Targeting Keloid Fibroblasts by Inhibition of Hypoxia Signaling

Keywords: Keloid; Abnormal fibroblasts; Metabolic reprogramming; Hypoxia; Glycolysis

Abstract
Keloids are persistent raised scars that are difficult to treat because attempts at removal typically result in recurrence. Keloid fibroblasts are the abnormal cell type responsible for the continuous scar protein deposition in this fibrotic skin disease that involves enhanced TGFβ activity. An effective therapy that can eliminate keloid fibroblasts and promote normal healing is needed. We examined the cellular and molecular differences between keloid and normal skin fibroblasts to identify characteristics that may be targeted for therapy. By limiting serum and/or glucose availability, we found that keloid fibroblasts are sensitive to glucose levels but not to serum withdrawal. Treatment with 2-deoxyglucose, a preclinical drug that blocks glycolytic metabolism, can reduce keloid fibroblast cell size. However, recovery occurs upon drug removal, indicating a cytostatic effect. To eliminate keloids, it would be necessary to induce cell death in order to disrupt the cycle of continuous fibrosis. Thus, we examined the role of HIF-1, a central transcription factor that regulates both glycolytic metabolism and fibrosis, to determine whether blocking its activity in keloid fibroblasts could yield a cytotoxic outcome. Using a chemical inhibitor against HIF-1, we observed a significant reduction in keloid fibroblast numbers. Although small molecule HIF-1 inhibitors have been developed for anemia and cancer therapy, their role in regulating fibrotic genes has led researchers to consider their potential use in the treatment of lung or kidney fibrosis. Since keloid disease may be considered a form of chronic skin fibrosis, reducing HIF-1 activity could provide a therapeutic strategy for keloid treatment.

Introduction
Keloids are dense, enlarged scars that extend beyond the borders of the original injury. They are prevalent in skin of color individuals and are difficult to treat [1] because removal generates another injury and the recurrence of an even larger keloid. Since they only occur in humans, animal models have not been sufficient to study this disease [1]. The keloid fibroblast (KF) is the abnormal cell type responsible for producing and depositing an excessive amount of matrix proteins into the scar. Research with KFs has identified enhanced TGFβ signaling, as well as features shared with cancer cells [2], such as a dependence on glucose for energy metabolism [3], elevated mTORC1 activation [4,5], and increased transcriptional activity of HIF-1 (Hypoxic Response Factor-1; [6,7]). In this study, we determined cell viability under conditions of limited glucose and/or serum [8], or upon treatment with inhibitors of glycolysis and/or hypoxia signaling. We found that treatment with a HIF-1a inhibitor effectively reduced KF viability, suggesting that targeting the hypoxia signaling pathway could provide a novel strategy that can eliminate pathological KF cells and promote normal healing of keloid scars.

Abbreviations
KF: Keloid Fibroblast; NF: Normal Fibroblast; HIF-1: Hypoxia Responsive Factor-1; 2-DG: 2-Deoxyglucose; qPCR: quantitative Polymerase Chain Reaction; TGFβ: Transforming Growth Factor-beta; mTORC1: mechanistic Target of Rapamycin Complex 1; ANCOVA: Analysis of Co-Variance

Materials and Methods

Fibroblast assays
Keloid fibroblasts (CRL1762) and race-matched normal fibroblasts (CRL2439) were purchased from the American Type Culture Collection (ATCC; 217 Perry Parkway Gaithersburg, MD 20877, USA) and cultured in DMEM (Gibco cat. no. 11965-065; Life Technologies Limited 3 Fountain Drive Paisley, PA49RF, UK) containing 10% FBS, 1:500 amphotericin B, and 1:100 penicillin/streptomycin (Gibco cat. no. 26140-079, 15290-026, 15140-122; Life Technologies Corporation 3175 Staley Road Grand Island, NY 14072, USA) as described [9]. Cell viability was determined by hemocytometry and Trypan Blue dye exclusion. Equal cell numbers were plated at day 0, then treated under 8 distinct conditions. Chemical inhibitors used were: 2-deoxyglucose (2-DG; Sigma-Aldrich cat. no. D6134; 3050 Spruce Street St. Louis, MO 63103, USA) at 10 mM for 2-16 h, or 40 µM KC2F7 for 16 h, NVP-AEW541, MLN117 and BEZ235 (Selleck Chemicals cat. no. S7946; 14408; no. S1034, S8581, S1009; W Sylvanfield Drive Houston, TX 77014, USA). For immunofluorescence, cells were fixed, permeabilized and stained with an anti-SMA or anti-fibronectin antibody (Sigma-Aldrich/MillporeSigma, cat. no. A5228, MAB1926; 400 Summit Drive, Burlington, MA 1803, USA), Alexa Fluor 594 phalloidin, Alexa Fluor 488 phallidin, Hoechst-33342 (Life Technologies cat. no. A12349, A12381, H1399; 29851 Willow Creek Road Eugene, OR 97402, USA), and/or DAPI (Molecular Probes, Inc. cat. no. 62248; 3747 N Meridian Road Rockford, IL 61101, USA). For live imaging, propidium iodide (BioSure, Inc. cat. no. 1032, 1050 Whispering Pines Lane, STE F, Grass Valley, CA 95945, USA) is used to stain dying cells and Hoechst-33342 is used to stain nuclei. Western blot analysis was performed as previously described [9]. Twenty micrograms of cell lysates per lane was used in standard western analysis. Blots were incubated with the indicated primary antibodies at 4 °C, overnight, the appropriate secondary horseradish peroxidase (HRP)-conjugated IgG (Life Technologies) used at 1:4000 in TBST for 1 h at RT. Immunoreactive protein bands were detected using Western

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blot luminol reagent (Bio-Rad cat. no. P10026378/ P10026379, 1000 Alfred Nobel Drive Hercules, CA, USA). Primary antibodies were: phospho-SMAD2 (pSMAD2) and phospho-S6 (pS6) (Cell Signaling Technology, Inc. cat. no. 18338T and no. 4857S, respectively; 3 Trask Lane, Danvers, MA 01923, USA).

**Statistical analysis**

The results were expressed as the mean ± Standard Deviation (SD). Statistical analyses were performed using the SPSS 22.0 software, one-way ANCOVA for group comparisons or a paired T-test for pairwise comparisons. A value of p ≤ 0.05 was considered statistically significant.

**Results and Discussion**

Keloids are often compared with cancers in terms of their chronic and persistent growth. Some of these features are preserved in keloid fibroblasts (KF); therefore, with the absence of an animal model, fibroblasts derived from keloid scars provide an important cell-based model that retains key features of keloids. Of the 10 hallmarks of cancer, keloids share 6 of them [2] (Table 1). To validate some of these characteristics [1], we used western and immunofluorescent analyses to examine the expression of relevant proteins and activated pathways in keloid fibroblasts (KF), but not in normal skin fibroblasts (NFs; Figure 1A). KFs typically express high levels of Smooth Muscle Actin (SMA), a recognized property of the transitional myofibroblast [10]. As reported by others [11], we also noted expression of TGF-β and detection of phospho-SMAD2 (pSMAD2) indicating chronic pathway activation in unstimulated KFs. In addition, mTORC1 (mammalian or mechanistic target of rapamycin complex 1) activity is also elevated in KFs, as shown by phospho-S6 (pS6) detection (Figure 1A). To demonstrate specificity of these activated signals, we also used IGF1 stimulation, either alone or in the presence of highly selective chemical inhibitors to block the signaling pathway at several nodes: IGF1 receptor (using NVP-AEW541), PI3Kα (phosphoinositide 3-kinase-alpha, using the PI3Kα inhibitor MLN1117), or both PI3K and mTORC1 (using the dual inhibitor NVP-BEZ235 [12,13]). In addition, we also observed an enlargement in KF cell size, in accord with enhanced mTORC1 activity, shown by immunofluorescent staining of SMA [5] (Figure 1B). Although the expression of some proteins is highly elevated, others are not. For example, fibronectin expression is similar in both types of fibroblasts (Figure1A and 1B).

Previous studies have shown a preference for glycolytic metabolism in KFs, where chemical inhibitors against this pathway were used to evaluate their potential use in keloid therapy [3,14]. In a comprehensive study by Vincent et al. (2008); [3]) changes in lactate or ATP production were reduced when glycolysis was impaired by 3 structurally distinct inhibitors. In another study, Li et al. (2018; [14]), 2-Deoxyglucose (2-DG) was shown to reduce KF viability over 4 days. Although both studies suggested that chemical glycolysis blockade could be targeted in treating keloids, it is not known whether KFs can be selectively inhibited while allowing NFs to survive.

To examine this further, we investigated whether limiting glucose availability could reveal a difference in viability between NF and KF. Thus, we determined live cell numbers when these fibroblasts were grown in media with reduced glucose and/or serum, as compared with...
complete media. For each type of fibroblast, at least 300,000 cells per flask were seeded in triplicate on day 0. Cell numbers were counted from days 3-7 using hemocytometry (Figure 1C). To highlight glucose-specific effects, we presented a simplified graph to show that both NF and KF cells were affected at similar rates when glucose levels are reduced to 22% or 0%. Thus, limiting glucose availability or antagonizing enzymes in the glycolysis pathway may not provide an effective means for keloid treatment. This observation was confirmed using an Analysis of Co-Variance (ANCOVA), to show that lowering glucose and/or serum levels did not yield statistical differences between cells (p > 0.05; Figure 1C).

We also examined whether preclinical drugs antagonizing overactivated pathways in KFs could destroy this abnormal cell type (Figure 1, data not shown). These include 2-DG to inhibit glycolysis, MLN-1117 to inhibit PI3K, BEZ235 to antagonize both PI3K and mTORC1, as well as KC7F2 (inhibiting HIF-1α; [15]). We reasoned that antagonizing the HIF-1 signaling pathway would limit the expression of genes for both glycolysis and fibrosis, which would be beneficial in keloid treatment (Figure 2A). HIF-1 targeting has also been considered for the treatment of muscle and organ fibrosis [16-18]. However, KFs are cultured under normoxic conditions; therefore, an abnormal activation of the hypoxia signaling pathway needs to be present in order for inhibitors to work. Elevated HIF-1 activity has been reported in KFs. In the Vincent et al. (2008) study, overactivation of hypoxia signaling in KFs was confirmed as these cells were still able to generate ATP in the presence of 2 hypoxia mimics: desferrioxamine and cobalt chloride [3]. We examined HIF-1α transcript levels using qPCR to find an ~2-fold increase. Using 3 distinct sets of primers, we confirmed HIF-1α overexpression in KFs at approximately ~2-fold, with a range of 1.6 to 2.3-fold, compared with NFs. Although HIF-1α mRNA was not highly overexpressed, HIF-1 regulated genes, GLUT1 (glucose transporter-1) and VEGF-A (vascular endothelial growth factor-A) transcripts, were increased in excess of 5-fold (Figure 2 and Table 2; see supplemental information). Thus, using protein or transcript detection, we have demonstrated increased TGF-β, GLUT1 and VEGF levels in the basal, unstimulated state in KFs. Since overactivation of these pathways can impact fibrosis, metabolism and angiogenesis, antagonizing HIF-1 function could affect several pathways that are abnormally activated in KFs (Figure 2A, 2B).

To determine whether a combined blockade of HIF-1α and glycolysis might be effective in eradicating KFs, we used a HIF-1α inhibitor, KC7F2 [15], either alone or in combination with 2-DG, on KF viability. Within 24 h of treatment, KFs were eliminated. Therefore, we refined conditions to obtain ~50% reduction in KF viability (Figure 3A-3C). When 2-DG was present alone, cell viability was only reduced to 80%. In the combined treatment, KF cell numbers remained similar to ~50%, as observed with KC7F2 treatment alone, suggesting that 2-DG did not contribute to enhancing KF elimination (Figure 3C). Furthermore, vital staining showed that increased cell death, indicated by propidium iodide, occurred in the presence of the KC7F2 HIF-1α inhibitor (Figure 3A). In addition, immunofluorescent analysis demonstrated severe alterations in cell morphology in cells treated with the HIF-1α inhibitor but not with 2-DG (Figure 3B).

HIF-1 is a master regulator that controls essential processes including metabolism, fibrosis, angiogenesis and erythropoiesis (Figures 2A and 3D). Small molecule inhibitors that alter the functional levels of the HIF-1α subunit have been in preclinical and clinical development for cancer treatment and anemia [19,20]. Currently, one of the HIF prolyl hydroxylase inhibitors, FG-4592 (Roxadustat, clinically approved in China in December 2018), has
been successful in clinical phase II and III trials for treating anemia in patients with chronic kidney disease [21]. As more HIF-1α inhibitors become clinically approved, their additional use for keloid disease could be accelerated based on clinical safety data. For keloid disease in particular, these inhibitors can be locally administered through an intralesional injection. Thus, direct targeting of KFs may be achieved without inducing an additional skin injury.

**Conclusion**

Our study provides a novel drug target and treatment strategy for patients with keloid disease. This method may be extended to other skin disorders such as localized scleroderma, hidradenitis suppurativa, nephrogenic fibrosing dermopathy, and eosinophilic fasciitis, where localized fibrotic lesions persist.

**Data Availability**

Data presented in this manuscript is available to the research community without restriction at the time of online publication.

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