Targeting non-coding RNAs with the CRISPR/Cas9 system in human cell lines

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ABSTRACT
The CRISPR/Cas has been recently shown to be a powerful genome-editing tool in a variety of organisms. However, these studies are mainly focused on protein-coding genes. The present study aims to determine whether this technology can be applied to non-coding genes. One of the challenges for knock-out of non-coding genes is that a small deletion or insertion generated by the standard CRISPR/Cas system may not necessarily lead to functional loss of a given non-coding gene because of lacking an open reading frame, especially in polyploid human cell lines. To overcome this challenge, we adopt a selection system that allows for marker genes to integrate into the genome through homologous recombination (HR). Moreover, we construct a dual guide RNA vector that can make two cuts simultaneously at designated sites such that a large fragment can be deleted. With these approaches, we are able to successfully generate knockouts for miR-21, miR-29a, IncRNA-21A, UCA1 and AK023948 in various human cell lines. Finally, we show that the HR-mediated targeting efficiency can be further improved by suppression of the non-homologous end joining pathway. Together, these results demonstrate the feasibility of knock out for non-coding genes by the CRISPR/Cas system in human cell lines.

INTRODUCTION
It is well known that the human genome is actively transcribed; however, there are only about 20,000 protein-coding genes (1), accounting for about 2% of the genome, and the rest of the transcripts are non-coding RNAs including microRNAs and long non-coding RNAs (lncRNAs). For instance, over 56,000 human lncRNAs have been identified to date (2); however, the biological function for most of them is not known. Therefore, there is a critical need for functional studies of lncRNAs. A most commonly used approach for gene functional study is knockdown by RNA interference (RNAi) which is mainly functional in the cytoplasm where RISC complexes are located (3). However, many lncRNAs are localized to the nucleus (4), which may make it difficult to achieve robust knockdown. Thus, genetic editing at the genomic level provides a better alternative because it targets the genomic DNA. There are several genetic tools available for this purpose, including zinc finger nucleases (ZFN) (5) and transcription activation-like element nucleases (TALEN) (6). For example, type II restriction enzyme FokI is often used as a cleavage domain in ZFN (7); similarly, engineered TAL effectors can also be fused to the cleavage domain of FokI to create TALENs for genome editing (8). Recently, a novel genetic engineering tool called clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated (Cas) system is more advanced because of easy generation and high efficiency of gene targeting (9). Importantly, it only requires changing the sequence of the guide RNA (gRNA); and it can be directly delivered into embryos, to generate sequence-modified animals (10–13). Furthermore, multiplexing capability of CRISPR/Cas makes it possible to target multiple genes simultaneously (12).

As a bacterial adaptive immune defense mechanism, the CRISPR/Cas9 system has been known for a long time. There are three types of CRISPR/Cas systems (14). Type II system has been well studied originally because it offered practical applications in the dairy industry to generate phage-resistant Streptococcus thermophilus strains (15). In addition, type II uses single effector enzyme such as Cas9 to cleave dsDNA. Because of this unique feature, CRISPR/Cas9 system has been extensively studied as a genetic engineering tool from bacteria to mammals.

Since the first report in the bacterial system (16), a large number of papers have been published on the utility of
CRISPR/Cas system in various eukaryotic organisms including yeast, drosophila, zebrafish, mouse and human cells (11,13,17–22). In the first landmark paper, Jinek et al. demonstrated that the dual-tracrRNA:crRNA, which occurs in nature, can be engineered as a single RNA chimera to function as just like dual-tracrRNA:crRNA, and is able to direct sequence-specific Cas9 dsDNA cleavage (16). Mali et al. reported a knockdown approach using human codon optimized Cas9 (hCas9) with custom gRNA (a chimeric form) in human cells (21). Using the endogenous AAVS1 locus as an example, they demonstrated that targeting rates reach from 2 to 25% depending on cell types used. A similar study showed that Cas9 nucleases can be directed by short RNAs to induce precise cleavage at endogenous genomic loci in human and mouse cells (19).

However, all of these studies are focused on protein-coding genes. Information about targeting non-coding RNAs is scarce. Although two reports used CRISPR/Cas9 to knockout microRNA in the zebrafish genome (23) and to target a maternally expressed lncRNA, Rian, in mouse model (24), none of them used homologous recombination (HR)-based selection system or a dual gRNA approach with two gRNAs expressed in the same vector. Therefore, the present study aims to explore an applicability of CRISPR/Cas system through an HR-based selection and/or dual gRNA approach to non-coding genes in human cell lines.

MATERIALS AND METHODS

Reagents

Primary antibodies were obtained as follows: PDCD4 monoclonal from Sigma-Aldrich (St. Louis, MO, USA); GAPDH from ProteinTech (Chicago, IL, USA). Secondary antibodies conjugated with IRDye 800CW or IRDye 680 were purchased from LI-COR Biosciences (Lincoln, NE, USA). Polymerase chain reaction (PCR) primers were purchased from IDT (Coralville, IA, USA) and siRNAs from Dharmacon (Lafayette, CO, USA).

Cell culture

HEK293, HEK293T, HCT-116, MCF-7 and LNCaP were obtained from ATCC (Manassas, VA, USA) and cultured in Dulbecco’s modified Eagle’s medium (DMEM) or RPMI-1640 with 10% fetal bovine serum (FBS) (Sigma). All media contained 2 mM glutamine, 100 units of penicillin/ml and 100 mg of streptomycin/ml. Cells were incubated at 37°C and supplemented with 5% CO₂ in a humidified chamber.

Construction of plasmids

The high fidelity enzyme Phusion (NEB, Ipswich, MA, USA) was used to amplify respective DNA fragments by PCR to make these constructs. gRNA design was based on CRISPR design (http://crispr.mit.edu/) or CHOPCHOP (https://chopchop.rc.fas.harvard.edu/). For single gRNA cloning we purchased a set of self-complementary oligos from IDT and then ligated into PrecisonX™ Cas9 Smart-Nuclease (Cat# CAS900A-1, System Biosciences, Mountain View, CA, USA) according to manufacturer’s protocol.

For dual gRNA cloning, we introduced dual gRNAs under H1 and U6 promoter along with gRNA scaffold downstream of hCas9 to generate a dual gRNA vector using multiplex gRNA cloning kit (Cat# CAS9-GRNA-KIT, SBI). To construct a donor vector for miR-21 or miR-29a, we used MCS1-LoxP-EF1a-GFP-T2A-Puro-P2A-hsvTK-pA-LoxP-MCS2 (Cat#: HR210PA-1, SBI). Left and right arms (~800 bp each) were amplified from genomic DNA using primers miR-21-L-Spe-5.1; miR-21-L-Bsa1–3.1; and miR-21-R-Bbs-5.2 and miR-21-R-Sal-3.2 (Supplementary Table S1), and sequentially cloned into the vector by ColdFusion cloning kit (Cat# MC101B-1, SBI). To construct a donor vector for UCA1, lncRNA-21A or AK023948, we used a pSK backbone carrying LoxP-EF1a-GFP-T2A-Puro-pA-LoxP with restriction enzyme BamH1 on the left side and EcoR I on the right side. Left and right arms were then sequentially cloned into the vector using ColdFusion cloning kit. All PCR products were verified by DNA sequencing.

Transfection of siRNAs/plasmid DNAs and selection

Cells were transfected with siRNAs using RNAfectin reagent (Applied Biological Materials, Vancouver, Canada) following the manufacturer’s protocol, as described previously (25). siRNA sequences have been described for Ku70 (26) and Lig4 and XRCC4 (27). The selection procedure was listed in Figure 1D. In brief, constructs carrying hCas9 and single gRNA or dual gRNAs, along with donor vector, were introduced into cells by transfection using DNAfectin (Applied Biological Materials). Next day, the cells were re-plated with fresh medium, and re-seeded when the cells became confluent. One week later, the transfected cells were subject to puromycin selection at concentrations of 0.5, 1.0 or 1.5 µg/ml depending on the type of cells used; the purpose was to maximize recombination and minimize the random insertion. Two weeks after selection, surviving cells were sorted by fluorescence-activated cell sorting (FACS) based on GFP signal into individual wells of 96-well plates and incubated for 2–3 weeks. The sorted clones were then expanded in 12-well plates, followed by extraction of genomic DNA or total RNA for genomic PCR or quantitative reverse transcription (qRT)-PCR using standard methods. To detect HR events, we performed junction PCR using primers derived from outside sequence and the donor vector sequence, and then verified them by DNA sequencing of junction PCR products.

RNA preparation and RT-PCR

For RT-PCR, we isolated total RNA using Direct-zol™ RNA MiniPrep (Zymo Research, Irvine, CA, USA) per the manufacturer’s protocol and used 0.5 µg RNA to synthesize cDNA by RevertAid Reverse Transcriptase (ThermoFisher) with random primers. Finally, the resultant cDNA was used for PCR reactions. PCR annealing temperature varied depending on the primers used. To detect miR-21, we used the polymerase A method, followed by SYBR Green qPCR as described previously (28). RNU1 and 5s RNA were used as an internal control.
Figure 1. Knockout of miR-21 in HEK293 cells. (A) Strategy for making Cas9 all-in-one vector targeting miR-21 precursor. Three gRNA target regions are shown at the top panel; a single miR-21 gRNA each was cloned into the vector at the bottom panel as detailed in the Materials and Methods section. (B) Detection of indels in miR-21 knockout cells by T7 endonuclease 1 or Surveyor nuclease assay, as detailed in the Materials and Methods section. The black arrow indicates wild-type band and the red arrows indicate mutant bands, with relative cleavage frequency shown at the bottom. (C) A donor vector for targeting miR-21 through homologous recombination (HR). pA, poly A signal. (D) A procedure of selection and identification of knockout clones. (E) miR-21 knockout increased colony number, as determined by colony formation assay for a mixed pool after puromycin selection. Values are means of ± SE (n = 3). **P < 0.01.

Western blotting

Cells were harvested and protein was extracted from cells as previously described (25). The protein concentration was determined using a protein assay kit (Bio-Rad, Hercules, CA, USA) and samples were separated in sodium dodecyl sulphate polyacrylamide gels.

Detection of mismatched duplexes by T7 endonuclease assay

HEK293 cells were transfected with hCas9 carrying miR-21 gRNAs as described above and then genomic DNA was isolated 72 h post transfection. Genomic region surrounding the target site for miR-21 was PCR amplified using Phusion high-fidelity DNA polymerase, and products were purified using Zymoclean™ Gel DNA Recovery Kit (Zymo Research, Irvine, CA, USA) following manufacturer’s protocol. A total of 200 ng of the purified PCR products were mixed with 2 µl 10 X Buffer 2 (NEB) and ultrapure water to a final volume of 20 µl, and subject to a re-annealing process to enable heteroduplex formation: 95°C for 10 min, 95–85°C ramping at –2°C/s, 85–25°C at –0.25°C/s and 25°C for 1 min. After re-annealing, products were treated with 0.5–µl T7 endonuclease I (NEB) at 37°C for 15 min and analyzed on 1.5–2% agarose gels.

Excision of loxP by Cre

Cre mRNA was synthesized using mRNAExpress mRNA Synthesis Kit (Cat# MR-KIT-1, SBI) according to user manual’s protocol. In brief, the Cre coding region was cloned into pMRNA vector. Then T7 promoter carrying Cre fragment was PCR amplified using primers the kit provided. Later, purified PCR product was used as a template to synthesis Cre mRNA. Cells were seeded at ~20% and grown overnight, and then transfected with 0.5–1 µg Cre
mRNA per well in a 12-well plate. Three days later the trans-
sected cells were transferred to large dishes (10 cm), and in-
dividual colonies were formed within 2 weeks. The excision
efficiency is dependent on transfection efficiency. Based on
our experience, about 50–60% colonies had lost the marker
genes, revealing non-green and puromycin sensitive.

Statistical analysis

Comparisons between groups were analyzed using the Stu-
dent’s t-test (two groups) or a one-way analysis of vari-
ance (ANOVA) followed by post hoc Tukey test (multiple
groups). Differences with \( P \) values less than 0.05 are con-
sidered significant.

RESULTS

Knockout of protein-coding genes is relatively easy because
a small deletion or insertion can disrupt the open read-
ing frame so that no functional gene product is produced.
However, non-coding genes are very different. Especially
for IncRNAs, a small deletion or insertion may not neces-
sarily cause functional knockout. To test the feasibility of
non-coding gene knockout with CRISPR
/sorry/ we chose two microRNAs, i.e. miR-21 (28) and miR-29a (29),
and three IncRNAs, i.e. UCA1 (30), IncRNA-21A (31) and
AK023948 (32) and tested them in various human cell lines.
For miR-21, we adopted an all-in-one system, i.e. hCas9
and a single gRNA in the same vector. Thus, three individ-
ual gRNAs were designed to target pre-miR-21 (Figure 1A,
top) and each of them was separately cloned into the vector
(Figure 1A, bottom). After introducing a single gRNA or
mixed three gRNAs into HEK293 cells by transfection, we
performed T7 endonuclease assay and found that each of
the three individual gRNAs or combination produced mis-
mached bands with targeting frequency of 17–39% (Figure
1B), suggesting that this approach is robust and miR-21 se-
quence is altered. However, we encountered a difficulty in
identifying complete knockout clones. There was no com-
plete miR-21 knockout among over 100 individual colonies
screened. Possible reasons for this are: (i) if transfection ef-
niciency is not 100%, those non-transfected cells will increase
the screening effort; (ii) many established cell lines are poly-
ploid. For example, HEK293 cells are hypotriploid. In this
case, the cut may have to be simultaneously made on three
chromosomes in order to get a complete knockout.

To facilitate the selection of complete knockout clones,
we generated a donor vector in such a way that target
fragment is replaced by marker genes (GFP and PU, the
puromycin resistance gene, and TK, thymidine kinase gene)
once it is integrated into the genomic DNA by HR (Fig-
ure 1C). GFP enables visualization of the transfected cells
and cell sorting, and puromycin resistance gene allows se-
lection of positive stable transfectants carrying targeted se-
quencies by HR. TK is for selection of positive clones af-
after the marker genes are excised at loxP sites by Cre when
it is needed; it also helps to eliminate some remaining
clones with random integration. Outside the markers, we
introduced an ∼800-bp DNA fragment from targeting sites
(left arm and right arm) for HR (Figure 1C). These vec-
tors were simultaneously introduced into cells by transfec-
tion and then subject to puromycin selection as outlined
in Figure 1D. To reduce the possibility that the donor se-
quence randomly integrate into the genome, we did not start
puromycin selection until 1 week after transfection. We first
tested targeting efficiency by counting surviving colonies in
the presence of puromycin (1 \( \mu \)g/ml). Co-transfection of
miR-21 gRNAs and donor vectors in HEK293 cells resulted
in an ~6-fold increase of colony number compared to vec-
tor control cells (Figure 1E).

These surviving cells were further subject to cell sorting
by FACS based on GFP signal into 96-well plates, followed
by expansion in 12-well plates. We recovered 10 individual
clones from a 96-well plate. qRT-PCR analysis revealed that
five clones were complete miR-21 knockouts because over a
1000-fold reduction in miR-21 level was detected for each of
them (Figure 2A). Since the donor vector carries two loxP
sites, we introduced Cre mRNA into each of them. As ex-
pected, excision of the marker genes generated non-green
cells (Figure 2B). To determine the nature of sequence al-
terations for these five clones, we designed primers miR-
21–5.1 and miR-21–3.1 which would produce an amplicon
size of 614 bp for wild type (WT). As shown in Figure 2C,
clones #1 and #2 each generated two bands, whereas the
parental cells produced a single band (WT). DNA sequenc-
ing showed that these two clones were duplicates because
the sequences for each of the two bands were exactly the
same for clones #1 and #2. The top band carried a 14-bp
deletion (ATGGGCTGTCTGAC) along with a 46-bp in-
sertion, leading to a 32-bp net insertion; bottom band car-
died a 72-bp deletion (pre-miR-21), and one copy of loxP
site (34 bp) was retained (Figure 2D). For clones #3 and #4,
we detected two different sequences (Supplementary Fig-
ure S1) in addition to the loxP site copy, whereas clone #5
carried an 88-bp insertion. These results are consistent with
the triploidy nature of this cell line. While one copy was re-
placed by donor vector through HR (i.e. the loxP copy), the
rest two copies (# 3 and #4) had a deletion or insertion,
suggesting that these cells have a strong non-homologous
end joining (NHEJ) activity. Although these clones are ‘het-
erozygous’ by nature, they are complete knockouts for miR-
21. In addition to miR-21, we also tested miR-29a knock-
out in HEK293 cells with the same approach and we were
also able to obtain complete miR-29a knockout clones (not
shown).

We then tested miR-21 knockout in colon cancer HCT-
116 cells that are mainly diploid (www.atcc.org). miR-21
knockout caused over a 5-fold increase in colony num-
ber compared to vector control (Supplementary Figure
S2A and B), which is comparable to the number of
HEK293 knockout cells (Figure 1E). After cell sorting, the
puromycin-resistant GFP-positive clones were picked out
and expanded. Among over 20 clones, we identified seven
clones with over 90% reduction of miR-21 by qRT-PCR
(Figure 3A). Clones #2 and #17 were subject to further
characterization. DNA sequencing of PCR products re-
vealed that clone #2 carried a 198-bp deletion but with a
195-bp insertion; clone #17 carried a 36-bp deletion (Fig-
ure 3B; Supplementary Figure S2C). To determine the func-
tion of miR-21 knockout, we performed colony forma-
tion assays and found that cell growth of these clones was
substantially slower than that of the control cells after 1-
week incubation. Representative pictures of crystal violet-
Figure 2. Identification of miR-21 KO clones and elimination of marker genes. (A) Detection of miR-21 level by qRT-PCR. Over a 1000-fold reduction was detected in all of the five miR-21 KO clones with $P$ value <0.01 for all five clones. (B) Excision of marker genes in miR-21 KO clone #1 cells by Cre. Shown here are representative images. (C) Both miR-21 KO clones #1 and #2 carry distinct PCR products using primers miR-21–5.1 and miR-21–3.1. (D) DNA sequences for each of the PCR products from clones #1 and #2. Both clones revealed identical sequences. Insertion sequences are shown in blue and loxP sequences are in red.

stained colonies are shown in Supplementary Figure S2D. The percentage of colony formation of miR-21 knockout clones decreased to 21.1%, and 33.9% in clones #2 and #17, respectively, compared to control (Figure 3C). Moreover, we found that two known miR-21 target genes, growth arrest-specific 5 (GAS5) (33) and programmed cell death 4 (PDCD4) (34), were significantly induced in the miR-21 knockout clones, as detected by qRT-PCR (Figure 3D) and western blot (Figure 3E), respectively, demonstrating that they are functional. Finally, to determine HR events, we performed junction PCR using primers miR-21-recom-5.2 and EF1-seq-3.2 (Supplementary Figure S3A) and found that while no PCR product was detected for donor + vector, HR rate was about 74% (14/19) for donor + miR-21 gRNAs (Supplementary Figure S2A). DNA sequencing of three randomly selected PCR products confirmed the HR event and all three products revealed identical sequences (Supplementary Figure S2B).

In addition, we also made miR-21 knockout in prostate cancer LNCaP and breast cancer MCF-7 cells. Knockout of miR-21 in LNCaP cells led to a 1.67-fold increase in colony number (Supplementary Figure S4A). Expression of miR-21 was decreased by 39.6% for a mixed pool, as detected by qRT-PCR (Supplementary Figure S4B). Similarly, knockout of miR-21 in MCF-7 cells resulted in a 2.77-fold increase in colony number (Supplementary Figure S4C) and miR-21 was decreased by 37.5% for a mixed pool (Supplementary Figure S4D).

Having demonstrated the successful knockout of miR-21, we then determined whether the CRISPR/Cas system can be applied to lncRNA genes, including UCA1, lncRNA-21A and AK023948. UCA1 has been shown to be a potential oncogene (30) and it contains three exons. We initially designed a single gRNA targeting UCA1 exon 1 (Supplementary Figure S5A). Colony formation assays with a mixed pool revealed that there was a 1.98-fold increase in colony number in UCA1 knockout compared to vector con-
Figure 3. Characterization of miR-21 KO clones in HCT-116. (A) Detection of miR-21 in miR-21 KO clones by qRT-PCR; *P value <0.01 for all seven clones compared to vector control. miR-21 KO #2 and #17 (in green) were selected for further analysis. (B) Detection of alterations at the target region of miR-21 KO #2 and #17 by DNA sequencing. (C) miR-21 knockout suppresses cell growth, as determined by colony formation assays. (D) miR-21 knockout increases the miR-21 target GAS5, as detected by qRT-PCR. (E) miR-21 knockout increases the miR-21 target PDCD4, as detected by western blot. Values in (C) and (D) are means of ± SE (n = 3). *P < 0.05 and **P < 0.01.

trol (Figure 4A). Genomic PCR identified seven potential clones among 50 randomly picked single clones; qRT-PCR revealed that UCA1 level for clones #2 and #4 was reduced by about 90% while UCA1 level for clones #14, #16 and #19 was reduced by about 50% (Figure 4B). However, genomic PCR using primers derived from exon 1 region (UCA1-inside-5.3 and UCA1-inside-3.3) (Supplementary Figure S5B) revealed a positive band for all five clones (Figure 4C). DNA sequencing of this band identified a single nucleotide insertion in red (Figure 4D) for clones #2 and #4, respectively, whereas the band from clones #14, #16 and #19 was wild type, again, suggesting that NHEJ takes place in addition to HR.

These findings suggest that although at least one copy of donor sequence is integrated into genome by HR, not all cells have homozygous recombinants at UCA1 site. Instead, we were still able to detect the targeted UCA1 region by genomic PCR (Figure 4C), suggesting that the cells carry one copy of recombinant and one copy wild type or altered sequence at UCA1 site if it is diploid. It is evident that altered copy due to deletion or insertion may still produce UCA1, as suggested by qRT-PCR. Since microRNA is relatively short, a small deletion can cause disruption of mature microRNA sequence. However, for lncRNAs, such small deletions or insertions may not be sufficient to completely disrupt their expression.

To overcome this problem, we constructed a dual gRNA system (Figure 5A, top), which carries two tandem gRNAs, under H1 and U6 promoters, respectively, in the same single vector, allowing co-expression of two gRNAs and hCas9. While gRNA1 targeted toward 5' end of UCA1 exon 1 (same as single gRNA as shown in Supplementary Figure S5A), gRNA2 targeted toward 3' end of exon 3 such that almost the entire UCA1 sequences can be deleted through NHEJ or replaced by HR mechanism (~5.6 kb; Figure 5A, bottom). In addition, we made a second single gRNA (same as gRNA2). Co-transfection of donor and UCA1 dual vector resulted in a 2.73-fold increase in colony number compared to control (Figure 5B). Of interest, the colony number from the UCA1 dual gRNA system was higher than ei-
Figure 4. Generation of UCA1 knockout in HCT-116 cells. (A) Colony formation for vector and UCA1 KO for a mixed pool after puromycin selection. (B) Detection of UCA1 expression in UCA1 KO clones by qRT-PCR. (C) Detection of UCA1 target region in all five clones using genomic DNA as a template and primers UCA1-inside-5.3 and UCA1-inside-3.3. (D) Detection of a single nucleotide insertion (in red) in UCA1 KO#2 and #4, by DNA sequencing. The arrow indicates the cleavage site by Cas9. Underlined GGG is PAM sequence. Values in (A) and (D) are means of ± SE (n = 3). **P < 0.01.

We also tested two additional lncRNAs, IncRNA-21A and AK023948. We designed a dual gRNA targeting a 475-bp fragment covering the IncRNA-21A sequence (Supplementary Figure S6A). Co-transfection of donor and IncRNA-21A dual vectors into HCT-116 cells resulted in a 16.3-fold increase in colony number compared to control (Supplementary Figure S6B). Using primers derived from either outside or inside the IncRNA-21A dual-gRNA-targeted sides (Supplementary Figure S6C) we detected one copy was knockout in clone S7; genomic PCR showed that wild-type IncRNA-21A was completely lost in clones M2 and S38 (Supplementary Figure S6D). Junction PCR confirmed that all three clones carried HR fragments and they all had identical sequences, as determined by DNA sequencing (Supplementary Figure S6E). Similarly, we were able to knockout AK023948 in MCF-7 cells (Supplementary Figure S7). Therefore, our dual gRNA approach increases the targeting efficiency by CRISPR/Cas9 system, demonstrating its feasibility for targeting lncRNAs with ei-
Figure 5. Targeting UCA1 with a dual gRNA approach. (A) Strategy for targeting UCA1 with dual gRNAs. (B) Colony formation for vector and UCA1 KO after puromycin selection. (C) The dual gRNA approach increases colony number, as compared to single or two individual gRNAs in combination. (D) Detection of UCA1 deletion by genomic PCR using primers UCA1-outside-5.5 and UCA1-outside-3.5. The red arrow (2.8 kb) indicates a PCR product corresponding to a replacement and the black arrow indicates a deletion (227 bp). Clones #3, #12 and #18 are expected to have a 2.8-kb fragment, but missing here possibly because a deletion band is much smaller, and thus it is more favorably amplified. Indeed, junction PCR with primers UCA1-Recom-5.1 and EF1−seq-3.2 revealed that clones #3, #5 and #12 all carried the HR fragment (bottom). (E) Detection of UCA1 expression in UCA1 KO clones by qRT-PCR. P value <0.01 for all six clones, as compared to vector control. (F) No PCR product (bottom) except for vector control (220 bp) was detected in UCA1 KO clones using primers derived from the region as indicated on top (UCA1-intron-5.1 and UCA1-intron-3.1). Values in (B) and (C) are means of ± SE (n = 3). *P < 0.05 and **P < 0.01.

These findings further support the notion that established human cell lines have a strong NHEJ activity relative to HR because even under selection pressure, we are still able to see a significant frequency of deletion. Therefore, we asked if we can increase the HR targeting frequency by suppressing genes involved in NHEJ. We chose three key genes, Ku70, Lig4 and XRCC4, that have been implicated in NHEJ pathway (35–37). Thus, we used siRNA to transiently knockdown Ku70, Lig4 and XRCC4 simultaneously. First, we confirmed that the expressions of Ku70, Lig4 and XRCC4 were knocked down. The siRNAs resulted in at least 23.2% or 37.3% decrease in the expression of Ku70, Lig4 and XRCC4 compared to control siRNA in HEK293T cells (Supplementary Figure S8A–C) or HCT-116 cells (Supplementary Figure S8D–F). To determine effect of suppression of NHEJ pathway on the colony formation, we conducted either microRNA (miR-21 gRNAs) or lncRNAs (UCA1 dual gRNA and lncRNA-21A dual gRNA) transfection 24 h following the siRNA treatment. As anticipated, knockdown of Ku70, Lig4 and XRCC4 by siRNAs led to a 2.4-fold increase in Cas9-mediated colony number compared to control siRNA for miR-21 (Figure 6A and B). To determine whether the colony number reflects HR events, we performed junction PCR using primers miR-21-recom-5.2 derived from sequence outside the HR event and EF1-Seq-3.2 derived from the donor vector (Supplementary Figure S3A). DNA sequencing of the junction PCR products confirmed the HR events (Figure 6C). We obtained 75% HR rate for control siRNA and 78% HR rate for KLX siRNAs. Similarly, UCA1 dual gRNA knockout under the suppression of NHEJ pathway resulted in a 2.2-fold increase in colony number in HCT-116 cells (Figure 6D and E). DNA sequencing of the junction PCR products also indicated roughly the same HR rate for siRNA or KLX siRNAs (Figure 6F), suggesting that relative colony number can represent HR events. LncRNA-21A dual gRNA knockout fol-
following the transient knockdown of Ku70, Lig4 and XRCC4 resulted in a 1.2-fold increase in colony number in HCT-116 cells (Supplementary Figure S9). Together, these results suggest that the Cas9-mediated targeting frequency through HR can be improved by suppressing the NHEJ pathway.

**DISCUSSION**

Non-coding genes represent the vast majority of human transcripts. However, little is known about the physiological functions for most of them, especially lncRNAs. Thus, the present study aims to explore the applicability of CRISPR/Cas9 to non-coding genes for their functional studies. Unlike protein-coding genes, non-coding genes such as lncRNAs lack open reading frames, although their molecular weight is comparable to that of protein-coding genes. Because of this, the commonly used approaches of CRISPR/Cas9 system may have limited utility for knock-out of non-coding genes. Here we demonstrate that a selection approach with a donor vector for HR can significantly
increase targeting efficiency. In particular, the dual gRNA method allows us to target various sizes of a given DNA segment. Moreover, suppression of NHEJ pathway can further increase HR to improve target efficiency with our dual gRNA and donor vector. In addition to non-coding genes, this platform can also be applied to protein-coding genes. Given that about 30% of chance for any small insertion or deletion does not change the downstream open reading frames, screening effort for knockout clones can be significantly reduced by this approach. Finally, our platform can be easily adapted to other types of cells of different organisms.

Compared to lncRNAs, microRNAs are relatively easy to knock out. Our results suggest that the knockout efficiency of microRNAs is higher than that of lncRNAs (Figures 2A and 4B). This may be attributed to the fact that microRNA genes are much smaller in size than lncRNAs; mature microRNAs are ~22 nucleotides in length. Of interest, although we target pre-miR-21, DNA sequencing reveals that alteration of the precursor can also lead to knockout of miR-21, which may be due to disruption of microRNA processing. On the other hand, lncRNAs are much larger than microRNAs. A previous study has shown that MALAT1 can be knocked out using ZFN or TALEN by introducing a poly A to terminate the downstream transcription (38). Although our donor vector also carries a poly A signal, apparently, this is not sufficient because a strong NHEJ activity can produce deletions or insertions. This ‘heterozygous’ feature is more prominent in polyplloid cells, including most cancer cell lines.

Our results further suggest that HR combined with a donor vector can greatly increase the knockout efficiency. Due to the selection that kills non-transfected cells, transfection efficiency is no longer an important factor. This feature is critical because there are many cell lines with a relative low transfection even with different transfection reagents. Moreover, this HR system may reduce off-target frequency by CRISPR/Cas9 (39,40) because the donor is less likely to combine into the genome at off-target sites due to lack of homologous flanking DNA regions and thus, those cells carrying off-target sites, but without donor vector, are expected to be lost during puromycin selection.

Although we used FACS-sorted single cell colonies for most of our experiments, we noticed that manually picked single colonies in a 10 cm dish can also generate relatively pure clones as long as the starting cell density is low. The benefit of this approach is evident, i.e., cost saving and less stress to the cells.

One feature of this study is a dual gRNA approach which is especially useful for lncRNA knockout. For instance, we detected a significant portion of deletion with the dual gRNAs at the pre-designed sites. However, with a single gRNA or combination of two gRNAs in separate vectors, we rarely detected measurable deletions, as determined by genomic PCR. In consistent with this, targeting efficiency as measured by surviving colony number with the dual gRNA approach is higher than that with two single gRNAs which have the same target sequences. Although there are several possibilities for this, one of them may be that two gRNAs in the dual system are always present in the same cells. Additional advantage of using the dual gRNA system is the significant improvement of screening procedure. With appropriate primers, it is much easy and fast to identify potential knockout clones. For example, we are able to obtain at least 74% HR rate for miR-21, UCA1 and lincRNA-21A (Supplementary Figure S10).

Since our donor vector carries two loxP sites, it is possible to make a double or triple knockout sequentially. This can be accomplished by excising out the marker genes in the previous knockout such as miR-21 knockout (Figure 2B). Once excision, the cells are no longer green and they are sensitive to puromycin, which makes the selection and cell sorting possible again.

Finally, our results suggest that the targeting efficiency can be further improved by suppression of NHEJ pathways. This approach has been successfully studied in human somatic cells. For instance, suppression of Ku70 and XRC4 by RNAi can greatly enhance gene targeting efficiency (41). We suppress three genes (Ku70, Lig4 and XRC4) simultaneously and the targeting efficiency is significantly increased, suggesting that decreased efficiency in NHEJ-mediated knockout can potentially benefit the HR-mediated knockout. Nevertheless, it remains to be determined whether targeting efficiency can be further increased by knockdown of various NHEJ genes individually or in different combinations.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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