for cells to efficiently migrate to the center of the wound and begin remodeling the extracellular matrix. Our findings here show that FL2 knockdown influences collagen organization within the wound zone, reducing the ratio of collagen III to collagen I in treated wounds (Fig. 6). While increased deposition of collagen III is known to reduce the density of blood vessels within the wound, collagen I is an optimal substrate for endothelial cells, and this may explain the increase in angiogenesis seen in SPD-FL2-siRNA–treated wounds (Fig. 5). Additional cell types such as epidermal stem cells also utilize this newly formed scaffold to migrate into the wound zone. We found that only FL2 siRNA–treated wounds showed stem cell–derived hair follicles within the wound zone, suggesting that these wounds were further progressed in the regeneration process (Fig. 3). Re-epithelialization also impacts local cellular signaling within the wound zone. Increasingly abundant keratinocytes release VEGF, recruiting endothelial cells into the wound zone, and may provide an alternative explanation for the increased angiogenesis seen in SPD-FL2-siRNA–treated wounds. Additional studies will be needed to define these processes more completely.

In conclusion, we present a new SPD-siRNA platform that can be applied topically and used to treat excision wounds and burns. Treatment with SPD-FL2-siRNA depletes FL2 levels, which expedites re-epithelialization and ultimately improves dermal structure compared with treatment with SPD alone. Due to its favorable chemical composition and ease of application, an SPD-FL2-siRNA product could improve the lives of patients suffering from a variety of dermal injuries.

**INNOVATION**

Here we incorporated siRNA into SPD for use on cutaneous burns and excisions. To our knowledge this is the first time siRNA in collagen microparticles has been incorporated into SPD. While previous formulations of particles have been used to encapsulate siRNA, the platform presented herein has several advantages. SPD is biocompatible,
nonionic, hydrating, and able to remain at the site of application. SPD also has noted anti-biofilm properties, helping to ward off infection and allowing wounds to heal. Combining SPD with the synergistic FL2 siRNA allows for the therapeutic use of both healing mechanisms.

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**AUTHOR DISCLOSURE AND GHOSTWRITING**

B.O.R. is an employee of MicroCures, Inc. J.N. is a scientific advisor for MicroCures, Inc. D.J.S. is the founder and Chief Science Officer of MicroCures, Inc. D.J.S. is the owner of U.S. Patent #20130022667 entitled “Fidgetin-like 2 as a target to enhance wound healing,” which has been licensed by MicroCures. No other author has a conflict of interest. No ghostwriters were used to write this article.

**KEY FINDINGS**

- FL2 siRNA can be incorporated into current treatment options to further improve healing outcomes.
- The use of SPD improves the ease of application of FL2-siRNA.
- Collagen microparticles in SPD is a viable platform for testing alternative therapeutic siRNAs in wound-healing contexts.

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Abbreviations and Acronyms

ANOVA = analysis of variance
CD45 = cluster of differentiation 45
FL2 = Fidgetin-like 2
mRNA = messenger RNA
MT = microtubule
PBS = phosphate buffered saline
PCNA = proliferating cell nuclear antigen
PECAM1 = platelet endothelial cell adhesion molecule-1
qPCR = quantitative PCR
SEM = standard error of the mean
siRNA = small interfering RNA
SPD = surfactant polymer dressing