A TALEN-based strategy for efficient bi-allelic miRNA ablation in human cells

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ABSTRACT

Significant progress in the functional understanding of microRNAs (miRNAs) has been made in mice, but a need remains to develop efficient tools for bi-allelic knockouts of miRNA in the human genome. Transcription activator-like effector nucleases (TALENs) provide an exciting platform for targeted gene ablation in cultured human cells, but bi-allelic modifications induced by TALENs alone occur at low frequency, making screening for double knockouts a tedious task. Here, we present an approach that is highly efficient in bi-allelic miRNA ablation in the human genome by combining TALENs targeting to the miRNA seed region with a homologous recombination donor vector and a positive selection strategy. A pilot test of this approach demonstrates bi-allelic miR-21 gene disruption at high frequency (∼87%) in cultured HEK293 cells. Analysis of three independent clones showed a total loss of miR-21 expression. Phenotypical analysis revealed increased miR-21 target gene expression, reduced cell proliferation, and alterations of global miRNA expression profiles. Taken together, our study reveals a feasible and efficient approach for bi-allelic miRNA ablation in cultured human cells and demonstrates its usefulness in elucidating miRNA function in human cells.

Keywords: double knockout; gene-editing; homology-directed repair (HDR); microRNA; transcription activator-like effector nucleases (TALENs)

INTRODUCTION

MicroRNAs (miRNAs) are short noncoding RNAs that negatively regulate gene expression in mammalian cells by targeting the 3′ untranslated region of mRNAs. In recent years, it has become increasingly apparent that miRNAs are of crucial functional importance for normal development and physiology, as well as a factor in various diseases in mammals (Svoboda and Flemr 2010; Pauli et al. 2011; Amiel et al. 2012; Gommans and Berezikov 2012; Iorio and Croce 2012b). For instance, miR-21 was found to be overexpressed in virtually all types of human cancers and thus has emerged as an important therapeutic target in cancer treatment (Huang et al. 2012; Iorio and Croce 2012a; Li et al. 2012a,b). Recently, miR-21 has been shown to play crucial regulatory roles in cell growth, proliferation, and apoptosis, as well as in autoimmune and cardiovascular diseases (Buscaglia and Li 2011; Liu et al. 2013; Xu et al. 2013).

Despite their fundamental importance, the exact functions of miRNAs in the context of human development and disease processes remain largely unknown. This is, in part, due to a lack of effective methods for completely abolishing the expression of miRNAs in human cells and disease-relevant models (Park et al. 2010, 2012). Although targeted gene knockdown by short-interfering RNAs (siRNAs) provides a rapid and inexpensive tool to functionally study most protein-coding genes, it cannot be used to reduce mature miRNAs in a sensible way at the cellular level. Alternatives to siRNAs include small molecule inhibitors, antisense oligonucleotides, anti-miR vectors (miRZips), and miRNA sponges (Krutzenfeldt et al. 2005; Zhu et al. 2011; Hu et al. 2013b). The major limitations of these methods are (1) the transient nature of their effects, and (2) a high risk of off-target effects and resulting toxicity (Jackson et al. 2003; van Dongen et al. 2008; Khan et al. 2009), limitations highlighted by reports on discrepancies between the effect of miRNA inhibitors and genetic knockouts (Patrick et al. 2010; Park et al. 2012).

Transcription activator-like effector nucleases (TALENs) are powerful gene editing tools for uncovering gene functions...
efficient bi-allelic miRNA ablation in human cells

(Bogdanove and Voytas 2011; Clark et al. 2011; Miller et al. 2011; Joung and Sander 2013; Sun and Zhao 2013). Though relatively new, TALENs have been successfully employed in a broad variety of systems (Christian et al. 2010; Hockemeyer et al. 2011; Carlson et al. 2012; Tong et al. 2012; Sung et al. 2013; Zhang et al. 2013). TALENs are custom endonucleases that work as dimers to create double-strand breaks in their target DNA sequences (Clark et al. 2011; Miller et al. 2011; Joung and Sander 2013; Sun and Zhao 2013). Each TALEN typically recognizes a binding site of ∼20 bp, providing a high specificity of gene targeting. Coupled with the ability of context-independent binding (Cermak et al. 2011; Moore et al. 2012; Ansai et al. 2013; Kim et al. 2013a), TALENs are uniquely suited to precisely edit very small genes such as miRNAs (Hu et al. 2013a; Kim et al. 2013b).

A number of approaches have been developed for rapid assembly of custom TALENs (Cermak et al. 2011; Briggs et al. 2012; Sanjana et al. 2012; Sakuma et al. 2013; Uhde-Stone et al. 2013). With these advances, TALEN pairs can be generated easily and economically in a matter of days. Many protein-coding genes or genetic loci with long genome sequences have been documented for the purpose of gene editing, but the short and noncoding nature of miRNA presents different challenges. Hu et al. have reported disruption of miRNA genes in human cells by TALENs. While it is possible to disrupt genes in the human genome with TALENs alone at a frequency of typically 2%–40%, averaging around 16% for mono-allelic disruptions (Kim et al. 2013a; Sakuma et al. 2013), isolating cells carrying rare bi-allelic disruptions requires time-consuming single-cell-derivation and subsequent screening. Our approach differs from other strategies (Hu et al. 2013a; Kim et al. 2013b) in that we combine TALENs targeting to the miRNA seed region with a homologous recombination (HR) donor vector carrying a selectable marker. Our strategy enables convenient positive selection, and the combination of NHEJ with stem–loop deletions results in efficient bi-allelic miRNA gene ablation, which is especially valuable for loci that may be difficult to target. Additionally, by using HR donors, endogenous loci can be potentially modified with custom sequences (such as IRES-fluorescent proteins) to allow functional assessment of endogenous gene expression and regulation (Hockemeyer et al. 2011).

As proof of concept, we targeted the human miR-21 seed region in cultured human HEK293 cells, and successfully selected bi-allelic miRNA knockouts with high efficiency (87%). Quantitative RT-PCR analysis of three independent clones confirmed complete loss of mature miR-21 expression. Phenotypical analysis confirmed increased protein levels of the miR-21 target gene PDCD4, reduced cell proliferation, and changes in global miRNA expression profiles. Re-expression of miR-21 in a miR21-knockout line restored miR-21 function, demonstrated by a decrease in target gene expression, further supporting the validity of our approach. Taken together, the high efficiency and ease of a positive selection protocol for bi-allelic miRNA deletions in the human genome provides a powerful tool for elucidating miRNA function in humans to realize, in the long-term, their full therapeutic potential.

results

system design and knockout strategies

TALENs have almost no restriction in regard to their target sequences. To take advantage of this feature, we designed a TALEN pair flanking the ∼6-bp seed region of the human miR-21, directing FokI cleavage to the seed region (Fig. 1A). TALEN-induced double-strand breaks can achieve gene knockouts in two main ways: by homologous recombination of exogenous donor DNA, or by nonhomologous end joining (NHEJ) repair. In the case of HR events, the donor replaces the entire miR-21 precursor with an RFP and a puromycin-resistance expression cassette (Fig. 1B). To completely knock out a gene of interest in diploid cells, the other allele must be disrupted by a simultaneous HR event or, more likely, by NHEJ-induced mutations. Due to the two selectable marker genes embedded in the HR donor vector (RFP and puromycine resistance), we can easily isolate the clonal populations of cells in which an HR event occurred. Because NHEJ is the predominant repair mechanism induced by double-strand breaks, selecting for HR events will most likely identify clones that harbor bi-allelic modifications, with the second allele carrying a NHEJ-mediated mutation.

junction PCR and sequencing confirms seamless donor integration by HR at miR-21 locus

To evaluate correct TALEN-mediated HR donor integration in the human miR21 genomic locus (chromosome 17q23.2; 55273409–55273480, adjacent to the coding gene TMEM49), we conducted transient transfection studies using TALENs in combination with an HR donor vector in HEK293 cells. Junction PCR with a combination of genome and vector-specific primers (Fig. 1C) was performed as specified below. The successful amplification of the correct PCR products was obtained only in cells cotransfected with both TALENs and HR donor vectors (Fig. 2A) but not in cells transfected with the HR donor alone, demonstrating the importance of TALENs in mediating the HR event. Sequence analysis of the junction-PCR products demonstrated the seamless fusion of donor DNA with miR-21 genomic sequences (Fig. 2B), confirming the gene targeting ability of the designed TALENs and the HR donor vector.

enrichment and isolation of miR-21 knockout candidates from single cell-derived clones

To enrich and isolate miR-21 precursor-deleted cells, HEK293 cells cotransfected with the miR-21-specific TALEN pair and HR donor vector were put under positive selection.
Potential miR-21 HR-candidate cells were selected by puromycin resistance and RFP expression (Fig. 2C). Out of 96 single cell-derived clones, 30 lines survived in the presence of puromycin and remained RFP-positive and puromycin-resistant for >2 mo. In the group transfected solely with the HR donor vector, only six colonies were initially formed, and none of these clones survived after prolonged puromycin treatment.

**Genotyping after selection reveals highly efficient generation of bi-allelic knockouts**

We next determined the genotypes of the selected, single cell-derived clones. As shown in Figure 2D, correctly targeted colonies were revealed by the presence of puromycin and remained RFP-positive and puromycin-resistant for >2 mo. In the group transfected solely with the HR donor vector, only six colonies were initially formed, and none of these clones survived after prolonged puromycin treatment.

To determine whether a simultaneous modification of the other allele by NHEJ had occurred, we sequenced the PCR products spanning the targeted seed region of miR-21. Among 11 mono-allelic HR clones analyzed, a significant portion of the non-HR alleles (91%; 10/11) were NHEJ-modified (Fig. 2E). Important, and consistent with previous reports (Cermak et al. 2011), almost all modifications were centered on the targeted region, resulting in at least 4-bp deletions or complete changes of the seed sequences of miR-21. Taken together, these results show an 87% efficiency of our approach in generating bi-allelic modifications of miR-21 in the human genome.
whether custom TALEN-mediated miR-21 targeting may lead to any changes in the neighboring gene TMEM49 (also referred to as VMP1). We performed quantitative RT-PCR to examine the mRNA levels of TMEM49 in three miR-21 knockout cell lines and the parental control. We found that there was no significant change in the mRNA levels of TMEM49, compared to the control (Fig. 3B), confirming that neither TALEN-mediated gene replacement nor small deletions caused significant changes in the expression of this neighboring gene.

Loss of miR-21 causes an increase in target gene expression that can be rescued by re-expression of miR-21

To test whether miR-21 ablation caused corresponding changes in the expression of tumor suppressor gene PDCD4 (Programmed cell death 4), a known target of miR-21 (Liu et al. 2012), we performed Western blot analyses. As expected, while the basal PDCD4 protein level is barely detectable, it was drastically elevated following miR-21 knockout (Fig. 3C, lanes 1,2). To further validate whether the elevated expression of PDCD4 was due to the loss of miR-21, we transduced miR-21 knockout cells with lentiviral particles containing a cassette to express miR-21 and GFP. As shown in Figure 3E, more than ~80% of miR-21 knockout cells (RFP-positive) were also GFP-positive, suggesting expression of miR-21 in most transduced cells (Fig. 3D). This was confirmed by quantitative RT-PCR showing an ~3000-fold increase in miR-21 RNA expression in the transduced cells vs. the control (Fig. 3E). Following restoration of miR-21 expression in miR-21 knockout cells, PDCD4 protein was partially reduced (Fig. 3C).

miR-21 ablation results in inhibition of cell proliferation and alterations in global miRNA expression

We next analyzed cell proliferation rates following miR-21 ablation in HEK 293. As shown in Figure 3F, the three independent miR-21-ablated lines tested all exhibited reduced cell proliferation compared to the parental control. The inhibition in cell proliferation occurred as early as Day 3 after plating and became more prominent with time, lasting as long as 8 d. We further tested proliferation of cell line KO#5 with reintroduced miR-21. However, proliferation of KO#5 expressing miR-21 was not restored to the wild-type level, possibly due to the relatively lower expression levels of miR-21 compared to wild type (Fig. 3E), or to complex, nonreversible changes in the KO cell line. Nevertheless, we can’t rule out that the miR21 phenotype is due to possible off-target effects.

To examine whether the reduced proliferation was associated with changes in other miRNAs, we profiled global miRNA expression in three independent bi-allelic miR-21 knockout lines (clone #1, #5, #7), compared to the parental control. Seventeen out of 760 miRNAs tested (~2%) showed more than threefold up- or down-regulation in all three mutant lines, demonstrating strong concordance between the independent lines (Table 1). A BLASTn search against the NCBI nucleotide database confirmed that these miRNA do not display sequence homology to the TALEN target sequences, indicating that the observed changes are secondary in nature and not off-target effects.

DISCUSSION

Gene targeting of miRNAs in mammalian cells by homologous recombination is inefficient (Bollag et al. 1989), which has limited the use of human disease models in elucidating miRNA functions and exploring their therapeutic potential. To overcome these limitations, we developed an approach for efficient bi-allelic miRNA ablation combining TALENs with a selectable HR donor vector. We demonstrate that
First, we achieved robust bi-allelic gene modifications with an efficiency of 87%. By simple puromycin selection, we were able to establish more than 20 bi-allelic modified clones from one transfection experiment with a starting cell number of $2 \times 10^5$. Although most of the bi-allelic clones harbor one complete hairpin deletion and one seed region disruption, we speculated that the expression of miRNAs was effectively abolished due to the precise disruption of the miRNA seed region. In fact, both bi-allelic modified clones tested showed a complete loss of miR-21 expression, similar to those of the bi-allelic hairpin deletion. Another advantage of our HR donor vectors was the potential to induce mutations in the human genome at a frequency averaging around 16% (Kim et al. 2013a; Sakuma et al. 2013) by creating targeted double-strand breaks and subsequent NHEJ, the majority of mutations are mono-allelic and require time-consuming single cell-derivation and subsequent screening (Hu et al. 2013a). For example, an average mono-allelic efficiency of 16% would result in $\sim 0.04\%$ bi-allelic modifications. A low-efficiency TALEN of 2% mono-allelic disruption would result in rare bi-allelic modifications of $\sim 0.04\%$. To overcome these limitations, we employed an approach of combining TALENs targeting the miRNA seed region with an HR donor vector for deleting the entire miRNA stem–loop structure. The HR donor vector was engineered to include an insulated cassette for RFP and puromycin markers to allow positive selection by drug treatment and/or fluorescent-activated cell sorting. Our combined approach of miRNA gene targeting via TALEN and HR donor vectors was highly efficient and produced some intriguing results:

Figure 3. Expression of miR-21 and associated changes in TALEN-mediated knockout lines. (A) Complete loss of miR-21 expression in three independent lines. The constitutive expression of miR-21 in parental HEK293 cells was high with a CT value of 18–21. Quantitative PCR confirmed a $\sim 1000$-fold decrease in miR-21 levels in all three lines tested compared to those of the parental control. (B) No significant change in the mRNA expression level of TMEM49, the closest neighbor of MIR21 on chromosome 17, as assessed by real-time RT-PCR. (C) Expression of miR-21 was associated with corresponding changes in the expression of its targeted gene PDCD4; GAPDH served as loading control. Western blot was conducted with equal amount of proteins extracted from parental HEK293, miR-21 double knockout line, and double knockout line re-expressing miR-21. (D,E) Confirmation of miR-21 re-expression in the miR-21-knockout line. (D) The upper panel shows cells positive for RFP due to presence of the HR donor cassette (bi-allelic). Re-expression of miR-21 was confirmed by co-expression of a GFP marker (lower panel) and real-time RT-PCR (E). (F) Reduced proliferations in all three miR-21 knockout lines tested. Growth curves were plotted for the parental HEK293 cells as well as the three independent knockout lines.

Programmable nucleases, such as zinc finger nucleases (ZFNs), meganucleases, and, more recently, TALENs and CRISPR/Cas9-guided DNA endonucleases have emerged as powerful tools for targeted genome editing (Rahman et al. 2011; Hafez and Hausner 2012; Perez-Pinera et al. 2012; Cho et al. 2013; Cong et al. 2013; Gaj et al. 2013; Mali et al. 2013). Among these, TALENs offer several advantages for miRNA perturbation, including the ability to bind any sequence in the genome, which enables specific targeting of small sequences, such as the miRNA seed sequence ($\sim 6$ bp). Several recent studies indicate high specificity and low cytotoxicity of TALENs, compared to ZFNs and the CRISPR/Cas9 system (Fu et al. 2013; Kim et al. 2013b; Sun and Zhao 2013). In addition, their simple, modular design is easy to implement, due to their availability as an open-source technology.

However, there are some key limitations in the use of programmable nucleases to achieve gene ablation in diploid mammalian cells. While TALENs have the potential to induce mutations in the human genome at a frequency averaging around 16% (Kim et al. 2013a; Sakuma et al. 2013) by creating targeted double-strand breaks and subsequent NHEJ, the majority of mutations are mono-allelic and require time-consuming single cell-derivation and subsequent screening (Hu et al. 2013a). For example, an average mono-allelic efficiency of 16% would result in $\sim 2.6\%$ bi-allelic modifications.

Second, targeted miRNA inactivation is a powerful method capable of providing conclusive information about gene function in a cell-type or disease-specific manner. This will help to establish the crucial link between genotype and phenotype, thus yielding functional- and clinical-relevant knowledge. As expected, miR-21 knockout lines displayed an increased protein level of the miR-21 target gene.
PDCD4. Re-expression of miR-21 in miR-21-knockout cells restored its functional role, demonstrated by a decrease in protein expression of the target gene PDCD4, further confirming the validity of miR-21 knockout lines for functional analysis. Moreover, we observed a pronounced attenuation of cell proliferation in all three independent miR-21-ablated cell lines tested. Strikingly, miRNA profiling by qRT-PCR revealed large-scale changes of miRNA expression in the three independent knockout lines tested, highlighting the promise of this approach for future dissection of miRNA regulatory circuits. With the ease and efficiency of bi-allelic mutant generation, combined with the advantage of a stable phenotype, we envision that this approach will broaden our knowledge in deciphering the role of miRNAs in human physiology and disease.

**MATERIALS AND METHODS**

**Cell culture and transfection**

The human embryonic kidney cell line HEK293 cells were maintained in high glucose Dulbecco’s Minimal Essential Medium (DMEM) supplemented with 10% FBS, 2 mM GlutaMax (Life Technologies), 100 units/ml penicillin and 100 units/ml streptomycin. All transfections were performed using Purefection transfection reagent according to the manufacturer’s manual (Cat# LV750A-1; System Biosciences Inc.). Transfected cells were incubated in a 37°C incubator with 5% CO₂.

**TALEN design and TALEN expression plasmids**

To target the seed region of the human miR-21 locus (Fig. 1A), a TALEN pair was designed using an online tool, TAL Effector Nucleotide Targeter 2.0 (Doyle et al. 2012). To streamline the design of miRNA targeted TALENs, we followed these criteria: (1) TALEN binding sites were set to 20 bp, including the first T, to ensure high specificity of gene targeting; (2) spacer lengths of 15–25 bp were chosen to maximize cleavage efficiency; (3) the miRNA seed sequence was centrally situated within the spacer to direct cleavage to the seed region. Following these criteria, designed TALENs were assembled into a CMV-driven expression cassette using the EZ-TAL Assembly Kit (Cat# GE100A-1; System Biosciences Inc.), and the final constructs were confirmed by DNA sequencing.

**miR-21 gene targeting and screening**

A knockout HR donor vector that bears homologous arms of 509 bp (left arm) and 600 bp (right arm) was designed and constructed (Fig. 1B). In the genome, the regions represented on the vector arms are spaced 202 bp apart, flanking the miR-21 stem–loop region. The TALEN cut site is located 52 bp downstream from the left vector arm, within the miR-21 seed region. In the HR donor vector, the two homologous arms flank an insulated cassette to express dual selectable markers, RFP and puromycin resistance proteins. After transfection, RFP-positive and puromycin-resistant single cells were isolated and expanded. Briefly, a total of ~4 × 10⁵ cells were transfected with 1 µg of donor plasmid and 1 µg of each TALEN-encoding plasmid. Cells were trypsinized 2 d after transfection and subsequently plated on 10-cm culture dishes for 24 h. Puromycin was then added to the culture medium (final concentration 4 µg/mL), to allow single cell-derived colony formation. After 2 wk of puromycin selection, RFP-positive clones were picked and expanded for further analysis.

**Genotyping for HR and NHEJ events**

To confirm miR-21 gene disruption, the targeted gene loci were PCR-amplified from genomic DNA prepared from transfected cells or candidate clones. To detect HR events, two pairs of primers were used to perform junction PCR. PCR products with sizes of 830 bp and 1281 bp were expected in the case of successful HR, using the following primers—P2f: 5′-TGATGCATTGACACCCCTGG-3′; P2r: 5′-GAAAGATCTGATGCATTGCAGGAGG-3′; P3f: 5′-AGCCCGGAGATCTGACATTGCAGGAGG-3′; P3r: 5′-TGATGCATTGACACCCCTGG-3′ (Fig. 1C).

To determine whether mono- or bi-allelic events occurred in single cell-derived clones, a triple-primer PCR strategy was devised. As shown in Figure 1C, two forward primers (P3f1: 5′-TGATGCATTGACACCCCTGG-3′; P3f2: 5′-GAAAGATCTGATGCATTGCAGGAGG-3′; P3f3: 5′-AGCCCGGAGATCTGACATTGCAGGAGG-3′) and a common reverse primer (P3r: 5′-TGATGCATTGACACCCCTGG-3′) were used to amplify a predicted 1067-bp fragment of the wild-type allele and a 1281-bp fragment in the case of HR, respectively.

Even cells that carry only one HR event may carry bi-allelic modifications of the miRNA if the miRNA seed region of the other allele is modified by NHEJ. To test for small NHEJ-induced sequence changes on the other allele, genomic PCR was performed to amplify the targeted sites using the primers P1f and P1r. The amplified PCR products were determined to be consistent with the expected sizes for the wild-type allele.

**TABLE 1. Global miRNA profiling revealing 17 miRNAs differentially expressed by at least threefold in three independent miR-21 knockout lines tested**

| miR-21 | miR-21 | miR-21 | Up- or down-regulated |
|--------|--------|--------|-----------------------|
| KO line#1 | KO line#2 | KO line#3 |
| hsa-miR-7 | 1.9 | 2.4 | 3.2 | Up |
| hsa-miR-7 | 1.8 | 2.6 | 3.0 | Up |
| hsa-miR-7 | 15.8 | 14.2 | 8.8 | Down |
| hsa-miR-34b | 1.6 | 2.0 | 2.0 | Up |
| hsa-miR-34 | 1.9 | 1.6 | 2.4 | Up |
| hsa-miR-134 | 4.3 | 4.5 | 3.4 | Down |
| hsa-miR-220c | 1.7 | 2.2 | 2.0 | Up |
| hsa-miR-302e | 3.9 | 6.6 | 4.2 | Up |
| hsa-miR-340 | 2.5 | 3.5 | 6.1 | Up |
| hsa-miR-376a | 3.4 | 3.4 | 4.5 | Up |
| hsa-miR-376c | 2.3 | 2.0 | 3.2 | Up |
| hsa-miR-429 | 2.3 | 1.7 | 3.7 | Down |
| hsa-miR-454 | 3.5 | 2.6 | 2.2 | Down |
| hsa-miR-487b | 2.6 | 4.4 | 2.9 | Up |
| hsa-miR-513a-3p | 4.2 | 5.0 | 3.8 | Up |
| hsa-miR-513b | 5.7 | 5.5 | 4.8 | Up |
| hsa-miR-521 | 2.2 | 3.9 | 4.2 | Up |

1.6-fold after log₂ transformation, compared to wild-type HEK293 cells.

*Data are shown as normalized, log₂-transformed fold-change of expression.
products were subjected to agarose gel electrophoresis and DNA purification using the QIAquick Gel Extraction Kit (QIAGEN), and the purified DNA was sequenced.

Western blot analysis
Whole-cell extracts were prepared using M-PER lysis buffer (Pierce). The concentration of proteins was measured by Bradford assay (Amresco), and equal protein amounts were used for SDS-PAGE. Briefly, proteins from whole-cell extracts were separated and transferred onto nitrocellulose membranes. The membranes were blocked with 1× TBS-T with 5% nonfat dry milk (Bio-Rad), followed by incubation overnight at 4°C with 1:1000-diluted primary anti-rabbit PDCD4 antibody (Cell Signaling). For protein loading controls, rabbit anti-human GAPDH antibody (Abcam) was used at 1:2500 dilution in Superblock T20 buffer (Pierce). The blot was then probed with 1:10,000-diluted goat anti-rabbit IgG secondary-HRP antibody (Pierce). The signal was detected by Super Signal West Femto ECL (Pierce).

miR-21 rescue
To re-express miR-21 in a knockout line, we used a lentiviral gene delivery system. Briefly, an expression vector containing the miR-21 precursor construct (PMIRH21PA-1; SBI) was packaged into pseudo-lentiviral particles using LentiSuite (LV300A-1; SBI) (Mendenhall et al. 2012). The lentiviral particles (MOI = 5) were transduced into miR-21-knockout cells in the presence of the virus-transduction reagent Transdux (LV850A-1, SBI). The re-expression of miR-21 was confirmed by both co-expression of GFP and quantitative RT-PCR measuring mature miR-21 expression levels as follows.

Quantitative RT-PCR and global miRNA profiling
Total cellular RNA was prepared from cells using Trizol Reagent (Life Technologies) according to the manufacturer instructions. Total RNA was converted to cDNA using RNA-Quant (Cat# RA430A-1; SBI), and qRT-PCR was performed. A miR-21-specific forward primer (CSRA 640A-1, SBI) was used in combination with a universal reverse primer (cat# RA420AU-3, SBI). For the miR-21-neighboring gene TMEM49 (also referred to as VMP1), gene-specific forward and reverse primers (TMEM49-F: 3′-CCG CATAGGTCCATCTGAGC-5′ and TMEM49-R: 5′-TCAACAAA TCCAGGACACCGAG-3′) were used to evaluate mRNA expression levels. To perform global expression profiling of cellular miRNAs, mRNA-specific primers were obtained from the hsa-miRNome miRNA Profiler kit (Cat# RA660A-1; SBI) in combination with a universal reverse primer (SBI) according to the manufacturer’s instructions. We used the comparative threshold cycle (Ct) method to quantify the expression levels. The Cts were normalized to three housekeeping RNAs (human U6 snRNA, RNU43 snoRNA, and Hm/Ms/Rt U1 snRNA).

Phenotypic growth analysis
Parental HEK-293 cells and miR-21-knockout cell lines were seeded at 10,000 cells/well in a 24-well plate in culture medium as described. Growth was monitored by counting cells with a hemocytometer from each cell line on Day 3, 4, 5, and 8. Culture medium was changed at Day 3 and 6. The experiments were performed in triplicate.

Data collection and presentation
For live cell monitoring, cultured cells were monitored at various times under a fluorescence microscope. RFP or GFP live cell images were taken using the same exposure conditions and magnification within the group of comparison. For qPCR assays, all data are presented as mean ± SD (n = 3), unless stated otherwise.

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