Identification of regulators of the myofibroblast phenotype of primary dermal fibroblasts from early diffuse systemic sclerosis patients

Loubna Chadli¹, Britt Sotthewes¹, Kejie Li², Stefan N. Andersen¹, Ellen Cahir-McFarland², Marc Cheung¹, Patrick Cullen², Annemarie Dorjée¹, Jeska K. de Vries-Bouwstra³, Tom W. J. Huizinga¹, David F. Fischer¹, Jeroen DeGroot¹, Joanne L. Viney², Timothy S. Zheng², Jamil Aarbiou¹ & Agnes Gardet²

Systemic sclerosis (SSc or scleroderma) is an auto-immune disease characterized by skin fibrosis. While primary cells from patients are considered as a unique resource to better understand human disease biology, the effect of *in vitro* culture on these cells and their evaluation as a platform to identify disease regulators remain poorly characterized. The goal of our studies was to provide insights into the utility of SSc dermal fibroblast primary cells for therapeutic target discovery. The disease phenotypes of freshly isolated and *in vitro* cultured SSc dermal fibroblasts were characterized using whole transcriptome profiling, alpha smooth muscle actin (ASMA) expression and cell impedance. SSc dermal fibroblasts retained most of the molecular disease phenotype upon *in vitro* culture for at least four cell culture passages (approximately 10 cell doublings). We validated an RNA interference high throughput assay that successfully identified genes affecting the myofibroblast phenotype of SSc skin fibroblasts. These genes included *MKL1*, *RHOA* and *LOXL2* that were previously proposed as therapeutic anti-fibrotic target, and *ITGA5*, that has been less studied in fibrosis biology and may be a novel potential modifier of SSc fibroblast biology. Together our results demonstrated the value of carefully-phenotyped SSc dermal fibroblasts as a platform for SSc target and drug discovery.

Systemic sclerosis (SSc) is a chronic auto-immune disease associated with high mortality and high morbidity¹. Initially, the organ most strongly affected is the skin, but the fibrotic process can progress to other organs such as the respiratory system and the gastro-intestinal tract. Both inflammatory and fibrotic processes are thought to contribute to the disease pathogenesis². Among the disease effector cells, activated dermal fibroblasts or myofibroblasts are considered to play a major role in the development of the skin fibrosis, due to their ability to produce large amounts of extracellular matrix and support tissue contractility³. Indeed, analyses of SSc skin biopsies have shown that the increase of myofibroblasts and large collagen fibrils correlates with modified Rodnan Skin Score (mRSS), which is a clinical score to estimate skin thickness⁴,⁵. Whole transcriptome analyses of skin biopsies from SSc patients also highlighted fibroinflammatory signatures; with a prominent fibroblast signature and an increase of the expression of extracellular matrix transcripts⁶–¹⁰. Pathway analysis using these transcriptional data suggested that the TGFβ, PDGF, WNT and the interferon pathways may be involved in the abnormal skin transcriptome of SSc patients¹¹–¹³. Among them, the TGFβ pathway is thought to play a critical role in the disease pathogenesis, as it acts as a potent inducer of fibroblast activation and extracellular matrix production¹⁴,¹⁵.

¹Charles River Nederland BV, Leiden, The Netherlands. ²Biogen, Cambridge, MA, USA. ³Department of Rheumatology, Leiden University Medical Center, Leiden, The Netherlands. Jamil Aarbiou and Agnes Gardet jointly supervised this work. Correspondence and requests for materials should be addressed to J.A. (email: Jamil.Aarbiou@crl.com) or A.G. (email: agnes.gardet@biogen.com)
**Characteristics of the Donors.**

Table 1. Characteristics of the Donors.

| Criteria/Donor                  | SSco1 | SSco2 | SSco3 | SSco4 | SSco5 | SSco6 | SSco7 | SSco8 | HS (n = 11) |
|--------------------------------|-------|-------|-------|-------|-------|-------|-------|-------|-------------|
| Gender                         | F     | F     | F     | M     | M     | F     | F     | F     |             |
| Ethnicity                      | Caucasian | Caucasian | Caucasian | Caucasian | Caucasian | Caucasian | Caucasian | Caucasian | Caucasian (10/11) |
| Patient age                    | 37    | 63    | 81    | 54    | 48    | 54    | 39    | 50    | 47.7±9.5  |
| Disease Duration (months)      | 11    | 14    | 5     | 2     | 4     | 4     | 2     | NA    |             |
| First non-Raynaud symptoms     | Yes   | Yes   | Yes   | Yes   | Yes   | Yes   | Yes   | Yes   | NA         |
| (months prior skin collection) | 11    | 15    | 5     | 7     | 4     | 10    | 4     | 6     | NA         |
| Modified Rodnan skin score     | 7/51  | 31/51 | 13/51 | 29/51 | 37/51 | 14/51 | 21/51 | 18/51 | NA         |
| Auto-antibodies                | PmScl | fibrillarin/U3 RNP | Topoisomerase (Scl-70) | Topoisomerase (Scl-70) | Topoisomerase (Scl-70) | Topoisomerase (Scl-70) | fibrillarin | RNA pol III | NA         |
| Treatment                      | Hydroxychloroquine | Prednisone | Methotrexate | No immuno-suppressive therapy | No immuno-suppressive therapy | No immuno-suppressive therapy | Methotrexate (3 wks) | No immuno-suppressive therapy | Methotrexate (2 months) | Methotrexate (day of biopsy collection) | NA         |

Similar to the findings from studies with SSc skin biopsies, cultured primary SSc dermal fibroblasts also appear to show a profibrotic phenotype at the transcriptional level. They produce increased amounts of extracellular matrix, including collagen 1 and fibrillin, and they express higher levels of SMAD3 and TGF-activating integrins, such as α; 55 and 31, supporting the relevance of the TGFβ pathway for their abnormal phenotype. In addition, a proportion of dermal SSc fibroblasts seem to be differentiated into myofibroblasts, with an increased expression of CTGF and ASMA as well as a more pronounced contractile phenotype, which is thought to contribute to the stiffness of the fibrotic tissue. Besides their profibrotic nature, SSc dermal fibroblasts have also been proposed to contribute to the inflammation in the SSc skin as they secrete higher amounts of IL-6 compared to dermal fibroblasts isolated from healthy donors at baseline or in response to proinflammatory stimulations, such as TNFα or a combination of TNFα and IFN-γ. Together, the discoveries of these abnormal SSc fibroblast phenotypes motivated multiple efforts to identify modifiers of fibroblast biology as an anti-fibrotic strategy, including targeting the TGFβ pathway, the extracellular matrix organization and myofibroblast contractility.

Human primary fibroblasts isolated from various organs are often used to support fibroblast biology studies and multiple potential fibroblast therapeutic targets have emerged from these efforts. Although characterization and functional studies have highlighted the abnormal behavior of SSc dermal fibroblasts, there are still limited reports on how potential disease modifiers may affect primary SSc fibroblast abnormal biology likely due to the difficult access to the primary cells. In this study, we aimed to investigate whether human primary SSc dermal fibroblasts could be used as a platform to support target identification of regulators of SSc disease phenotype and whether they would be suitable for RNA interference comprehensive screens. Thus, we characterized the disease phenotypes of dermal fibroblasts isolated from healthy skin biopsies using transcript profiling, ASMA expression as a myofibroblast marker, and cell impedance as a surrogate for cell morphology differences. Using an adenoviral shRNA platform, we confirmed that a high throughput RNAi screen could be achieved with human primary SSc dermal fibroblasts with adequate robustness and throughput (384-well format). We successfully identified top regulators of ASMA expression in primary SSc dermal fibroblasts among a set of 87 genes enriched for genes relevant to myofibroblast biology. Overall, this study highlights the need to carefully understand and define the disease phenotypes of primary disease cells and how these phenotypes are maintained in an in vitro system in order to assess the value patient primary cells for target discovery and drug discovery.

**Results**

Fibroblasts isolated from SSc skin biopsies retain part of SSc transcriptional signature up to at least four in vitro culture passages. Skin biopsies were obtained from 10 healthy donors and from 6 donors affected by early diffuse SSc from clinically affected or non-affected skin area (Table 1 provides a summary of the characteristics of the donors, Supplementary Table 1 provides the information on what data were collected for each donor). Microarray analyses revealed that skin biopsies from SSc donors showed different transcript profiles than skin biopsies obtained from healthy donors. Principal component analysis confirmed that SSc samples clustered separately from healthy samples (Supplementary Fig. 1A). There were 1178 probes differentially expressed between the SSc skin biopsies and the healthy skin biopsies (Supplementary Fig. 1B and Table 2). Pathway analysis revealed that SSc differentially expressed genes were enriched in genes involved in extracellular matrix organization and immune pathways as well as an interferon signature previously associated with SSc skin (Supplementary Fig. 1C–E). Within the SSc samples, the skin biopsies obtained from disease affected skin area could not clearly be differentiated from the ones obtained from non-affected skin area as shown by the principal
Figure 1. Microarray gene expression analyses of freshly isolated and in vitro cultured primary SSc dermal fibroblasts. Microarray gene expression data from fibroblasts from Passage 0 to Passage 4 from 5 SSc patients (isolated from disease affected skin (SSc_d) or non-disease affected skin (SSc_n)) and 7 healthy donors were analyzed. (A) Principal Component Analysis. (B) Z-score heatmap showing the gene expression profiles of the differentially expressed probes between SSc dermal fibroblasts at P0/P1 and healthy dermal fibroblasts at P0/P1. (C) Overlap of the differentially expressed genes from SSc dermal fibroblasts P0/P1 compared to healthy dermal fibroblasts and from SSc dermal fibroblasts P4 compared to healthy dermal fibroblasts. (D) Quantitative PCR validation data from fibroblasts from at least 5 SSc patients isolated from non-affected skin (orange) or affected skin (red) and 3 healthy donors (black). Statistical significance was assessed using Mann-Whitney test with *p < 0.05 and **p < 0.01. (E) Gene Set Enrichment Analysis of a published in vitro TGFβ gene signature in the SSc signature. Top panel shows a Venn diagram showing the number of unique and overlapping differentially expressed genes. Bottom panel is a plot showing correlation between the log2 fold changes of the union of the differentially expressed genes.

Component analysis (Supplementary Fig. 1A). Only 2 transcripts were detected to be statistically differentially expressed with a lower expression in biopsies obtained from affected site vs non-affected site (HOXB-AS3, HOXB7). This was consistent with the previous studies reporting the challenges of identifying differences at the transcriptional levels between SSc skin biopsies obtained from clinically affected site vs non-affected skin area. Overall, microarray transcriptomic analyses confirmed that the skin biopsies that were used to isolate the SSc primary fibroblasts recapitulated the disease signatures previously described by various groups.

Dermal fibroblasts were isolated from skin biopsies from SSc patients and healthy individuals and cultured up to four cell culture passages, which corresponds approximately to 10 cell doublings from the freshly isolated. Two microarray experiments were conducted to analyze the transcriptomes of these primary cells at different passages. Since the microarray data were collected from 2 different experiments, data were first subjected to batch correction before differential gene expression analyses (Supplementary Fig. 2). The microarray gene expression data could differentiate freshly isolated SSc skin fibroblasts from healthy skin fibroblasts with the identification of 926 differentially expressed unique genes (1365 differentially expressed probes) between SSc skin fibroblasts (passage 0 and passage 1, P0/P1) and healthy skin fibroblasts (Passage 0 and 1, P0/P1) (Fig. 1A–C and Supplementary Table 3). Both principal component analysis and a heatmap showing the gene expression profiles of the SSc fibroblast signature highlight that SSc skin fibroblasts from passage 0 (P0) to passage 4 (P4) showed very similar gene expression profiles (Fig. 1A,B). No transcript passed the 1.5-fold change threshold and FDR-adjusted p-value of less than 0.01 when directly comparing SSc P4 and P0P1 groups (Supplementary Table 4). The SSc skin fibroblasts showed upregulation of the typical fibrotic genes such as components of the extracellular matrix (e.g., collagens, tenascin, decorin, lumican, aggrecan transcripts and metalloproteases). Quantitative PCR data confirmed the upregulation in SSc fibroblasts of several transcripts of genes known to be associated with fibrosis and myofibroblast biology and detected in our microarray analysis: ACTA2 (encoding for ASMA), extracellular matrix associated genes (COL11A1, LOXL2, CTGF), a known myofibroblast transcription factor (GLI2) and a
Fibroblasts isolated from SSc skin biopsies show a higher alpha smooth muscle actin expression level for up to four in vitro culture passages compared to fibroblasts isolated from healthy skin.

Fibroblasts isolated from chronic fibrotic tissues are known to express higher alpha smooth muscle actin (ASMA) levels leading to an increased contractility and cellular morphological change. Using ASMA immunofluorescent staining, we confirmed that overall SSc dermal fibroblasts expressed a higher amount of ASMA protein than healthy dermal fibroblasts for up to four in vitro cell culture passages (Fig. 2A,B). At passage 2, SSc fibroblasts from 7 donors out of 7, either isolated from clinically affected skin or non-affected skin, showed higher ASMA expression levels than healthy dermal fibroblasts (4 donors at passage 2 (Fig. 2C)). At passage 3, SSc fibroblasts from 7 donors out of 7 donors (5 cultures isolated from non-affected area and 3 from affected area) showed higher ASMA expression levels than healthy dermal fibroblasts (Fig. 2C). At passage 4, SSc fibroblasts from 3 donors out of 7 (2 lines isolated from non-affected area and 1 from affected area) showed higher ASMA expression levels than healthy dermal fibroblasts from 4 donors (Fig. 2C). Noteworthy, in vitro culture led to an overall increase of ASMA levels in healthy fibroblasts likely due to the time of culture on plastic cell culture flasks, which present as a matrix with high stiffness (Fig. 2C). Consistent with a higher expression of ASMA and functional effect of cell cytoskeleton, the impedance readout of monolayers of SSc dermal fibroblasts showed statistical difference compared to the impedance measured from healthy dermal fibroblasts after four in vitro cell culture passages (Fig. 2D).

Cell expansion is necessary to obtain sufficient number of cells for functional studies. Together our data showed that expanded SSc dermal fibroblasts up to four in vitro cell culture passages from multiple donors maintained about 77% disease transcriptional signature compared to freshly isolated disease cells, in addition to a high ASMA expression level and abnormal cell impedance. This suggests that SSc dermal fibroblasts may be used as a platform to identify regulators of disease fibroblast phenotypes after a limited cell culture expansion.

Adenovirus delivery of shRNAs in SSc dermal fibroblasts leads to efficient gene expression knockdown and enables to identify regulators of ASMA expression in SSc dermal fibroblasts. Using adenoviral delivery system, shRNAs were transduced into SSc primary fibroblasts. Gene expression knockdown was assessed with quantitative PCR after transduction of 103 shRNAs targeting 31 different genes with 2–4 shRNAs per gene into dermal fibroblasts from 2 different SSc donors. In cells from both donors, approximately 80% of shRNAs led to more than 60% target gene expression knockdown and approximately 65% of shRNAs led to more than 75% target gene expression knockdown (Fig. 3A). This demonstrated that the adenovirus delivery of shRNAs was an efficient RNAi platform for primary SSc dermal fibroblasts. Protein knockdown was also successfully achieved after transduction of shRNAs targeting either ASMA or fibronectin, as assessed using ASMA immunofluorescence staining or EDA-Fibronectin MSD assay (Fig. 3B).

Using ASMA immunofluorescent staining data from SSc dermal fibroblasts isolated from clinically affected or non-affected skin area from 6 different donors and transduced with shRNAs targeting either ASMA or GFP (negative control), we evaluated how well the assay could measure inhibition of ASMA expression. Based on 120 data points for each donor and each shRNA, the difference in ASMA expression level from the ASMA shRNA conditions showed that the assay window was consistently 2–4 fold and the assay Z-prime values were mostly between 0.3–0.6 (Fig. 3C), demonstrating this assay would be robust for high throughput screening to identify inhibitors of ASMA expression in SSc dermal fibroblasts.

Noteworthy, there were two observations suggesting that, although the assay had the potential to identify regulators of the myofibroblast phenotype of SSc cells, the assay conditions did alter the SSc dermal fibroblasts. First, the viral transduction seemed to induce an increase of ASMA protein expression of SSc dermal fibroblasts, but not FN1 (Fig. 3B). Second, an additional transcriptomic analysis using cells at the time of the ASMA assay read-out highlighted that part of the SSc disease signature was affected by the 7-day cell culture condition, which contained low fetal bovine serum (0.5% vs 10% FBS in the expansion culture medium) (Supplementary Fig. 3). Using 2 SSc fibroblast lines and microarray gene expression profiling, we determined that approximately 43% of the differentially expressed genes detected in SSc dermal fibroblasts at P4 were maintained through the 7-day function assay (Supplementary Fig. 3A,B). Although the differential expression of some of the genes may not have passed statistical significance, there was still a statistically significant correlation between the log2 fold changes of the differentially expressed genes from SSc dermal fibroblasts P4 versus healthy fibroblasts and SSc dermal fibroblasts after the 7-day array compared to healthy fibroblasts (Supplementary Fig. 3C). Gene set enrichment analysis confirmed that the published in vitro TGFβ3 signature remained enriched in differentially expressed genes between the SSc dermal fibroblasts P4 after the 7-day assay and the healthy fibroblasts, suggesting that the pathway is still active in the 7-day culture assay (Supplementary Fig. 3D).

First, the viral transduction seemed to induce an increase of ASMA protein expression of SSc dermal fibroblasts, but not FN1 (Fig. 3B). Second, an additional transcriptomic analysis using cells at the time of the ASMA assay read-out highlighted that part of the SSc disease signature was affected by the 7-day cell culture condition, which contained low fetal bovine serum (0.5% vs 10% FBS in the expansion culture medium) (Supplementary Fig. 3). Using 2 SSc fibroblast lines and microarray gene expression profiling, we determined that approximately 43% of the differentially expressed genes detected in SSc dermal fibroblasts at P4 were maintained through the 7-day function assay (Supplementary Fig. 3A,B). Although the differential expression of some of the genes may not have passed statistical significance, there was still a statistically significant correlation between the log2 fold changes of the differentially expressed genes from SSc dermal fibroblasts P4 versus healthy fibroblasts and SSc dermal fibroblasts after the 7-day array compared to healthy fibroblasts (Supplementary Fig. 3C). Gene set enrichment analysis confirmed that the published in vitro TGFβ3 signature remained enriched in differentially expressed genes between the SSc dermal fibroblasts P4 after the 7-day assay and the healthy fibroblasts, suggesting that the pathway is still active in the 7-day culture assay (Supplementary Fig. 3D).
Identification of regulators of ASMA expression in SSc dermal fibroblasts. After we confirmed that the adenovirus shRNA platform in primary SSc dermal fibroblasts could achieve a robust gene knockdown efficiency and allow the identification of inhibitors of ASMA expression, we wanted to provide biological validation of the assay by testing 667 shRNAs targeting a set of 87 genes enriched for genes predicted to regulate myofibroblast biology (Supplementary Table 3). The 667 shRNAs were tested in dermal fibroblasts obtained from two different SSc donors. The level of ASMA expression was measured by immunofluorescent staining and the percentage of inhibition of ASMA expression was reported for each shRNA tested with dermal fibroblasts from the two SSc donors (Fig. 4A). Multiple shRNAs targeting ASMA and MKL1 showed the strongest inhibition of ASMA in dermal fibroblasts from 2 different SSc donors, confirming the ability of the assay to identify strong regulators of the myofibroblast phenotype (Fig. 4A). The percentage of inhibition of ASMA expression for all the shRNAs was averaged from the values obtained from the two SSc donors and ranked based on the percentage of ASMA expression inhibition (Fig. 4B). MKL1 shRNA was identified as a very strong hit with 8 different shRNAs...
yielding to more than 35% inhibition. S100A8, AKT1, EDN1, PALLD, RHOA shRNAs were considered as strong hits with at least 2 different shRNAs leading to more than 35% inhibition. JTGAS, LOXL2, CTHRC1, MAP3K7 were considered as moderate hits with at least 1 shRNA with more than 35% inhibition and 1 shRNA with less than 30% inhibition, but more than 25% inhibition. Finally, CCL18, IRF5, MAPK8, PDGFRα were considered as weak hits with at least 3 shRNAs with more than 25% inhibition, but less than 35% inhibition.

In order to confirm the modulation of the SSc myofibroblast phenotype, we investigated the effect of MKL1 knockdown on the expression of other fibroblast SSc-upregulated genes in our gene signature and associated with fibrosis. We confirmed that 2 different shRNAs targeting MKL1 were able to decrease in SSc fibroblasts the expression of ACTA2 (encoding for ASMA), extracellular matrix associated genes (COL11A1, ACTA2_v1, ACTA2_v2, ACTA2_v3), a known myofibroblast transcription factor (GLI2) and a known profibrotic secreted factor (IGFBBP) (Fig. 4C). Note that, as reported above, the low FBS assay condition affected the SSc transcriptional signature, such that the expression fold changes between the SSc fibroblast cells and healthy cells is lower during the low FBS assay condition than what we measured from the cells after passage 0 and passage 4 in Fig. 1D.

Together, these results demonstrate our RNAi assay with carefully phenotyped primary dermal fibroblasts from SSc donors can be used to identify regulators of the myofibroblast phenotype of SSc dermal fibroblasts.

**Discussion**

Primary dermal fibroblasts from patients affected with skin fibrosis have been considered to be an attractive system to explore the potential of anti-fibrotic drugs. It is commonly thought that these primary fibroblasts could be used up to 10 cell culture passage to avoid senescence effect or to maintain their abnormal collagen production. Our results provide the first comprehensive transcriptomic characterization of freshly isolated vs in vitro expanded primary SSc dermal fibroblasts after approximately four cell culture passages that correspond to approximately 10 cell doublings (30–40 million cells). Fibroblasts were obtained from patients diagnosed with early diffuse cutaneous SSc with less than 18 months since the first non-Raynaud symptom which is thought to be a disease stage where fibrosis disease modifiers could have an impact. Our results highlight that these primary cells maintain about 77% of their transcriptional disease signature for at least four in vitro culture passages. Interestingly, we also confirmed that this disease fibroblast signature is detected in fibroblasts obtained from both clinically affected skin and non-affected skin, consistent with a previous reports of skin biopsy and fibroblast analysis. It is important to note that low serum culture conditions often used in typical fibroblast functional assays may further affect the maintenance of SSc disease fibroblast transcriptional phenotype, as we observed a further drift of the SSc fibroblast signature after a 7-day assay in low serum medium culture condition. Overall, while cultured primary SSc dermal fibroblasts do not retain completely the disease transcriptional signature, they still retain a large part of it as well as additional disease phenotypes, such as a higher expression of ASMA and an abnormal cell impedance. Our data support the recommendation to monitor the SSc fibroblast disease transcriptional signature to confirm that the primary cells used in SSc functional assays retain their disease phenotype. Our results suggest avoiding the use of cells after 10 cell doublings as they start to lose their disease transcriptional signature. Note that our results also highlight that ASMA expression started to increase in healthy fibroblasts at passage 4/5, such that high ASMA expression after these passages may be due to cell culture conditions rather...
than the original disease phenotype. It is possible that this ASMA expression increase is due to the culture on standard tissue culture plastic ware, which provides a high stiffness substrate that is thought to promote myofibroblast differentiation. Together these findings supported the use of low cell passaged and carefully-phenotyped primary cells for functional investigations to identify regulators of the SSc dermal fibroblast disease phenotypes.

Figure 4. Identification of regulators of ASMA expression in primary SSc dermal fibroblasts. (A) Inhibition of ASMA protein expression was determined using ASMA immunostaining after image quantification and normalization with the number of nuclei after transduction of shRNAs into dermal fibroblasts obtained from 2 different SSc donors. ASMA expression level for each shRNA was compared to the ASMA expression level after the negative control GFP shRNA transduction. Plot shows the percentages of ASMA inhibition for the 667 different shRNAs transduced into dermal fibroblasts obtained from 2 different donors. (B) Percentage of ASMA inhibition was averaged between the fibroblasts from 2 different donors and ranked from highest average inhibition to lowest inhibition values. (C) Quantitative PCR gene expression data of SSc upregulated genes after transduction of negative control shRNA or two different MKL1 shRNAs (cells from 2 healthy donors and 1 SSc donor). Statistical significance was assessed using two tailed Student's t-test with *p<0.05, **p<0.01, ***p<0.001 and ****p<0.0001.
One limitation of our study was the limited number of patients that could be enrolled in that study (n = 8), given the low frequency of the disease and that we restricted the study to early diffuse disease. Nevertheless, the transcriptional signature observed in disease fibroblasts was consistent and robust enough such that it could be clearly detected by sample clustering and passed statistical significance (almost 1,000 genes with fold change > 1.5 and FDR-adjusted p-value < 0.01, from cells isolated from 6 SSc patient enrolled at the time of the analysis).

Importantly, the results are similar to previous studies, including the detection of a TGFβ3 signature (Fig. 1E), further providing additional confidence in the data collected. As for functional assays, to our knowledge, this is the first original study describing multiple functional readouts from at least fibroblasts from 5 different SSc patients given the difficulty to enroll these patients and obtain skin biopsies (ASMA expression (n = 7), cell impedance (n = 5), shRNA performance with ASMA (n = 6)). Our results were reproducible enough that they passed statistical significance; however, the robustness of these phenotypes across patients remained to be further observed from future independent studies from other groups, given our relatively limited access to patients.

Using an adeno virus RNAi platform46, we confirmed that primary SSc dermal fibroblasts could be used a system to identify regulators of their myofibroblast phenotype using ASMA expression as a readout. The assay performance was robust with cells obtained from 6 different SSc donors. Beside technical validation, the biological validation of that assay was also achieved as we successfully detected shRNAs directed at genes that are currently considered as promising anti-fibrotic targets among the strongest inhibitors of ASMA expression in dermal fibroblasts obtained from 2 different SSc donors. MKL1 and RHOA are key regulators of the actin cytoskeleton and are being studied intensively as a potential avenue for anti-fibrotic drug targets22,37–39. In particular, MKT- A protein, encoded by MKL1, was shown to be increased in fibroblasts of SSc skin tissue and MKL1 knockdown was known to decrease contractility and Type I collagen production22. Another strong inhibitor of ASMA expression in SSc dermal fibroblasts identified in our assay was LOXL2, which is a lysyl oxidase that promotes the cross-linking of collagen and elastin40. Its role in the regulation of the extracellular matrix has been thought to be critical for the development and maintenance of fibrosis, as LOXL2 inhibitors have shown benefit in liver and lung fibrosis mouse models and serum LOXL2 levels have been associated with poor prognosis in patients suffering from idiopathic pulmonary fibrosis (IPF)41–43. However, a clinical trial that tested an anti-LOXL2 mAb in IPF failed to show efficacy44. Lysyl oxidase activity was shown to be increased in SSc skin and lysyl oxidase serum concentrations correlated with the mRSS clinical score, which evaluates patient's skin thickness45,46. It remains to be investigated whether a strategy to block LOXL2 may be beneficial in SSc skin fibrosis.

Among the 87 genes we tested, it is noteworthy that the shRNAs targeting the SMAD family were not identified as the strongest regulators of the ASMA expression in SSc dermal fibroblasts: 1 SMAD2 shRNA led to more than 35% inhibition of ASMA expression; 1 SMAD3 shRNA and 2 SMAD4 shRNAs led to more than 25%, but less than 35% inhibition of ASMA expression. This was somewhat surprising, given that i) the TGFβ3 pathway is considered to be a major driver of myofibroblast differentiation and ii) we confirmed the enrichment of the TGFβ3 gene signature in the SSc dermal fibroblast transcriptomes. This result may be due to redundant role of SMAD2 and SMAD3 and the fact that our assay did not include the addition of exogenous recombinant TGFβ3, which would have introduced a bias toward the identification of modulators of the TGFβ3 pathway as the strongest hits.

Several shRNAs targeting Irf5 showed a weak inhibitory effect on the myofibroblast phenotype of the primary SSc dermal fibroblasts. Genetic polymorphisms in Irf5 have been associated with multiple auto-immune diseases, including SSc47. Although one may speculate that this result may have been linked to the use of a viral vector to deliver the shRNAs, there are several reports suggesting that Irf5 may be involved in myofibroblast or fibroblast biology: fibroblasts from Irf5 KO mice showed a decrease production of collagen and other ECM components, which may contribute to the protection against disease seen with Irf5 deficiency in the bleomycin skin mouse model and with the Irf5 inhibition in the tight skin mice48,49. Our results combined with these reports suggest that additional studies are needed to confirm the role of Irf5 in the SSc dermal fibroblast disease phenotypes using non-viral inhibitor strategies.

As discussed above, LOXL2, RHOA and MKL1 have been already extensively studied for their role in fibrosis. The integrin α5, encoded by ITGAV5, has not yet been examined closely for its role on skin fibrosis. It is known as a receptor of fibronectin, but has not been shown to promote TGFβ3 activation like some αv integrin heterodimers. The increase α5 expression is associated with poor prognosis of several types of solid tumors and functional studies have demonstrated that α5 promotes invasiveness, metastasis and angiogenesis40. However, volociximab, an anti-αvβ3 blocking antibody, has yet to show efficacy in the treatment of solid tumors40. In the context of fibrosis, there are limited studies exploring the role of α5 integrin biology. In vitro studies have shown that α5 can be upregulated by CTGF and TGFβ3 and that blockade of α5β1 seems to inhibit TGFβ3-induced fibronectin matrix deposition from airway smooth muscle cells51–53. Dioscin, a molecule shown to have anti-fibrotic activity in liver and lung fibrosis mouse models, has been speculated to act through α5 integrin54. Thus, studies with a more specific inhibitor will be needed to definitely establish the role of α5 in the development of fibrosis. Our results combined to the reports linking α5 to fibronectin, TGFβ3 and CTGF biology would warrant further translational and in vivo investigations of the role of α5 in SSc skin fibrosis.

The phenotyping of the SSc dermal fibroblasts combined with the RNAi high throughput assay we described offer a unique opportunity to investigate regulators of SSc fibroblast biology. There are still very limited studies using primary disease patient cells that report the effect of potential SSc fibroblast disease-modifiers. These studies are often independent, which makes comparisons between different SSc fibroblast disease-modifiers difficult, as they are rarely compared directly to each other's in a same study. In addition, there are only a few reports of screens to identify potential anti-fibrotic molecules and these studies used either cell lines or rat primary hepatic stellate cells combined with stimulation with TGFβ3 or other profibrotic agonists22,55–57. Our system is the first successful report of the use of primary human disease fibroblasts for high throughput screening which will allow a comprehensive identification of regulators of dermal fibroblast disease phenotypes and provide the opportunity to compare and rank potential regulators to prioritize further efforts on the most potent pathways.
While we believe our primary screen to be a robust approach to identify regulators of the SSc myofibroblast phenotype, these results relied mainly on the measurement of ASMA expression levels and validation by a set of SSc myofibroblast gene markers. Additional efforts will be needed to confirm the effects of hits on other readouts of SSc myofibroblast biology, such as contractility and a production of extra-cellular matrix components and ultimately disease mouse models. Together our results provide a comprehensive characterization of primary SSc dermal fibroblasts to highlight both the potentials and limitations of this platform to explore SSc fibroblast disease biology. Our study is the first report of the use of phenotypically characterized primary human patients’ cells for identification of modifiers through screening. We developed a robust high throughput RNAi screen that can identify regulators of the myofibroblast phenotype of primary SSc cells, confirming the relevance of known potential targets already considered for skin fibrosis as well as several potential genes of interest that could be the focus of additional investigations.

Patients and Methods
Patient population and skin biopsy collection. The skin biopsies were performed after informed consent and according to European and Dutch regulation, following approval by the medical ethical committee of the Leiden University Medical Center, Leiden, The Netherlands (protocol reference P13.182, NL46199.058.13). Skin biopsies from healthy subjects were collected following informed consent by subjects remaining skin after corrective resection surgery. Eight SSc patients enrolled the study and were diagnosed with diffuse cutaneous SSc with less than 18 months since the first non-Raynaud symptom. Demographics and skin biopsy locations for all healthy or SSc subjects as well as disease characteristics and treatment information for each SSc patients are available in Table 1. Most patients were positive for anti-topoisomerase antibodies and had not received immunosuppressive therapy. When possible, biopsies were taken from different locations of the skin. Skin biopsies were used to isolate primary fibroblasts. Note that all the assays could not be performed for all the fibroblast cultures from all the donors; information on what SSc fibroblasts were evaluated for each assay is reported in Supplementary Table 1.

Dermal fibroblast isolation and culture. Skin biopsies were minced in small pieces and treated with 2.5 U/mL dispase II (Roche Diagnostics) overnight at 4 °C to allow separation of the epidermis and dermis. Fibroblasts were then extracted from the dermis compartment after incubation for 2 hours at 37 °C in 2.5 U/mL dispase II and 750 U/mL collagenase II (Gibco). After centrifugation, cells were resuspended in DMEM with 10% fetal bovine serum (to support cell survival and proliferation) and 2% penicillin-streptomycin. Cells (designated passage 0) were plated in tissue culture-treated plates or flasks and incubated at 5% CO2 in a humidified atmosphere at 37 °C. When reaching an approximate 70% confluence, cells were trypsinized using 0.05% trypsin-EDTA for 5 minutes at 37 °C and seeded at 1,000 cells/cm2 in 96-well plates or in tissue culture flasks up to passage four (P4), which corresponds to approximately 10 cell doublings from the freshly isolated cells.

RNA extraction and micro-array data collection. Isolation of RNA from cultured cells was performed by addition of RLT buffer (RNeasy kit, Qiagen) with 1% β-mercaptoethanol. Isolation of total RNA from the cell lysates was performed using the RNeasy Plus 96 or RNeasy Mini kit (Qiagen) and from frozen biopsies using the RNeasy fibrous kit (Qiagen). Total RNA samples were quantified on the Nanodrop 2000 (Thermo Scientific). RNA quality was assessed using TapeStation 2200 (Agilent Technologies). Samples demonstrating low total RNA concentrations were processed with RNA clean & concentrator kit (Cat. R1080, Zymo Research).

Sample labeling and hybridization were performed using The GeneChip™ HT HG-U133 Plus 96 Array Plate (Affymetrix Inc). Washing and staining of the hybridized arrays were completed as described in the GeneChip Expression analysis technical manual for HT plate arrays using the GeneChip Array Station (Affymetrix Inc). The processed GeneChips plate arrays were scanned using GeneTitan scanner (Affymetrix Inc). RNA integrity was assessed using the HT RNA reagent kit (Part number 760410, Caliper Life Sciences) using a LabChip GX (PerkinElmer). RNA samples with a RQ score of >8.0 were considered high quality for downstream microarray processing. All microarray transcript profiling data are deposited in the Gene Expression Omnibus (GEO) database (GSE# TBD).

Analysis of Affymetrix data. Affymetrix scans were subject to quality control (QC) measures, including visual inspection of the distribution of raw signal intensities and assessment of RNA degradation, relative log expression (RLE), and normalized unscaled standard error (NUSE). CEL files were subjected to GC-content-based Robust Multi-array Average (GCRMA) normalization. Expression levels were log (base 2) transformed. All calculations and analyses were carried out using R and Bioconductor computational tools. For the fibroblast dataset, a strong batch effect was observed as the data came from two different profiling experiments. Nine replicate samples were included in both experiments to control for batch effect and after batch correction. ComBat (R version 3.3.0, Bioconductor sva package version 3.20.0) was used to adjust for batch effects, which uses empirical Bayes frameworks for adjusting data for batch effects, by inputting the GCRMA normalized data. Disease status was included as the covariate of interest.

Analyses were applied to discover genes that were differentially expressed (DEGs). Based on principal component analysis, a few samples were considered to be outliers and removed for downstream analyses (outlier in the biopsies dataset: SSC07_A, outliers in the fibroblast dataset: HS10_P1, SSC04_A_P0 and SSC04 NA_P0). The 9 replicate samples present also in the second fibroblast profiling experiment and that were included to support technical batch effect evaluation and correction were also removed for downstream gene expression analysis. To identify differentially expressed genes between groups of samples, we applied the Linear Models for Microarray (LIMMA) data analysis and paired LIMMA when analyzing difference between affected vs non-affected samples or passage 4 vs passage 0 and passage 1 samples as these were a collection of paired samples originating...
Adenoviral transduction of dermal fibroblasts. Adenoviruses were produced using PerC6 cells as described earlier. Dermal fibroblasts were seeded at 2,000 cells/well in 384-well plates at 8,000 cells/well in 96-well plates in 10% FBS-containing FBS. FBS levels in media were titrated to determine an optimal FBS concentration that supports cell survival while not affecting ASMA expression (not shown). Based on the results from the FBS titration, cells were serum-deprived in 0.5% FBS-containing DMEM two days after seeding and subsequently transduced with adenoviral shRNAs at a multiplicity of infection (MOI) of 20 to 40 infectious units (IU) per cell. One day later (Day 4) the adenoviral vectors were removed and cells were refreshed with 0.5% FBS-containing DMEM. Three days later (day 7) supernatants were harvested to measure ASMA expression levels.

Dermal fibroblast impedance measurements. Healthy donor and SSc patient-derived dermal fibroblasts were seeded at 8,000 cells/well in 10% FBS-containing DMEM on E-plate CardioECR 48-well plates (ACEA Biosciences). The next day cells were serum-deprived in 0.5% FBS-containing DMEM and impedance, presented as the staining intensity (density; D) multiplied by the stained area (A) and normalized for the number of DAPI-positive nuclei (DxA/nuclei).

References
1. Elhai, M., Meune, C., Avouac, J., Kahan, A. & Allanoire, Y. Trends in mortality in patients with systemic sclerosis over 40 years: a systematic review and meta-analysis of cohort studies. *Rheumatology (Oxford)*, 51, 1017–26 (2012).
2. Cossu, M., Beretta, L., Mosterman, P., de Hair, M. J. H. & Radstake, T. R. D. J. Unmet Needs in Systemic Sclerosis Understanding and Treatment: the Knowledge Gaps from a Scientist's, Clinician's, and Patient's Perspective. *Clin. Rev. Allergy Immunol.*, https://doi.org/10.1007/s12016-017-8636-1 (2017).
3. Schulz, J.-N. et al. New developments on skin fibrosis - Essential signals emanating from the extracellular matrix for the control of myofibroblasts. *Matrix Biol.* https://doi.org/10.1016/j.matbio.2018.01.025 (2018).
4. Kissin, E. Y., Merkel, P. A. & Lafyatis, R. Myofibroblasts and hyaline collagen: markers of skin disease in systemic sclerosis. *Arthritis Rheum.* 54, 3655–60 (2006).
5. Perlish, J. S., Lemlich, G. & Fleischmajer, R. Identification of collagen fibrils in scleroderma skin. *J. Invest. Dermatol.* 90, 48–54 (1988).
6. Gardner, H. et al. Gene profiling of scleroderma skin reveals robust signatures of disease that are imperfectly reflected in the transcript profiles of explanted fibroblasts. *Arthritis Rheum.* 54, 1961–73 (2006).
7. Whitfield, M. L. et al. Systemic and cell type-specific gene expression patterns in scleroderma skin. *Proc. Natl. Acad. Sci. USA* 100, 12319–24 (2003).
8. Hinchcliff, M. et al. Molecular signatures in skin associated with clinical improvement during mycophenolate treatment in systemic sclerosis. *J. Invest. Dermatol.* 133, 1979–89 (2013).
9. Assassi, S. et al. Dissecting the heterogeneity of skin gene expression patterns in systemic sclerosis. *Arthritis Rheumatol.* (Hoboken, N.J.) 67, 3016–26 (2015).
10. Pendergrass, S. A. et al. Intrinsic gene expression subsets of diffuse cutaneous systemic sclerosis are stable in serial skin biopsies. *J. Invest. Dermatol.* 132, 1563–73 (2012).
11. Mahoney, J. M. et al. Systems level analysis of systemic sclerosis shows a network of immune and profibrotic pathways connected with genetic polymorphisms. *PLoS Comput. Biol.* 11, e1004005 (2015).
12. Taroni, J. N. et al. A novel multi-network approach reveals tissue-specific cellular modulators of fibrosis in systemic sclerosis. *Genome Med.* 9, 27 (2017).
13. Sargent, J. L. et al. A TGFbeta-responsive gene signature is associated with a subset of diffuse scleroderma with increased disease severity. *J. Invest. Dermatol.* 130, 694–705 (2010).
14. Varga, J. & Whitfield, M. L. Transforming growth factor-beta in systemic sclerosis (scleroderma). *Front. Biosci. (Schol. Ed.)* 1, 226–35 (2009).
15. Tan, F. K. et al. Classification analysis of the transcriptosome of nonlesional cultured dermal fibroblasts from systemic sclerosis patients with early disease. *Arthritis Rheum.* 52, 865–76 (2005).
16. Fuzii, H. T. et al. Affected and non-affected skin fibroblasts from systemic sclerosis patients share a gene expression profile deviated from the one observed in healthy individuals. *Clin. Exp. Rheumatol.* 26, 866–74 (2008).
17. Zhu, L., Gao, D., Yang, J. & Li, M. Characterization of the phenotype of high collagen-producing fibroblast clones in systemic sclerosis, using a new modified limiting-dilution method. *Clin. Exp. Dermatol.* 37, 395–403 (2012).
18. Wallis, D. D. et al. Abnormalities in fibrillin 1-containing microfibrils in dermal fibroblast cultures from patients with systemic sclerosis (scleroderma). *Arthritis Rheum.* 44, 1855–64 (2001).
19. Mori, Y., Chen, S. J. & Varga, J. Expression and regulation of intracellular SMAD signaling in scleroderma skin fibroblasts. *Arthritis Rheum.* 48, 1964–78 (2003).
20. Asano, Y., Ihn, H., Yamane, K., Kubo, M. & Tamaki, K. Increased expression levels of integrin alphavbeta5 on scleroderma fibroblasts. *Am. J. Pathol.* 164, 1275–92 (2004).
21. Chen, Y. et al. Matrix contraction by dermal fibroblasts requires transforming growth factor-beta/activin-linked kinase 5, heparan sulfate-containing proteoglycans, and MEK/ERK: insights into pathological scarring in chronic fibrotic disease. *Am. J. Pathol.* 167, 1699–711 (2005).
22. Shiwen, X. et al. A Role of Myocardin Related Transcription Factor-A (MRTF-A) in Scleroderma Related Fibrosis. *PLOS One* 10, e0126015 (2015).
23. Chen, Y. et al. Contribution of activin receptor-like kinase 5 (transforming growth factor beta receptor type I) signaling to the fibrotic phenotype of scleroderma fibroblasts. *Arthritis Rheum.* 54, 1309–16 (2006).
24. Antonelli, A. et al. Systemic sclerosis fibroblasts show specific alterations of interferon-γ and tumor necrosis factor-α-induced modulation of interleukin 6 and chemokine ligand 2. *J. Rheumatol.* 39, 979–85 (2012).
25. Antonelli, A. et al. IFN-γ and TNF-α induce a different modulation of interleukin-6 in systemic sclerosis fibroblasts compared to healthy controls. *Scand. J. Rheumatol.* 40, 453–6 (2011).
26. Li, X., Zhu, L., Wang, B., Yuan, M. & Zhu, R. Drugs and Targets in Fibrosis. *Front. Pharmacol.* 8, 855 (2017).
27. Garret, S. M., Frost, D. B. & Feghali-Bostwick, C. The mighty fibroblast and its utility in scleroderma research. *J. scleroderma Relat. Disord.* 11, 69–134 (2017).
28. Milano, A. et al. Molecular subsets in the gene expression signatures of scleroderma skin. *PLOS One* 3, e2696 (2008).
29. Frost, J. et al. Differential gene expression of MMP-1, TIMP-1 and HGF in clinically involved and uninvolved skin in South Africans with SSC. *Rheumatology (Oxford).* 51, 1049–52 (2012).
30. Kramann, R. et al. Pharmacological GLI2 inhibition prevents myofibroblast cell-cycle progression and reduces kidney fibrosis. *J. Clin. Invest.* 125, 2935–31 (2015).
31. Liu, L. et al. Insulin-like growth factor binding protein-7 induces activation and transdifferentiation of hepatic stellate cells in vitro. *World J. Gastroenterol.* 15, 3246–53 (2009).
32. Vaidani, S., Bansal, R. & Prakash, J. Drug targeting to myofibroblasts: Implications for fibrosis and cancer. *Adv. Drug Deliv. Rev.* 121, 101–116 (2017).
33. Wettlaufer, S. H., Scott, J. P., McEachin, R. C., Peters-Golden, M. & Huang, S. K. Reversal of the Transcriprome by Prostaglandin E2 during Myofibroblast Dedifferentiation. *Am. J. Respir. Cell Mol. Biol.* 54, 114–27 (2016).
34. Vuerio, T., Mäkelä, J. K. & Vuerio, E. Activation of type I collagen genes in cultured scleroderma fibroblasts. *J. Cell. Biochem.* 28, 105–13 (1985).
35. Hinz, B. Tissue stiffness, latent TGF-beta1 activation, and mechanical signal transduction: implications for the pathogenesis and treatment of fibrosis. *Curr. Rheumatol. Rep.* 11, 120–6 (2009).
36. Arts, G. et al. Adenoviral vectors expressing siRNAs for discovery and validation of gene function. *Genome Res.* 13, 2325–32 (2003).
37. Haak, A. J. et al. Targeting the myofibroblast genetic switch: inhibitors of myocardin-related transcription factor/serum response factor-regulated gene transcription prevent fibrosis in a murine model of skin injury. *J. Pharmacol. Exp. Ther.* 349, 480–6 (2014).
38. Velasquez, L. S. et al. Activation of MRTF-A-dependent gene expression with a small molecule promotes myofibroblast differentiation and wound healing. *Proc. Natl. Acad. Sci. USA* 110, 16850–5 (2013).
39. Zhou, Y. et al. Inhibition of mechanosensitive signaling in myofibroblasts ameliorates experimental pulmonary fibrosis. *J. Clin. Invest.* 123, 1096–108 (2013).
40. Reiter, K., McCormick, R. J. & Rucker, R. B. Enzymatic and nonenzymatic cross-linking of collagen and elastin. *FASEB J.* 6, 2439–49 (1992).
41. Barry-Hamilton, V. et al. Allosteric inhibition of lysyl oxidase-like-2 impedes the development of a pathologic microenvironment. *Nat. Med.* 16, 1009–17 (2010).
42. Ikemura, N. et al. Selective targeting of lysyl oxidase-like 2 (LOXL2) suppresses hepatic fibrosis progression and accelerates its reversal. *Gut* 66, 1697–1708 (2017).
43. Chien, J. W. et al. Serum lysyl oxidase-like 2 levels and idiopathic pulmonary fibrosis disease progression. *Eur. Respir. J.* 43, 1430–8 (2014).
44. Baghu, G. et al. Efficacy of simtuzumab versus placebo in patients with idiopathic pulmonary fibrosis: a randomised, double-blind, controlled, phase 2 trial. *Lancet. Respir. Med.* 5, 22–32 (2017).
45. Chanoki, M. et al. Increased expression of lysyl oxidase in skin with scleroderma. *Br. J. Dermatol.* 133, 710–5 (1995).
46. Rimar, D. et al. Brief report: lysyl oxidase is a potential biomarker of fibrosis in systemic sclerosis. *Arthritis Rheumatol. (Hoboken, N.J.)* 66, 726–30 (2014).
47. Gardet, A., Zheng, T. S. & Viney, J. L. Genetic architecture of human fibrotic diseases: disease risk and disease progression. *Front. Pharmacol.* 4, 159 (2013).
48. Saigusa, R. et al. Multifaceted contribution of the TLR4-activated IRF5 transcription factor in systemic sclerosis. *Proc. Natl. Acad. Sci. USA* 112, 15136–41 (2015).
49. Wehrbrauch, D. et al. An IRF5 Decoy Peptide Reduces Myocardial Inflammation and Fibrosis and Improves Endothelial Cell Function in Tight-Skin Mice. *PLOS One* 11, e0151999 (2016).
50. Schaffner, E., Ray, A. M. & Dontenwill, M. Integrin α5β1, the Fibronectin Receptor, as a Pertinent Therapeutic Target in Solid Tumors. *Cancers (Basel).* 5, 27–47 (2013).
51. Weston, B. S., Wahab, N. A. & Mason, R. M. CTGF mediates TGF-β1-induced fibroblast matrix deposition by upregulating active alpha2beta1 integrin in human mesangial cells. *J. Am. Soc. Nephrol.* 14, 601–10 (2003).
52. Moir, L. M., Burgess, J. K. & Black, J. L. Transforming growth factor beta 1 increases fibroblast deposition through integrin receptor alpha 5 beta 1 on human airway smooth muscle. *J. Allergy Clin. Immunol.* 121, 1034–9.e4 (2008).
53. Chen, G. & Khalil, N. TGF-beta1 increases proliferation of airway smooth muscle cells by phosphorylation of map kinases. *Respir. Res.* 7, 2 (2006).
54. Xu, L. et al. Dioscins, a potent ITG5 inhibitor, reduces the synthesis of collagen against liver fibrosis: Insights from SILAC-based proteomics analysis. *Food Chem. Toxicol.* 107, 318–328 (2017).
55. Gordon, G. M. et al. A cell-based screening assay to identify pharmaceutical compounds that enhance the regenerative quality of corneal repair. *Wound Repair Regen.* 24, 89–99 (2016).
56. Bollong, M. J. et al. Small molecule-mediated inhibition of myofibroblast transdifferentiation for the treatment of fibrosis. *Proc. Natl. Acad. Sci. USA* 114, 4679–4684 (2017).
57. Oh, R. S. et al. RNAi screening identifies a mechanosensitive ROCK-JAK2-STAT3 network central to myofibroblast activation. J. Cell Sci., https://doi.org/10.1242/jcs.209932 (2018).
58. Irizarry, R. A. et al. Summaries of Affymetrix GeneChip probe level data. Nucleic Acids Res. 31, e15 (2003).
59. Li, C. & Hung Wong, W. Model-based analysis of oligonucleotide arrays: model validation, design issues and standard error application. Genome Biol. 2, RESEARCH0032 (2001).
60. Gentleman, R., Carey, V., Huber, W., Irizarry, R. A. & Dudoit, S. Bioinformatics and Computational Biology Solutions using R and Bioconductor. Springer Sci. Media (2005).
61. Johnson, W. E., Li, C. & Rabinovic, A. Adjusting batch effects in microarray expression data using empirical Bayes methods. Biostatistics 8, 118–27 (2007).
62. Ritchie, M. E. et al. limma powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acids Res. 43, e47 (2015).
63. Yeo, B. & Hochberg, Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J R Stat Soc B 57, 289–300 (1995).
64. Chen, E. Y. et al. Enrichr: interactive and collaborative HTML5 gene list enrichment analysis tool. BMC Bioinformatics 14, 128 (2013).
65. Kuleshov, M. V. et al. Enrichr: a comprehensive gene set enrichment analysis web server 2016 update. Nucleic Acids Res. 44, W90–7 (2016).

Acknowledgements
The authors are grateful to the individuals who have participated in this study and volunteered to provide skin biopsies. We thank Dr. Nathalie Franchimont for helpful discussions on inclusion/criteria for the patient recruitment and Dr. Richard Janssen (Galapagos, Leiden) for his input during the initial phases of this project. We thank Dr. Andrea Bertolotti-Ciarlet, Dr. Shelia Violette, Dr. Alan Buckler and Werner Meier for their critical reviews of the approach and data through the project. We thank Dr. Patrick Faloon for suggesting the impedance readout as well as Fleur Stevenhagen and Marijn Vlaming (Ncardia previously Pluriomics, Leiden) who has helped us develop the impedance assay.

Author Contributions
L.C., K.L., E.C.M., D.F.F., D.J., J.L.V., T.S.Z., J.A. and A.G. designed the studies. A.D., J.d.V.B. and T.W.J.H. overviewed patient enrollment. L.C., B.S., S.N.A., M.C. and P.C. performed the experiments. L.C., B.S., S.N.A., M.C., J.A. and A.G. analyzed and reviewed all the cell-based assay data. P.C. collected the microarray data; K.J. and A.G. analyzed the microarray data. L.C., J.A. and A.G. wrote the main manuscript text. All the authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-41153-w.

Competing Interests: K.L., E.C.M., P.C., J.L.V., T.S.Z. and A.G. are current or former employees of Biogen and shareholders. L.C., B.S., S.N.A., M.C., D.F.F., J.D. and J.A. are current or former employees of Charles River, a contract research organization. D.F.F. is a shareholder of Charles River. The authors, including A.D., J.d.V.B. and T.W.J.H. declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019