Research Article

Lentiviral-Mediated RNA Interference against TGF-Beta Receptor Type II in Renal Epithelial and Fibroblast Cell Populations In Vitro Demonstrates Regulated Renal Fibrogenesis That Is More Efficient than a Nonlentiviral Vector

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Received 29 March 2010; Revised 5 July 2010; Accepted 18 August 2010

Academic Editor: G. S. Stein

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Background. Lentiviral constructs reportedly can integrate into the genome of non-dividing, terminally differentiated cells and dividing cells, for long-term gene expression. This investigation tested whether a third generation lentiviral-mediated small interfering RNA (siRNA) delivered into renal epithelial and fibroblast cells against type II transforming growth factor-beta receptor (siRNA-TBRII) could better attenuate renal fibrogenesis in comparison with a non-lentiviral construct. Methods. HIV-derived lentiviral and non-lentiviral constructs were used to transfect cells with siRNA-TBRII or siRNA-EGFP control. Human embryonic kidney (HEK-293T), renal epithelial cells (NRK-52E) and renal fibroblasts (NRK-49F) were transfected and gene silencing quantified (fluorescence microscopy, Western blotting, fluorescence-activated cell sorting). Renal fibrogenesis was assessed using extracellular matrix protein synthesis (fibronectin and collagen-III; Western immunoblot), and α-smooth muscle actin (α-SMA) was analysed as a marker of fibroblast activation and epithelial-to-mesenchymal transdifferentiation (EMT). Results. Lentiviral-mediated siRNA-TBRII significantly suppressed TBRII expression in all cell lines, and also significantly suppressed renal fibrogenesis. In comparison with the non-lentiviral construct, lentiviral-mediated siRNA-TBRII produced stronger and more persistent inhibition of collagen-III in NRK-49F cells, fibronectin in all renal cell lines, and α-SMA in renal epithelial cells. Conclusions. Lentiviral vector systems against TBRII can be delivered into renal cells to efficiently limit renal fibrogenesis by sequence-specific gene silencing.

1. Background

Tubulointerstitial fibrosis is an almost invariable finding in the chronically diseased kidney, irrespective of the underlying disease or the originating compartment. The degree of fibrosis, determined by the relative interstitial volume caused by accumulation of extracellular matrix (ECM), is an important predictor of organ prognosis and kidney excretory function [1]. Renal tubulointerstitial fibrosis is believed to be the final common pathway for nearly all forms of kidney disease that progress towards end-stage renal disease [2].

Transforming growth factor-β (TGF-β), a protein regulator of cell growth and differentiation, is one of the primary mediators that induces accumulation of ECM in renal fibrogenesis [3, 4]. TGF-β increases the production and deposition of ECM proteins, reduces matrix degradation through decreased proteinase production and increased production of matrix degradation inhibitors, and stimulates synthesis of ECM protein receptors [5]. Upregulation of TGF-β is a universal finding in virtually every type of chronic kidney disease, in animal models and in humans [4]. In vitro, TGF-β alone can stimulate mesangial cells and interstitial...
fibroblasts to produce ECM and tubular epithelial cells to undergo epithelial-to-mesenchymal transformation (EMT) to become matrix-producing fibroblast-like cells. Expression of exogenous TGF-β, either via gene delivery in vivo or in transgenic mice, causes renal fibrosis [6]. TGF-β binds to specific receptors on most cells including renal epithelial and fibroblast cells and then initiates a signal cascade that results in production of profibrotic cytokines and inflammatory mediators. The type I and type II signalling receptors mediate the biological actions of TGF-β. The extracellular domain of the type II receptor (TBRII) binds TGF-β, causing the formation of heteromeric complexes [7, 8]. TBRII then transphosphorylates the type I receptor, activates its kinase, and initiates downstream signalling. In this respect, the TBRII appears to be essential for the biological activity of TGF-β signalling pathway [9], and its inhibition or modification is considered as a promising therapeutic strategy to inhibit renal fibrosis.

Modulation of TGF-β action on ECM suppresses tissue fibrosis [2, 10]. For example, biological inhibition of TGF-β protein with neutralising antibodies [11], decorin [12, 13], and soluble TBRII-IgG Fc chimera [14] suppressed the accumulation of ECM in models of renal fibrosis. Other studies have shown that expression of soluble TBRII could effectively block TGF-β signalling in vitro and in vivo, using various means of delivery [15, 16]. However, the methods used in previous studies have therapeutic limitations because the protein or gene is rapidly degraded by enzymes after administration in vivo [17, 18]. The short-term duration of TGF-β signalling inhibition is a major problem to be solved.

Lentiviral vectors can infect non-dividing cells, can be pseudotyped with retroviral envelopes, and so have a broad cell tropism, have no toxic effect, and have stable gene expression due to viral genome integration into cell chromosomes [19]. Despite having many of the same characteristics, retroviral vector-mediated gene therapies or nonviral vector-mediated gene deliveries have a limited or transient effect and can infect only dividing cells [19]. Melding the lentiviral vector-mediated gene delivery and the powerful tool of RNA inhibition (RNAi) could potentially provide targeted long-term gene silencing. RNAi is a mechanism for sequence-specific, posttranscriptional gene silencing triggered by double-strand RNA (dsRNA; referred to as small interfering RNA, silencing RNA or siRNA), which targets the degradation of complementary miRNAs [20]. The siRNA mediators of RNAi typically consist of 19–23 nucleotide RNA duplexes. During RNAi, the introduction of siRNA by transfection into a diverse range of organisms and cell types causes degradation of the complementary mRNA, thereby silencing gene expression.

Because of the ability of lentiviral vectors to deliver transgenes into tissues that are considered resistant to stable genetic manipulation, like well-differentiated adult renal cell populations with limited levels of mitosis, new possibilities for gene therapy are opening. Gusella et al. [21] successfully transduced a 1st-generation lentiviral vector into mouse kidneys. The current study used a cell culture model of renal fibrosis to compare the efficiency to limit renal fibrogenesis and the durability of expression of a lentiviral versus a nonlentiviral versus a lentiviral-delivered siRNA to TBRII (siRNA-TBRII).

2. Methods

2.1. Cell Culture. Rat renal epithelial (NRK-52E, ATCC CRL1571) and fibroblast (NRK-49F, ATCC CRL1570) cell lines were selected for comparison of fibrogenesis. Human embryonic kidney- (HEK-) 293T cells, a derivative of the HEK-293 cell line (ATCC CRL1573) were selected for plasmid transfection and lentiviral vector transduction. An enhanced green fluorescent protein (EGFP) stably expressing HEK-293T cell line (S7 cells, established at Dr. Ming Wei’s laboratory, Department of Medicine, Prince Charles Hospital, University of Queensland, Brisbane, Australia) was used for transfection to evaluate the silencing effect on EGFP by RNAi.

NRK-49F and NRK-52E cells were maintained routinely in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% bovine calf serum (BCS), 1,000 units per millilitre (U/mL) penicillin, and 1,000 μg/mL streptomycin (BioWhittaker, Edward Keller Pty Ltd, Silverwater, Australia). For HEK-293T cells and the EGFP stably expressing S7 cells, DMEM was supplemented with 10% fetal calf serum (FCS). Cells were grown in a humidified atmosphere of 95% air and 5% carbon dioxide at 37°C in a tissue culture incubator. The original cell seeding density was of 2–5 × 10^5 cells per mL. Cells were grown for 24–48 hours prior to treatment or until just subconfluent. Prior to treatment, all cells were washed twice in serum-free (SF) DMEM containing antibiotics only, to remove any traces of serum growth factors or supplements.

2.2. PCR Cloning and Plasmid Preparations. A 444 bp extra-cellular ectodomain including the critical soluble TBRII TGF-β binding domain comprising the amino acid DNA sequence from 284 bp to 727 bp (targeting the sequence between starting codon and the beginning of hydrophobic trans-membrane spanning domain) of Rattus norvegicus TBRII mRNA (Genbank s67770) [22] was generated by reverse transcriptase PCR from normal rat kidney cDNA using a QIAGEN Taq PCR core kit. The 444 bp DNA sequence of the TBRII gene was ligated into the pDrive vector (QIAGEN, Australia) at a molar ratio of 5–10 times more PCR product DNA than pDrive cloning vector DNA. The orientation of PCR product in the recombinant plasmid was confirmed by restriction enzyme digestion. After enzyme digestion, the recombinant plasmid pDrive-TBRII with the PCR insert of the 444bp soluble domain of the TBRII gene was mini-preped using a GenElute Plasmid Miniprep Kit (Sigma-Aldrich, Australia) and maxi-preped for a larger amount of plasmid production using a Plasmid Maxi Kit (QIAGEN, Australia). The constructed pDrive-TBRII with its PCR insert was further confirmed by DNA sequencing at the Australian Genome Research Facility (Gehrmann Laboratories, University of Queensland, http://www.agrf.org.au/).
2.3. Preparation of RNAi Expression Plasmids. Two complementary oligonucleotides necessary to create the hairpin insert for pPlasRi cloning vectors were designed following a software siRNA Wizard v2.5 available on line at http://www.sirnawizard.com/. The sequences of custom-designed oligonucleotides were as follows: TBRII RNAi short hairpin (sh) insert “A” targeting the first 100bp starting from ATG start codon: 5’-GAT CCC C ACGTTCAGAAGGCTGTTAA T TCA AGA GA TTAACCGACTGGAAGCT TTT TTG GAA A-3’ and 5’-TCGA TT TCC AAA AA ACGTTCAGAAGGCTGTTAA TCT CTT GAA TTAACCGACTGGAAGCT GTT G-3’, insert “B” targeting the last 100bp of codons of TBRII: 5’-GAT CCC C AGATTCCAGAAGACGGCTC T TCA AGA GA GAGCCGCTCTCCTCGGAATCT TTT TTG GAA A-3’ and 5’-TCGA TT TCC AAA AA AGATTCCAGAAGACGGCTC TCT CTT GAA GAGCCGCTCTCCTCGGAATCT GTT G-3’. The 19-nucleotide TBRII target sequences are underlined in these sequences. Two single-strand oligonucleotides synthesised for EGFP and used as experimental control siRNA were provided by Dr. Ming Q. Wei (one of the authors). All oligos were purchased in a lyophilised desalted form from Sigma Genosys Australia (Castle Hill, Australia). A pair of oligonucleotide inserts was annealed using the methods previously described [23]. After annealing of two-single stranded oligos into double-stranded inserts, siTBRII and siEGFP were ligated to linearised pPlasRi plasmid (provided by Dr. Ming Q. Wei, one of the authors) that contains the human H1 RNA Polymerase III promoter after BglII and SalI digestion. Compatible enzyme sequences were synthesized for EGFP and used as experimental control siRNA.

2.4. Production of Lentiviral RNAi Constructs. The third-generation replication-defective HIV-1-based lentiviral vector pHIV-CS (7488 bp) was provided by Dr. Ming Q. Wei, one of the authors. Both lentiviral RNAi constructs (pLenti-TBRII and pLenti-EGFP) were constructed by cloning the pHIV-CS backbone with siRNA inserts derived from pPlasRi-RBRII or pPlasRi-EGFP by BamHI and XhoI digestion, in which the RNA promoter was also transferred to the lentiviral RNAi plasmids.

2.5. Lentiviral Construct Transfection In Vitro. For in vitro transfection, 1 × 10⁵ cells were seeded per well in 12-well plates 16–18 hours before the experiment, by which time the cells reached 75–80% confluence. Plasmids DNA of 2 μg with a linear PEI derivative (ExGen 500 MBI Fermentas, Hanover MD, USA), at N/P ratio of 1:6 to 1:9, were incubated for 10 minutes at room temperature and then added to the wells. The transfection medium containing plasmid DNA was replaced 8 hours later with fresh growth medium and cells were further incubated for 16 hours or longer for up to 72 hours as required.

2.6. FACS Quantification of EGFP Expression. EGFP gene expression was analysed by direct fluorescence using a confocal microscope (Queensland Institute for Medical Research, Brisbane), and images were digitised. Fluorescence-activated cell sorting (FACS; Becton Dickinson and Co, Franklin Lakes NJ, USA) was performed using 10,000 fixed cells at various intervals after transfection to quantify the number and the mean fluorescence intensity of cells positive for EGFP. Untransfected HEK-293T and S7 cells were used to determine the cutoff values. Gene silencing effectiveness was evaluated by direct visualization under a fluorescent microscope (in the stably EGFP-expressing S7 cells which were transfected by EGFP RNAi plasmids) and quantified by FACS. Immunofluorescence (IF) microscopy or Western immunoblot was used for protein analysis of TBRII in NRK-49F and NRK-52E cells which were transfected by TBRII RNAi plasmids.

2.7. Immunofluorescence. For IF staining of TBRII on transfected HEK-293T cells, cells on coverslips were washed and fixed, and then nonspecific binding was blocked with 3% bovine serum albumin in Tris-buffered saline with Tween-20 (BSA in TBST) for 1 hr at room temperature. Coverslips were incubated with primary antibody against TBRII (TGFβ RI, 1:10, Santa Cruz) for 1 hour at room temperature (RT). Sections were washed 3 × 30 minutes in TBST, then incubated with a fluorescent secondary antibody mixture containing an FITC-labelled goat antirabbit IgG (Molecular Probes, Oregon, USA) at a dilution of 1:200 in 1% BSA in TBST for 1 hour at RT in a dark cupboard. Coverslips were washed 3 × 30 minutes in TBST, then mounted onto glass slides using DAKO antifade fluorescence mounting medium (DAKO, Australia), and allowed to set overnight in the dark at 4°C or over 2–3 hours at RT prior to confocal fluorescence microscopy.

2.8. Western Immunoblot. Cells were disrupted in ice-cold cell lysis buffer (50 mM Tris-Cl at pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 0.1% sodium dodecyl sulphate (SDS), 25 mM sodium fluoride, and 0.5 M ethylenediamine tetra-acetic acid) containing protease and phosphatase inhibitors (100 μg/mL phenylmethylsulfonyl fluoride, 20 μg/mL leupeptin, 20 μg/mL aprotinin, and 1mM sodium orthovanadate, all Sigma-Aldrich products) using a tissue homogenizer. Cell debris was removed by centrifugation at 16,000 g for 15 minutes at 4°C. Protein concentration was determined in each extract by a Bradford protein assay (Bio-Rad Pty Ltd, Sydney, NSW, Australia) and spectroscopy at 595 nm. 30 μg of total protein were loaded onto 12% SDS polyacrylamide gel, run at 100 V with a set current of 60 mamps for 1.75 hours, and then transferred to a polyacrylic membrane (Perkin Elmer, Melbourne, Australia) using a Bio-Rad Mini Protein 3 unit (Bio-Rad, Australia). Nonspecific binding of antibodies to the membrane was blocked by incubation in 4% skim milk powder in TBST buffer (blotto) for 1 hour with continuous rocking. Antibodies were added to blotto or 0.1% BSA in TBST (primary
antibody TGFβ RII Mouse mAb) and incubated with membranes overnight at 4°C on a rocking platform. Blots were washed repeatedly (4–5 times, 5–10 minutes each) in TBST and then incubated with the corresponding Horseradish peroxidase- (HRP-) conjugated secondary antibody (rabbit-α-mouse IgG-HRP) diluted in either 4% blotto or 0.1% BSA in TBST for 1 hour at RT on a rocking platform. Membranes were subjected to a second stage of vigorous and stringent washes (4-5) in TBST containing 0.1% Igepal CA-630, each for 5-minute duration. Membranes were incubated with SuperSignal West Pico chemiluminescent substrate (Pierce Biotechnology, Rockford, IL, USA) for 2 minutes with vigorous shaking. Protein bands were detected using enhanced chemiluminescence (ECL) imaging onto X-ray film (Kodak X-Omat AR-5 or Fuji Rx XR Film) and scanned using a Hewlett-Packard ScanJet 3200C at 300 dpi. Scion Image software (v4.0.2) was used to quantify the density of protein bands minus local background in arbitrary densitometry units. All protein bands were normalised against actin, or membranes were stained with Coomassie Brilliant Blue (Sigma-Aldrich: catalogue number B7920) to verify equal protein loading of lanes. Cell extract positive controls from antibody suppliers were routinely used for immunoblots.

2.9. Statistical Analysis. Data were analysed using standard statistical methods, linear regression, Students, t-test, one-way analysis of variance (ANOVA), and Tukey’s, Dunnett’s multiple comparison, or Bonferroni posttest using Graphpad software. Most data are presented in graphic form as the means ± SEM. Significance was assessed at P < .05.

3. Results

3.1. RNAi Expression Constructs for TBRII and EGFP. TBRII and EGFP RNAi expression constructs were constructed first, followed by the lentiviral RNAi construct. EGFP was used as a reporter gene. The RNAi effect was first tested by transfection of the expression plasmids in vitro before moving onto Lenti-RNAi plasmid construction. Once the gene silencing effect was confirmed, RNAi cassettes together with the upstream RNA promoter were transferred into a third-generation HIV-1 based lentiviral plasmid. Two single-stranded oligonucleotides were synthesized for TBRII and EGFP siRNA were annealed to be double-stranded inserts. Constructed RNAi expression plasmids pPlasRi-TBRII and pPlasRi-EGFP were confirmed by restriction digestion.

3.2. Inhibition of EGFP by RNAi Expression Construct and Lentiviral RNAi Construct In Vitro. EGFP RNAi expression construct pPlasRi-EGFP and lentiviral RNAi construct pLentiRi-EGFP were transfected into S7 cells, facilitated by the cationic polymer PEI. Silencing of EGFP expression by RNAi was evaluated by counting numbers of green fluorescent cells by FACS analysis at 24 hours, 48 hours, and 72 hours post-transfection. Empty pPlasRi and pLentiRi constructs without RNAi insert were used as negative controls. These results are demonstrated in Figure 1. There was 78.28%, 69.37%, and 40.09% inhibition of EGFP expression (P < .001) as measured by FACS at 24, 48, and 72 hours, respectively, post transfection of pPlasRi-EGFP into S7 cells. The negative control group did not show EGFP inhibition. The RNAi expression construct pPlasRi-EGFP gene silencing effect was however transient, peaking at 24 hours, and then starting to diminish. In comparison, persistent EGFP inhibition up to 72 hours was observed post lentiviral RNAi construct pLentiRi-EGFP transfection, with an 84.17%, 91.57%, and 95.91% decrease of EGFP expression achieved at 24, 48, and 72 hours, respectively, and EGFP silencing effect by lentiviral construct increased with time. There was no significant difference in gene silencing effect when comparing the transfection of the two constructs at 24 hours. The lentiviral RNAi construct, however, demonstrated significant advantages in posttranscriptional gene silencing than the expression construct at 48 and 72 hours. No EGFP inhibition was observed in the negative control group.

3.3. Posttranscriptional Sequence-Specific Gene Silencing of TBRII by RNAi In Vitro. Having proven successful EGFP gene silencing by RNAi-expressing and lentiviral-RNAi constructs in S7 cells, RNAi gene silencing of TBRII was evaluated in renal epithelial and fibroblast cell lines (NRK-52E and NRK-49F) in vitro. An embryonic renal cell line HEK-293T was also tested. Inhibition of ECM accumulation in renal cells as a result of downregulation of TGF-β signalling by RNAi was investigated.

3.4. TBRII Gene Suppression in HEK-293T by RNAi. TBRII RNAi expression construct pPlasRi-TBRII and lentiviral RNAi construct pLenti-Ri-TBRII were transfected into cultured HEK-293T cells, facilitated by transfection agent PEI. Inhibition of TBRII was examined by FITC-fluorescence immunohistochemistry and Western blot at 24, 48, and 72 hours. Empty pPlasRi and pLentiRi constructs without RNAi insert were used as negative controls. Inhibition of TBRII expression was observed after both pPlasRi-TBRII and pLentiRi-TBRII transfection. TBRII inhibition in both experimental groups persisted for up to 72 hours post-transfection. pLentiRi-TBRII construct transfection showed a significantly stronger inhibition of TBRII when compared to pPlasRi-TBRII at 72 hours (P < .01). Gene silencing effect of lentiviral RNAi construct increased with time (P < .01). These results are shown in Figure 2.

3.5. TBRII Gene Suppression by RNAi in Renal Epithelial and Fibroblast Cells. Having confirmed the RNAi effect in HEK-293T cells post-transfection of both RNAi expression construct and lentiviral RNAi construct, gene silencing of TBRII was tested in renal epithelial (NRK-52E) and fibroblast (NRK-49F) cells. Expression of TBRII protein was evaluated by Western blot analysis and semiquantified by densitometry at 24, 48, and 72 hours post-transfection of pPlasRi-TBRII and pLentiRi-TBRII. There was a significant decrease of TBRII expression in both cell lines at all tested
Figure 1: Continued.

Control

pPlasRi-EGFP 24 h

pPlasRi-EGFP 48 h

pPlasRi-EGFP 72 h
**Figure 1:** Gene silencing of EGFP by transfection of lentiviral and non-lentiviral expression constructs in vitro. RNAi EGFP gene silencing was tested at 24, 48, and 72 hrs in EGFP stably expressing S7 cells post transfection of an EGFP-RNAi expression construct (pPlasRi-EGFP) and a lentiviral RNAi construct (pLentiRi-EGFP). Fluorescence microscope images of cultured cells are demonstrated as well as the FACS analysis. The percent of EGFP positive (+) cells (mean ± SEM) was assessed using FACS results, and these are shown graphically. There was a significant decrease of EGFP 24 hrs after both transfections compared to controls (\( ** P < .01 \) in comparison with controls). The pPlasRi-EGFP plasmid showed a transient silencing effect in comparison with a persistent inhibition by the pLentiRi-EGFP, which produced a continuing inhibition of EGFP (+) cells at 48 and 72 hrs. Gene silencing of EGFP by the pLentiRi-EGFP plasmid remained at 95.91% at 72 hrs.
time points. The lentiviral construct achieved a higher level of inhibition and a longer time of gene silencing of TBRII in both renal epithelial and fibroblast cell lines. These results are demonstrated in Figure 3.

3.6. Inhibition of Fibrogenesis by TBRII RNAi in Renal Cells.

The biological effect of TBRII RNAi should be reduced fibrogenesis, represented by reduced ECM production and reduced EMT. Inhibition of fibrogenesis was evaluated by Western blotting in HEK-293T, NRK-52E, and NRK-49F cells. Collagen III, fibronectin (ECM production), and α-SMA expression (EMT) were tested and semiquantified by densitometry at 24, 48, and 72 hours post-transfection of the RNAi expression construct pPlasRi-TBRII and lentiviral RNAi construct pLentiRi-TBRII. These results are demonstrated in Figures 4 and 5. In Figure 4, there was a significant decrease of collagen III expression at 24 and 72 hours post-transfection with both constructs in HEK-293T cells, and there was significant inhibition of collagen
III expression in NRK-52E cells at 24 hours (P < .01). Collagen III inhibition was not apparent at 72 hours in NRK-52E cells. A significant decrease of collagen III expression was also observed in NRK-49F cell lines by lentiviral RNAi construct at 24 and 72 hours (P < .05). Fibronectin expression was significantly decreased in HEK-293T, NRK-52E, and NRK-49F cells at 24 and 72 hours transfection of either TBRII RNAi expression or lentiviral construct. The variation in responses of the different cell lines, including the time line responses, underlines the complexity of cellular contributions to fibrosis and the antifibrotic responses when planning therapies involving the gene therapies.

There was significantly reduced α-SMA expression in NRK-52E and NRK-49F cells post-transfection of both RNAi constructs (Figure 5). Both expression vectors significantly reduced α-SMA expression in NRK-49F cells at 72 hours. In the NRK-52E cells, α-SMA expression was markedly reduced at both time points, although the reduction in expression was most marked with the pLentiRi vector.

4. Discussion

Renal fibrosis is the common pathway of chronic renal disease that progresses to end-stage kidney disease. Chronic renal disease is characterized by persistent accumulation and deposition of ECM which leads to widespread tissue fibrosis [3, 4, 24]. While progress has been made in delineating the cellular and molecular pathogenesis during the last decades, the clinical therapies for chronic renal fibrosis remain extremely limited. Drug delivery systems that take advantage of recent RNAi technologies may open new pathways for therapeutic strategies. In the present study, a lentiviral construct-mediated RNAi system for posttranscriptional, sequence-specific gene silencing of TBRII for the treatment of renal fibrosis was demonstrated. Knockdown of the TBRII gene was observed in cultured renal epithelial and fibroblast cell lines, and consequently EMT and TGF-β signalling-regulated ECM accumulation was halted. The present study demonstrates that the introduction of TBRII siRNA with lentiviral construct to renal cell populations can effectively suppress the expression of TBRII. If applied in disease, this system could limit renal tubulointerstitial fibrosis.

As a new technology for silencing of a target gene, a pioneer study in the field of renal fibrosis has shown that siRNA targeting TBRII effectively inhibited renal interstitial fibrosis by using an RNAi expression construct with cationic gelatin, in an attempt to overcome the transient, insufficient transfection efficiency of chemically synthesised siRNAs [17]. TBRII expression and α-SMA expression were suppressed for up to 10 days in a unilateral ureteral obstruction (UUO) mouse model. A recent study using TGF-β1 siRNA showed successful delay of the tubulointerstitial fibrosis process by evaluating collagen accumulation in the UUO model of renal fibrosis, in mice [25]. In this study, an RNAi expression construct processing the U6 RNA promoter, pU6shX, was employed to demonstrate that RNAi targeting of the TGF-β signalling pathway could be a valuable tool for prevention and treatment of renal tubulointerstitial fibrosis.
Figure 4: Inhibition of ECM by TBRII RNAi in renal cell lines. Western immunoblots of protein extracts from human embryonic kidney cells (HEK-293T), rat renal fibroblasts (NRK-49F), and epithelial cells (NRK-52E) showed significantly reduced ECM production after TBRII RNAi plasmid transfection (fibronectin is abbreviated as “F’nectin” beside the Western immunoblots). There was significantly decreased collagen III and fibronectin expression post-transfection with pPlasRi-TBRII and pLentiRi-TBRII in NRK-49F and NRK-52E cell lines. In comparison with pPlasRi-TBRII, pLentiRi-TBRII produced a stronger and longer inhibitory effect on collagen III in NRK-49F and on fibronectin in all 3 tested cell lines (∗∗∗∗ P < .01 and ∗∗ P < .05 in comparison with controls).
In the present study, a similar RNAi construct, namely, pPlasRi, was used with the polymerase III H1-RNA gene promoter [23, 25, 26]. It produces a small RNA transcript lacking a polyadenosine tail and has a well-defined start of transcription and a termination signal consisting of five thymidines in a row [26]. Two pairs of 19-nt sequence-targeted short hairpin RNAi (shRNAi) cassettes were designed and cloned into the RNAi expression construct. EGFP was selected as the experimental control because fluorescence is easy to observe under the fluorescent microscope and quantified by FACS analysis [27, 28]. The result showed satisfactory EGFP suppression of 78.28% at 24 hours. The inhibition effect was however weakened to 69.37% and 40.09% at 48 hours and 72 hours, respectively, post transfection. This indicated that a gene silencing effect of EGFP could be achieved by an RNAi expression construct, but the effect is transient. Similarly, after construction of the TBRII RNAi expression construct and its transfection to HEK-293T cell lines, reduced TBRII expression was most pronounced at 24 hours post transfection. In comparison with 48 hours and 72 hours, the transient RNAi effect of the expression construct was again noted.

In principle, there are two general siRNA delivery methods. One uses chemically synthesised 19-21nt siRNA with apparent short-lived effect [28]. The other approach is vector-based siRNA, including the use of plasmid vectors and viral vectors. Vector-based siRNA has a more prolonged effect and is relatively inexpensive. Plasmid-mediated biological activity of RNAi still needs to be improved by various means. Several synthetic materials, including cationic liposomes, poly-L-lysine, and PEI, have been molecularly designed to allow successful transfection of plasmid DNA for mammalian cells both in vitro and in vivo. In Kushibiki et al.’s pSUPER TBRII RNAi system, cationic gelatin was used to facilitate transfection, and it showed relatively satisfactory gene silencing effect in the UUO model for up to 10 days [5]. Our previous experience indicated that a cationic polymer PEI could facilitate better transfection than poly-L-lysine [29], and this was confirmed in the renal cell lines tested. The results showed that there was RNAi effect on EGFP and TBRII in both renal epithelial and fibroblast cells after RNAi expression plasmid transfection.

A prerequisite for developing RNAi-based therapeutics is an efficient delivery vehicle to transfer the RNAi cassette into the target cells. Our results demonstrated the transient nature of the RNAi vectors and the ability of the lentiviral vector-mediated RNAi system for a sustained transduction effect, as had been demonstrated previously in terminally differentiated cells [30]. The advantages of lentivirus for gene transfer, especially to non-dividing cells, make the RNAi technique more accessible for specific in vitro and in vivo silencing in renal cell populations. The lentiviral construct showed significant silencing of EGFP in S7 cells. There was 84.17%, 91.57%, and 95.91% decrease of EGFP achieved at 24 hours, 48 hours, and 72 hours post PEI facilitated transfection. In comparison with the non-lentiviral RNAi expression construct, the gene-silencing effect of the lentiviral RNAi construct was stronger and persistent (P < .01). TBRII gene silencing in HEK-293T cells showed a similar result that favoured the lentiviral RNAi construct.
TBRII expression, ECM production (collagen III, fibronectin), and marker of EMT (α-SMA expression) were examined after gene suppression of TBRII. Both constructs achieved significant inhibition with reduced collagen III and fibronectin expression in HEK-293T, NRK-49F, and NRK-52E cells, and with the lentiviral RNAi construct producing a more robust inhibition of fibrogenesis and continuing inhibition of TBRII 72 hours post-transfection. Complete TBRII silencing was not achieved, even though a lentiviral construct was employed, and was facilitated by a cationic transfection aid PEI. During renal fibrogenesis in vivo, several events occur in parallel, including an influx of monocytes/macrophages, activation of myofibroblasts, accumulation of various ECM proteins, and EMT [31, 32]. As an indicator of fibroblast activation and tubular EMT, α-SMA expression was evaluated in renal fibroblasts and epithelial cells. Significant reduction of α-SMA was shown by Western blot analysis, indicating an inhibitory effect on fibroblast activation and EMT.

5. Conclusion

The present data demonstrated that the lentiviral-mediated RNAi in renal cell populations was effective not only in silencing TBRII gene, therefore, blocking TGF-β signalling, but also modulating activation of fibroblasts, development of EMT, and reducing ECM accumulation. Successful prolonged presence of RNAi in renal cells in vitro was also demonstrated. The lentiviral RNAi transfection facilitated by cationic polymer PEI was superior to the classical RNAi by cationic polymer PEI was superior to the classical RNAi in renal cell populations was effective not only in silencing TBRII gene, therefore, blocking TGF-β signalling, but also modulating activation of fibroblasts, development of EMT, and reducing ECM accumulation. Successful prolonged presence of RNAi in renal cells in vitro was also demonstrated. The lentiviral RNAi transfection facilitated by cationic polymer PEI was superior to the classical RNA-promoter-possessed plasmid-mediated RNAi. Thus, this technique of delivering RNAi into tubulointerstitial cells could be a practical therapeutic method for progressive renal interstitial fibrosis.

List of Abbreviations

ANOVA: Analysis of variance
BSA: Bovine serum albumin
DMEM: Dulbecco's modified Eagle's medium
dsRNA: Double-stranded RNA
ECM: Extracellular matrix
EGFP: Enhanced green fluorescent protein
EMT: Epithelial-to-mesenchymal transition
F’nectin: Fibronectin
γ-GT: Gamma-glutamyl transpeptidase
HEK: Human embryonic kidney
NRK: Normal rat kidney
PCR: Polymerase chain reaction
PEI: Polyethyleneimine
RNAi: RNA interference
SF: Serum-free
siRNA: Silencing RNA
TBRII: Transforming growth factor-beta receptor II
TBST: Tris-buffered saline-Tween
TGF-β: Transforming growth factor-beta.

Acknowledgments

For this work, T. Yang was supported by an International Postgraduate Research Scholarship from The University of Queensland and an Australian Postgraduate Award from the Australian Federal Government. The paper was also supported by a grant from Kidney Health Australia. All authors contributed to the development of the testable hypotheses, analysis of data, development of results concepts, and writing the paper. Most of the cell culture lentiviral work was carried out by Dr. T. Yang, with some help from Drs. B. K. Pat and B. Zhang.

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