Kelch-like protein 42 is a profibrotic ubiquitin E3 ligase involved in systemic sclerosis

Received for publication, December 3, 2019, and in revised form, February 7, 2020. Published, Papers in Press, February 17, 2020, DOI 10.1074/jbc.AC119.012066

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Systemic sclerosis (SSc) is an autoimmune disease that affects over 2.5 million people globally. SSc results in dysfunctional connective tissues with excessive profibrotic signaling, affecting skin, cardiovascular, and particularly lung tissue. Over three-quarters of individuals with SSc develop pulmonary fibrosis within 5 years, the main cause of SSc mortality. No approved medicines to manage lung SSc currently exist. Recent research suggests that profibrotic signaling by transforming growth factor β (TGF-β) is directly tied to SSc. Previous studies have also shown that ubiquitin E3 ligases potently control TGF-β signaling through targeted degradation of key regulatory proteins; however, the roles of these ligases in SSc–TGF-β signaling remain unclear. Here we utilized primary SSc patient lung cells for high-throughput screening of TGF-β signaling via high-content imaging of nuclear translocation of the profibrotic transcription factor SMAD family member 2/3 (SMAD2/3). We screened an RNAi library targeting ubiquitin E3 ligases and observed that knockdown of the E3 ligase Kelch-like protein 42 (KLHL42) impairs TGF-β–dependent profibrotic signaling. KLHL42 knockdown reduced fibrotic tissue production and decreased TGF-β–mediated SMAD activation. Using unbiased ubiquitin proteomics, we identified phosphatase 2 regulatory subunit B’ε (PPP2R5ε) as a KLHL42 substrate. Mechanistic experiments validated ubiquitin-mediated control of PPP2R5ε stability through KLHL42. PPP2R5ε knockdown exacerbated TGF-β–mediated profibrotic signaling, indicating a role of PPP2R5ε in SSc. Our findings indicate that the KLHL42–PPP2R5ε axis controls profibrotic signaling in SSc lung fibroblasts. We propose that future studies could investigate whether chemical inhibition of KLHL42 may ameliorate profibrotic signaling in SSc.

Systemic sclerosis (SSc) is an autoimmune rheumatic disease affecting numerous tissue and organ systems and is characterized by hardening and fibrosis of connective tissues. SSc manifests prominently in the skin through skin thickening or increased inflammation; however, the primary cause of mortality is SSc-associated interstitial lung disease (SSc-ILD) (1, 2). Over three-quarters of SSc patients develop lung fibrosis within 5 years of diagnosis, with a subset developing progressive fibrotic disease (3). SSc-ILD is characterized by decreased respiratory function because of excessive hardening and fibrotic scarring because of extracellular matrix deposition in the lung. In combination with an increased inflammatory response, epithelial injury, and cellular fibrotic transition, fibrous production and collagen deposition in the lung interstitium lead to respiratory failure (4, 5). Although new therapeutic agents are under evaluation, current treatment options that target the underlying pathophysiology are limited (6). Profibrotic cellular signaling pathways are associated with pathological remodeling of the lungs, leading to SSc-ILD, particularly that of the transforming growth factor β (TGF-β) pathway (4). Research has shown TGF-β cytokine and downstream signaling to have stark causal effects on SSc disease models (7).

TGF-β signaling proceeds through cellular recognition of the TGF-β cytokine by receptor complexes and activation of mothers against decapentaplegic homolog (SMAD) transcription factors (8). In canonical TGF-β/SMAD signal transduction, receptor-regulated SMAD2/3 protein is phosphorylated, complexed with the co-SMAD protein SMAD4, and shuttled to the nucleus to promote transcription of pro-fibrotic genes.

The abbreviations used are: SSc, systemic sclerosis; SSc-ILD, systemic sclerosis-associated interstitial lung disease; TUBE, tandem ubiquitin-binding entities; CHX, cycloheximide; KD, knockdown; SADe, mothers against decapentaplegic homolog; esiRNA, Endoribonuclease-prepared siRNA.
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the nucleus, where profibrotic transcription programs are activated (9). Nuclear localization and activating phosphorylation of SMAD transcription factors are essential for their signaling transduction, as numerous studies have shown inhibition of TGF-β signaling upon SMAD inactivation or nuclear exclusion (10–14). SMAD-dependent TGF-β signaling is tightly regulated through a variety of mechanisms, including posttranslational protein degradation pathways (15, 16).

Protein degradation is an evolutionarily conserved process for regulating cellular protein longevity. Dysfunctional or aberrant protein degradation is at the heart of many diseases, including fibrotic signaling in the lung. The major mechanism governing protein degradation is the ubiquitin proteasome system (17). Briefly, the small protein ubiquitin is conjugated to target substrate proteins through an enzymatic cascade, among which the ubiquitin E3 ligase enzyme class plays an essential role in substrate recognition and completion of the ubiquitination process. Ubiquitination is a crucial mechanism to regulate homeostasis in TGF-β signaling, as E3 ligase–mediated degradation of TGF-β receptors and of SMAD2/3 transcription factors helps to dampen signaling (18, 19). However, ubiquitination is associated with pathologic fibrotic signaling; we and other groups have observed ubiquitination proteins affecting multiple facets of fibrotic signaling in the lung, including the TGF-β pathway (15, 20–23). Further, profibrotic E3 ligases have also shown promise as targets for chemical inhibition to reduce deleterious fibrotic signaling (21, 24–26). To uncover new ubiquitin E3 ligases regulators of SSc fibrotic signaling, we utilized a high-throughput imaging system to screen an RNAi library targeting E3 ligases for their effect on SMAD2/3 translocation in SSc lung fibroblasts.

Here we report the development of SSc lung fibroblasts as a screening tool for TGF-β activation based on SMAD2/3 nuclear translocation. We identified the ubiquitin E3 ligase KLHL42 as a profibrotic mediator of TGF-β–SSc fibrotic signaling. KLHL42 was also shown to be a regulator of the stability of the protein phosphatase 2A regulatory subunit PPP2R5e and may regulate fibrotic signaling through the PP2A pathway. This study provides a new model of E3 ligase control for fibrotic signaling in SSc lung disease.

Results
Development of the SMAD2/3 translocation ratio screening assay

The Systemic Sclerosis Center of Research Translation at the University of Pittsburgh works with the University of Pittsburgh Medical Center for collection of tissue and explant samples from SSc patients for research. Through this center, we utilized primary SSc patient lung fibroblasts in culture as the basis for the high-content screening assay. Previous research has demonstrated that SSc cell cultures have increased TGF-β/SMAD signaling and exhibit a strongly profibrotic phenotype, which would make them ideal for RNAi loss-of-function screening (7, 27, 28).

We first sought to validate SSc cells as a proper tool for high-content screening in a 384-well plate format. Immunofluorescence studies of SSc lung fibroblasts (SSc cells) show responsiveness to TGF-β1 stimulation, leading to an increased SMAD2/3 fluorescence signal in the nucleus (Fig. 1A). As a surrogate for overall fibrotic activity, we calculated a translocation ratio metric based on nuclear SMAD2/3 immunostaining relative to cytoplasmic SMAD2/3. As SMAD transcription factors are shuttled to the nucleus for fibrotic signaling, we hypothesized that an increased translocation ratio represented a more fibrotic response. The translocation ratio metric showed robust results through titration of primary and secondary antibodies for both baseline and TGF-β1–stimulated treatments (Fig. 1B). We calculated a Z’ factor of 0.39 for this assay; this signal window proved to be adequate for screening (Fig. 1C) (29). Silencing of an essential protein for the TGF-β/SMAD signaling process would result in a decreased translocation ratio (Fig. 1D). We then proceeded to screen for key ubiquitin E3 ligase modulators.

The E3 ligase KLHL42 affects fibrotic signaling and SMAD activation in SSc

With the validation of SSc fibroblasts as a potential screening tool, we sought to screen an Endoribonuclease-prepared siRNA ( esiRNA) library (Sigma) against ubiquitination proteins for their effect on SMAD2/3 localization. Following siRNA knockdown and TGF-β1 stimulation, we collected and immunostained SSc fibroblasts for SMAD2/3 protein, conducted automated microscopy, and calculated a translocation ratio for each esiRNA (Fig. 2A). We observed that most siRNAs had little effect on the changing translocation ratio relative to control esiRNA, but we observed hits whose silencing reduced the SMAD2/3 translocation ratio (Fig. 2B). Given that knockdown of these targets resulted in a reduced translocation ratio (less nuclear SMAD2/3 protein), we hypothesized that these ubiquitination proteins functioned as profibrotic mediators. Of these hits, we observed Kelch-like protein 42 (KLHL42) to be a relatively undercharacterized E3 ligase. Previous studies have suggested that KLHL42 functions as part of the Cul-3 E3 ligase complex, in which KLHL42 functions as the key substrate-engaging protein to facilitate substrate ubiquitination (30, 31). However, the effect of KLHL42 on fibrotic signaling is unclear. We validated the initial screening data through confocal microscopy, as KLHL42 knockdown resulted in decreased SMAD2 activation upon TGF-β1. Knockdown of the TGF-βR1 receptor was used as a positive control, as loss of a major membrane receptor for TGF-β1 would impede TGF-β1–mediated activation of SMAD signaling (13) (Fig. 2, C and D). KLHL42 knockdown significantly reduced production of fibrotic proteins such as fibronectin (Fig. 2, E and F). Further, we observed, through an immunoblot assay, that KLHL42 knockdown affects SMAD2 activation (Ser465/467 phosphorylation) without changing total SMAD2 protein levels, suggesting that the effect of KLHL42 occurs upstream in the signaling process and not through degradation of the SMAD2 protein (Fig. 2, G and H). These initial studies suggest that KLHL42 may be a profibrotic E3 ligase.

Unbiased determination of PPP2R5e as a putative KLHL42 substrate

Ubiquitin E3 ligases primarily exert biological function through targeted ubiquitination and degradation of substrate proteins. To uncover the putative KLHL42 substrate, we con-
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We analyzed the subset of interest for any Gene Ontology terms that would be enriched relative to the total dataset (Fig. 3D), including protein phosphorylation. This suggests that proteins disproportionately represented in the control siRNA group (and potential KLHL42 substrates) may be involved in protein phosphorylation. Next we investigated the set of proteins detected to be directly ubiquitinated and only found in the control siRNA treatment group for their relevance to the enriched gene ontologies (49 proteins). Of these proteins, protein phosphatase 2A regulatory subunit ε (PPP2R5e) was one of the proteins to be detected as directly ubiquitinated and related to the process of protein phosphorylation. PPP2R5e functions as a regulatory subunit for protein phosphatase 2A (PP2A) (35, 36). Intriguingly, PP2A has been implicated in regulation of fibrotic and TGF-β signaling, including SSc patient samples, suggesting that PPP2R5e might function through this pathway in SSc and be a candidate substrate of KLHL42 (37–39).

**Detected PPP2R5e ubiquitination on Lys-84 controls protein stability**

The ubiquitin proteomics assay on SSc fibroblasts detected that PPP2R5e was directly ubiquitinated, with the highest probability on Lys-84 (Fig. 4A). Given the proximity of two other lysine residues to Lys-84, (Lys-86 and Lys-89), there was a chance that the detected ubiquitination was on these Lys sites instead of Lys-84. Initial data analysis used a minimum local-

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**Figure 1. Assay development for scleroderma lung fibroblasts.**

A, immunofluorescence analysis of SSc lung fibroblasts for SMAD2/3 localization without and with short-term TGF-β1 stimulation. B, optimization of assay conditions. Translocation ratios were calculated for titrations of primary and secondary antibody without and with TGF-β1 treatment. C, Z’ factor calculation of the SMAD2/3 translocation ratio with vehicle or TGF-β1 treatment (n = 64 wells/treatment). D, Schematic of the high-content screening assay to measure SMAD translocation upon treatment with ubiquitination RNAi library. SSc cells were treated with the siRNA library in 384-well glass plates prior to TGF-β1 treatment, fixation, and staining for SMAD2/3 and the nucleus. The nucleus was segmented based on nuclear counterstain signal, and the cytosolic region was defined through expansion of the nuclear segment to the threshold of the SMAD2/3 fluorescent signal. The translocation ratio was defined as the ratio of nuclear to cytosolic signal, with a higher signal suggestive of greater SMAD activation. Translocation ratios were calculated with Gen5 software (BioTek) or CellProfiler (49). Scale bar = 200 μm.

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ducted unbiased ubiquitin proteomics MS. We utilized Life-Sensors tandem ubiquitin-binding entities (TUBE) technology to purify ubiquitinated proteins from control or KLHL42 siRNA–treated SSc fibroblasts prior to MS (32, 33) (Fig. 3A). This technology uses multiple ubiquitin-binding moieties linked to resin to affinity-purify polyubiquitinated proteins from the lysate. Our hypothesis was that the putative KLHL42 substrate protein would be less ubiquitinated in KLHL42 siRNA–treated cells (KLHL42-depleted) relative to the control. Less ubiquitinated substrate would prevent its precipitation with the TUBE pulldown; thus, the candidate substrate would be less represented in the proteomics study. The study detected 2486 total proteins, among which 155 were detected to be directly ubiquitinated (Fig. 3B). Among the detected proteins, 291 unique proteins were detected in solely the KLHL42 knockdown, 464 were found only in the control sample, and 1731 were found in both samples (Fig. 3C). We focused our analysis on proteins disproportionately detected in the control siRNA treatment relative to KLHL42 siRNA (464 proteins), as these proteins may have been less ubiquitinated when KLHL42 was silenced.

We analyzed the subset of interest for any Gene Ontology terms that would be enriched relative to the total dataset using the Gene Ontology enrichment analysis and visualization tool (34). We observed that several ontology terms were significantly enriched in the control siRNA treatment group relative to the total dataset (Fig. 3D), including protein phosphorylation. This suggests that proteins disproportionately represented in the control siRNA group (and potential KLHL42 substrates) may be involved in protein phosphorylation. The study detected that PPP2R5e was directly ubiquitinated, with the highest probability on Lys-84 (Fig. 4A). Given the proximity of two other lysine residues to Lys-84, (Lys-86 and Lys-89), there was a chance that the detected ubiquitination was on these Lys sites instead of Lys-84. Initial data analysis used a minimum local-

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![Image](image_url)
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Figure 2. Depletion of ubiquitin E3 ligase KLHL42 impairs TGF-β1–mediated fibrotic signaling. A, screening of SSc fibroblasts with an esiRNA library targeted against ubiquitination proteins and E3 ligases by their effect on the SMAD2/3 translocation ratio. B, KLHL42 is a top hit from the SSc fibroblast translocation ratio screen. C, quantitative PCR analysis of KLHL42 knockdown by RNAi in SSc cells. D. knockdown of KLHL42 impairs SMAD2 nuclear translocation upon TGF-β1 treatment. E, immunoblot analysis of fibrotic products following KLHL42 knockdown and prolonged TGF-β1 treatment. F, quantification of the fibronectin signal from D. G. depletion of ubiquitin E3 ligase KLHL42 affects sclerotic signaling through PPP2R5e.

KLHL42 facilitates PPP2R5e polyubiquitination and degradation

Finally, to validate PPP2R5e as a bona fide substrate of KLHL42, we investigated the mechanism of KLHL42. Endogenous PPP2R5e immunoprecipitated from KLHL42 siRNA–treated SSc fibroblasts showed less ubiquitination relative to the control (Fig. 5A). As an orthogonal approach, we ectopically expressed KLHL42-HA with His-tagged PPP2R5e prior to His pulldown and ubiquitin blotting. We observed that KLHL42 coexpression enhanced the polyubiquitin signal detected upon PPP2R5e precipitation (Fig. 5B). As substrate polyubiquitination often signals for degradation, we probed whether KLHL42 depletion affected PPP2R5e protein levels and observed that KLHL42 knockdown led to a significantly increased PPP2R5e signal relative to the control (Fig. 5C). We utilized other airway cells, BEAS-2B, to ectopically express KLHL42 and observed a significant dose-dependent decrease in PPP2R5e protein levels (Fig. 5D). Further, the critical lysine site PPP2R5e mutant K84R proved to be resistant to coexpression with KLHL42 (Fig. 5F, lane 4), relative to WT PPP2R5e (Fig. 5F, lane 2), suggesting that KLHL42 mediates the protein stability of PPP2R5e through Lys-84. These data suggest that KLHL42 regulates PPP2R5e ubiquitination and protein stability. Finally, we investigated the role of PPP2R5e on fibrotic signaling in SSc cells (Fig. 5G–I). We observed significantly decreased PPP2R5e protein upon siRNA treatment (Fig. 5G and H). PPP2R5e KD resulted in increased fibrotic production upon TGF-β1 treatment, as measured by fibronectin protein relative to control siRNA (Fig. 5G and I). These data suggest a role of the KLHL42–PPP2R5e degradation axis in fibrotic SSc signaling, as KLHL42 functions as a profibrotic mediator degrading the PP2A-enhancing subunit PPP2R5e, resulting in increased fibrotic signaling (Fig. 5F).
Figure 3. The ubiquitin proteomics screen uncovers PPP2R5e as a potential KLHL42 substrate. A, schematic of the ubiquitin proteomics experiment to find KLHL42 substrates. SSc cells were treated with control (Ctrl) or KLHL42 siRNA and with TGFβ1 prior to lysis and precipitation of ubiquitinated proteins with TUBE technology. Precipitated proteins were then subjected to MS analysis. B, total proteins detected from ubiquitin proteomics. 155 unique proteins were detected to be directly ubiquitinated. C, overlap of unique proteins detected between control and KLHL42 siRNA treatments. Proteins in the control KD-only set were considered potential substrates of KLHL42. D, proteins detected in the control KD-only set were queried for significantly enriched Gene Ontology (GO) terms relative to the total dataset; several phosphorylation-related terms were significantly enriched. PPP2R5e was detected in the control KD-only set and directly detected to be ubiquitinated and is a candidate substrcete of KLHL42.

Figure 4. Lys-84 is a candidate ubiquitin acceptor site within PPP2R5e. A, the PPP2R5e peptide identified by ubiquitin proteomics spectrometry, with a high probability of a ubiquitin remnant on Lys-84. B, schematic of detected ubiquitin modifications of PPP2R5e from the posttranslational modifications database (Phosphosite). Potentially modified lysine sites from this study (Lys-84, Lys-86, and Lys-89) are included. C, immunoblot analysis of PPP2R5e-V5 protein and Lys-Arg mutants of suspected Lys sites from this study following a CHX time course. Data are from three to five independent experiments. D, immunoblot analysis of PPP2R5e-V5 protein and Lys-Arg mutants of Lys sites from other studies following a CHX time course. Data are from three independent experiments. *p < 0.05, NS, not significant (p > 0.05) by F test comparisons of nonlinear regression (one-phase decay), with the null hypothesis that the rate constant k = 0 (C and D).
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Figure 5. PPP2R5e is a KLHL42 substrate and affects TGF-β signaling in SSc. A, immunoblot validation of the ubiquitin proteomics assay. PPP2R5e protein was immunoprecipitated (IP) from SSc cells with control or KLHL42 siRNA treatments. WCL, whole-cell lysate. B, immunoblot analysis of the in vivo ubiquitination assay in SSc cells, as PPP2R5e-V5-His protein pull-down was subjected to ubiquitin immunoblotting. C, immunoblot analysis of PPP2R5e protein level following control or KLHL42 siRNA treatments. D, densitometry of C. Data are means ± S.D. of three independent experiments. E, immunoblot analysis of PPP2R5e protein following dose overexpression of the KLHL42-V5 plasmid. Data are means ± S.D. of four independent experiments. F, WT or K84R mutant PPP2R5e-V5 was expressed without or with coexpression of KLHL42-HA prior to blotting. G, SSc cells were treated with PPP2R5e siRNA prior to 18-h TGF-β1 treatment and immunoblotting. H and I, densitometry of PPP2R5e (H) and fibronectin (I) protein without and with PPP2R5e knockdown and 18-h TGF-β1 treatment. Data are means ± S.D. of three independent experiments. J, schematic of the proposed model, in which KLHL42 controls the ubiquitination (Ub) and stability of PPP2R5e. PPP2R5e affects PP2A antifibrotic activity and downstream fibrotic signaling, leading KLHL42 to be a profibrotic regulator in SSc. *, p < 0.05; ***, p < 0.001; two-tailed Student’s t test (D, H, and I). **, p < 0.01 by analysis of variance with Dunnett’s multiple comparisons test (E).

Discussion

This study demonstrated SMAD2/3 immunostaining and automated microscopy as a screening tool for SSc cells. We calculated a translocation ratio of the SMAD signal as a surrogate for TGF-β pathway activity. This assay system is robust and shows good assay parameters with a Z-factor of 0.39. We observed that knockdown of the E3 ligase KLHL42 acutely affects SMAD localization and affects downstream TGF-β/fibrotic signaling, suggesting that KLHL42 is a profibrotic E3 ligase. To uncover potential KLHL42 substrates, we conducted unbiased MS ubiquitin proteomics. Through bioinformatic analysis, PPP2R5e emerged as a putative substrate, and we confirmed Lys-84 as a critical site for PPP2R5e stability through an unbiased proteomics and mutagenesis assay. Finally, we demonstrated that KLHL42 facilitates polyubiquitination of PPP2R5e, leading to its degradation, and that knockdown of PPP2R5e increases fibrotic protein production, potentially through the PP2A pathway.

Phosphatase control of TGF-β signaling is a major mechanism in SSc. PPP2A has been shown to play a direct role in regulating fibrotic signaling in primary SSc disease models. TGF-β treatment was observed to affect PP2A modulation of ERK1/2 activation in SSc fibroblast autocrine signaling (41). Extracellular signal-regulated kinase activation has been directly implicated in dysfunction mechanisms from SSc fibroblasts and has been observed to be constitutively activated in fibroblasts (42, 43). Further, TGF-β treatment of dermal fibroblasts causes a decrease in PP2A transcript/protein levels (38). The regulatory subunit PPP2R5e belongs to the B56/B’ family and modulates overall PP2A activity (35). PPP2R5e expression leads directly to more phosphorylation of PP2A substrates, suggesting that PPP2R5e is an enhancer of PP2A activity (36). This is in line with our functional observations, as we observed that PPP2R5e knockdown led to increased fibrotic signaling. Most research has emphasized the role of PPP2R5e in neoplasia; this is the first study to suggest a function in SSc-ILD fibrotic signaling. More studies are needed to characterize the exact molecular mechanisms of KLHL42 control of PPP2R5e and PP2A signaling.

E3 ligases may play an important role in SSc-ILD signaling. Our previous studies have shown that E3 ligases can potentially control fibrotic signaling through targeted degradation of key signal transduction (24). There is evidence that this paradigm...
can be extended to SSC-ILD fibrotic signaling as well (44, 45). Further, therapeutic targeting of ubiquitination may show promise for lung fibrotic diseases. Recent studies have shown that inhibition of the E3 ligase Skp2 ameliorates fibrotic phenotype in mice (46), and pharmacological inactivation of Cullin-type E3 ligases is protective in experimental pulmonary fibrosis models (21). Current therapeutic options in SSC-ILD are limited, and ubiquitin E3 ligases may prove to be new targets for intervention. In conclusion, we utilized two unbiased approaches (siRNA screen and proteomics) to discover the ubiquitin E3 ligase KLHL42 as a mediator of fibrotic signaling in SSC-ILD through regulation of PPP2R5e ubiquitination and stability.

**Experimental procedures**

**Materials**

PPP2R5e siRNA (HsCD00735088) (47). The QuikChange II XL site-directed mutagenesis kit (200251) was from Agilent. BEAS-2B cells (CRL-9609) were from the ATCC. The anti-HA tag (6E2, 2367S), anti-ubiquitin (P4D1, 3936), anti-phospho-Smad2 (Ser-465/467, 138D4, 3108), and anti-Smad2/3 (D7G7, 8685) were from Cell Signaling Technology. 384-well glass-bottom tissue culture plates (P384-1.5H-N) were from Cellvis. The antibio–antimycotic supplement (15240062), DMEM (10569004), DMEM-F12 (11330057), FBS (26140079), and penicillin–streptomycin (15140122) were from Gibco. Lullaby siRNA transfection reagent (LL7) was from OzBiosciences. hTGF-β1 (100-21) was from PeproTech. Anti-PPP2R5e (NB1-79638) was from Novus. Anti-fibronectin (EP5, sc-8422) and anti-collagen type I (C24, sc-13800) were from Santa Cruz Biotechnology. 384-well glass-bottom plates (Cellvis, 5000 cells/well) and transfection were performed with Mission esiRNA-targeting ubiquitination proteins (Sigma-Aldrich) using XtremeGene siRNA transfection reagent (Roche). Knockdown proceeded for 48–72 h prior to cell collection and analysis. Sequences of dsiRNA used were as follows: KLHL42 siRNA 1, 5’-rCrUUrUrArUrCrArGrUrCrUrCrUrGrArCrArGATT3’ and 5’-rArUrCrUrGrUrCrArGrUrCrArGrUrArArGrArA3; KLHL42 siRNA 2, 5’-rArCrUrUrGrArArUrUrUrArUrUrArUrUrGrUrUrGrUrA-UGT-3’ and 5’-rArCrUrArUrGrArArUrUrUrUrGrUrGrGrArArUrArArUrArUrGrUrGrArGrUrU-3’; PPP2R5e siRNA, 5’-rCrArGrUrCrUrArUrUrGrArUrUrGrArArUrUrGrArGrUrA-UC3’ and 5’-rArGrUrGrUrArUrGrArArUrArUrUrGrArGrUrArGrArArUrArUrArArUrArGrArCrArC-3’; TGF-βR1 siRNA, 5’-rGrUrUrGrUrArUrUrCrUrCrArCrArGrCrArCTT-3’ and 5’-rArGrUrArUrGrUrUrGrGrUrGrArGrArArUrArUrArUrArCrArCrArC-3’.

**Immunoprecipitation and immunoblotting**

Cells were lysed in immunoprecipitation buffer (50 mM Tris-HCl (pH 7.6), 150 mM NaCl, and 0.25% v/v Triton X-100) and centrifuged at 10,000 × g for 10 min at 4 °C. The supernatant was incubated for 2 h at 4 °C with 1:100 dilution of immunoprecipitating antibody. The antibody was captured with protein A/G beads (Thermo Fisher) for an additional 2 h. The immunoprecipitate was washed three times with immunoprecipitation buffer and eluted in 1× Laemmli buffer at 88 °C for 5 min prior to immunoblot analysis. Immunoprecipitated samples were blotted with anti-mouse or anti-rabbit TrueBlot (Rockland).

**Cloning**

Recombinant DNA constructs were prepared through PCR cloning techniques and cloned into the pcDNA3.1D vector (Thermo Fisher) unless otherwise noted. Point mutants were generated with the QuikChange XL 2 site-directed mutagenesis kit (Agilent). All constructs were validated by DNA sequencing (Genewiz).

**Ubiquitination siRNA SMAD translocation screen**

Primary SSC patient lung fibroblast cells were seeded in 384-well glass-bottom plates (Cellvis, 5000 cells/well) and transfected with Mission esiRNA-targeting ubiquitination proteins (E1, E2, E3 ligases etc.) (Sigma-Aldrich) using XtremeGene siRNA transfection reagent (Roche). Knockdown proceeded for 72 h prior to TGF-β1 (Peprotech, 2 ng/ml) stimulation for 1 h. Cells were then fixed (4% paraformaldehyde), permeabilized (0.5% Triton X-100), and stained for endogenous SMAD2/3 (Cell Signaling Technology), followed by goat anti-rabbit Alexa Fluor 488 or 647 (Invitrogen) and nuclear counterstain with Hoechst 33342 (Invitrogen). The fluorescent signal was imaged using Cytation5 (BioTek) or Image Express (Molecular Devices). The SMAD2/3 translocation ratio (nuclear to cytosolic signal) was calculated and analyzed by CellProfiler (49) or Gen5 software (BioTek).

**Ubiquitin proteomics**

Ubiquitin proteomics and TUBE experiments were conducted at LifeSensors. Experimental and control samples were
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analyzed using biological duplicates. Use of biological replicates allowed better accounting for variability with each step of the MS workflow (cell culture, sample preparation, and analysis).

Samples were spun down, and duplicate 22-μl volumes of each sample were mixed with SDS loading buffer. Samples were heated at 90 °C for 5 min and run on SDS-PAGE prior to Coomassie Blue staining. The remaining sample was run on a different gel, resulting in each sample being run on a total of three gel lanes. The lanes were excised, reduced with tris(2-carboxyethyl)phosphine, alkylated with iodoacetamide, and digested with trypsin. Tryptic digests were analyzed using a 150-min LC run on a Thermo Q Exactive HF mass spectrometer. A 30-min blank was run between samples.

Ubiquitin proteomics resulted in four RAW files, two each for KLHL42 siRNA treatment and control siRNA treatment. MaxQuant 1.6.2.3 was used to query MS data against the Uniprot human database (2018_10_01); 196,371 entries were specifically searched. The first search peptide mass tolerance was set at 20 ppm; the main search peptide tolerance was set at 4.5 ppm. Fragment ion mass tolerance was set at 20 ppm. The protein, peptide, and site false discovery rate was set at 1%. Full trypsin specificity was used to generate peptides, with a maximum of three missed cleavages permitted. Carabimidomethyl (C) was set as a fixed modification, with acetyl (protein N-term), oxidation (M), and GlyGly (K) considered variable modifications.

Protein quantification was performed using Razor and unique peptides. Razor peptides are shared (nonunique) peptides assigned to the protein group with the most other peptides (Occam’s razor principle). Quantitation was based on the sum of the peptide MS peak areas for the protein (intensity) because of its increased accuracy compared with the MS/MS count. To account for the fact that larger proteins generate more peptides, the intensity values were adjusted by normalizing against the number of theoretical peptides for each protein (Intensity Based Absolute Quantification (iBAQ) intensity). We then sorted (largest to smallest order) on the iBAQ values to determine which proteins were more abundant in a sample. Data were then analyzed via the Gene Ontology enrichment analysis and visualization tool, querying the 464 proteins identified solely in the control siRNA treatments (and that may have lost their families for generous donation of the tissues used in this study).

Confocal microscopy

SSc cells were seeded in 35-mm MatTek glass-bottom dishes before siRNA knockdown and TGF-β1 treatment. Cells were washed with 1× PBS prior to fixation with 4% paraformaldehyde and permeabilization with 0.5% Triton X-100. Following blocking with 2% BSA in PBS, cells were exposed to SMAD2 primary antibody (1:500) overnight and then 1:1000 Alexa Fluor 488 secondary antibodies for immunostaining. The nucleus was counterstained with Hoechst 33342. Cells were visualized with a Nikon A1 confocal microscope using 405-, 488-, or 567-nm wavelengths. All experiments were done with a ×60 oil differential interference contrast objective lens.

Statistics

All statistical tests were calculating using GraphPad Prism 8. p < 0.05 was used to indicate significance. Densitometry was calculated using ImageJ (National Institutes of Health).

Data availability

Ubiquitin proteomics data have been uploaded through the MassIVE repository, ID: MSV00084800. The MSViewer Data can be accessed using the key axtpvmdao.

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