**sdRNA: siRNA with a DNA Seed for an Efficient and Target-gene Specific RNA Interference**

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**Abstract**

RNA interference (RNAi), a process through which small interfering RNAs (siRNAs) induce sequence-specific post-transcriptional gene silencing, is commonly recognized as a powerful tool not only for functional genomics but also for therapeutic applications. To achieve accurate target gene function and successful therapeutic applications, it is necessary to select an efficient and target gene-specific siRNA with minimal off-target effects. We found that the ability to induce off-target effects on unintended genes is strongly correlated to the thermodynamic stability of the duplex formed between the seed region (positions 2-8 from the 5' end of the siRNA guide strand) and target mRNA. Consistent with this property, we found that DNA-RNA chimeric siRNA (chiRNA) with deoxyribonucleotides in the 5' proximal eight nucleotides of the guide strand and the complementary nucleotides in the passenger strand exerted virtually no off-target effect due to low stability of the DNA-RNA duplex in seed-target base-pairing. However, the corresponding RNAi activities for primary target genes were also decreased to one-tenth on average by the DNA substitutions. Here, we report that siRNAs with seven deoxyribonucleotides exclusively in the seed region (sdRNA) may exhibit efficient target-specificity, but off-target effect-reduced RNAi activity.

**Keywords**: DNA modification; Off-target effect; Seed region; Small interfering RNA; siRNA; RNA interference; RNAi

**Introduction**

RNA interference (RNAi) is an evolutionarily conserved pathway induced by small interfering RNAs (siRNAs), which are 19 bp in length, double-stranded RNAs (dsRNAs) with two-nucleotide 3’ overhangs [1]. siRNAs incorporated into cells are transferred to an RNAi effector complex called an RNA-induced silencing complex (RISC) [2,3]. RISCs are assembled on one of the two strands of the siRNA duplex and are activated upon removal of the passenger strand [4-6]. Others reported that asymmetric features at both siRNA terminals are common to functional siRNAs [6-8]. An RNA strand with an unstable 5’ terminus is easily retained in a RISC. An activated RISC is a ribonucleoprotein complex consisting minimally of a core protein, Argonaute (Ago) and an siRNA guide strand [9-11]. The siRNA guide strand retained in an RISC recognizes mRNAs with perfectly complementary sequences and the Ago2 protein cleaves them.

The RNAi effect of a siRNA has been assumed to be extremely specific. However, accumulated evidence has revealed that siRNAs may downregulate many unintended genes with partial complementarities mainly in the seed region (nucleotides 2-8 from the 5’ end of the siRNA guide strand) [12-16]. This phenomenon is referred to as an off-target effect. The recognition mechanism of an off-target effect is known to be similar to that of microRNA-mediated gene silencing [13,15-21]; seed nucleotides are present on the Ago surface in a quasi-helical form to serve as the entry or nucleation site for small RNAs in the RISC [12,22,23]. To achieve accurate target gene function and successful therapeutic applications, it may be critical to select a target gene-specific siRNAs with minimal off-target effects.

We found that ability to induce an off-target effect is strongly correlated to the thermodynamic stability of the duplex formed between the seed region of siRNA guide strand and target mRNA (seed-target duplex) (Figure 1a) [12]. Furthermore, we have developed a DNA-RNA chimeric siRNA (chiRNA) with deoxyribonucleotides in the 5’ proximal eight nucleotides of the guide strand and the complementary nucleotides. The chiRNA showed virtually no off-target effects, probably because the stability of the seed-target duplex was reduced by the DNA substitutions in the siRNA seed region (Figure 1b) [16]. For target-specific RNAi with reduced off-target effects, we propose an improved chiRNA, referred to as an sdRNA, in which seven nucleotides in the seed region is exclusively replaced with DNA without including a nucleotide at the 5’ terminus (Figure 1c).

**Materials and Methods**

**siRNA**

RNAi efficiency in mammalian cells varies considerably depending on the siRNA sequence. We have shown that highly functional siRNAs for mammalian RNAi have A or U residues at nucleotide position 1 measured from the 5’ end of the guide strand, four to seven A/Us at nucleotide positions 7-17 (AU ≥ 57%) and a G/C at position 19 (Figure 1) [6]. In addition, a GC stretch of no more than nine nucleotides is contained in siRNA sequences. All of the double-stranded small RNAs used in this study shown in S1 simultaneously satisfied these conditions.

**Cell culture and silencing assay**

Human HeLa cells were cultured and subjected to gene silencing as described previously [6,12]. Briefly, a 1 ml suspension of human HeLa cells (1×10⁵ cells/ml) was inoculated in a 1.5 cm well 24 h prior to transfection. The cells were transfected with pGL3 or pGL2 control (1 µg; Promega), both of which encode firefly luciferase (luc), and the Renilla luc-encoding plasmid psiCHECK-completely matched (CM) or seed matched (SM) (0.1 µg; Promega) with siRNA, chiRNA, sdRNA, or sd/chiRNA. Lipofectamine 2000 regent (Invitrogen) was used for transfection. Each of siRNA, chiRNA, sdRNA, or sd/chiRNA (0.5, 5, 50, 500, 5,000, and 50,000 pM) was diluted in 50 µl of Opti-MEMI without fetal bovine serum (FBS) and mixed with 50 µl of Opti-MEMI containing 1µl of Lipofectamine 2000 for 30 min in room temperature. After the culture medium was changed with FBS-free medium, 100 µl of siRNA, chiRNA, sdRNA, or sd/chiRNA-Lipofectamine 2000 complex were added to each well. After 4 h incubation, the medium was replaced with fresh medium with FBS. The cells were harvested.

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24 h after transfection and their luc activity was measured using a Dual-Luciferase Reporter Assay System (Promega). The siRNAs used in this study were chemically synthesized (Sigma) (see S1). siGY-441, an siRNA for GFP knockdown, was used as a control. IC₅₀ was determined from the results shown in figures 2 and 3 by using the following formula:

\[ IC_{50} = 10^{\left(\log\left(\frac{A}{B}\right)\times(50-C)\times(D-C)\times\log(B)\right)} \]

where in A: of the two points on each Figure which bracket 50% inhibition, the higher concentration of siRNA, B: of the two points on each Figure which bracket 50% inhibition, the lower concentration of siRNA, C: inhibitory activity (%) at the concentration B, D: inhibitory activity (%) at the concentration A.
Plasmid construction

All plasmids constructed were derivatives of psiCHECK-1 (Promega). Chemically synthesized oligodeoxynucleotides, including one copy of the 23-bp CM target sequences (S2) or three copies of the SM target sequences (S3) with cohesive XhoI/EcoRI ends, were inserted into the psiCHECK-1 (Promega) XhoI/EcoRI site, which is situated in the region encoding the 3' UTR of Renilla luc mRNA; they are referred to as psiCHECK-CM and SM, respectively. The inserted CM or SM targets were expressed as part of the 3' UTR of Renilla luc mRNA in the transfected cells. The CM target completely matched the siRNA guide strand (Figure 2a), whereas the SM target consisted of two parts: the eight 3' terminal nucleotides of the target were complementary in sequence to the 5' end (nucleotide position 1) and the seed (positions 2-8) of the corresponding siRNA guide strand, while the remaining thirteen nucleotides were totally non-homologous to the guide strand (Figure 3a).

Results and Discussion

Designing sdRNA

We developed an improved method using siRNAs with DNA in
the seed region (sdRNA), in which ribonucleotides were replaced with DNA at positions 2-8 from the 5' end of the siRNA guide strand but a ribonucleotide at the 5' terminal end was not replaced with DNA. The complementary ribonucleotides of the guide strand seed region in the passenger strand were also replaced with DNA (Figure 1c). The sdRNA was expected to have strong target-specific RNAi activity and weak off-target effect activity, because it had a DNA seed sequence that could form a DNA-RNA seed-target duplex with low stability compared to that formed by an RNA-RNA duplex; the other positions (at 1 and 9-21) were ribonucleotides. In particular, A or U ribonucleotide residues at the 5' end of the guide strand might be important because the A/U nucleotide itself at the 5' terminal was shown to contribute strongly to RNAi activity. This observation might be explained by the fact that the nucleotide monophosphates AMP and UMP bind to Ago2 with up to 30-fold higher affinity than that observed with either CMP or GMP [24].

Comparison of the on-target RNAi activities of siRNAs, chiRNAs, sdRNAs, and sd/chiRNAs

To compare the RNAi activities of siRNAs, chiRNAs, sdRNAs, and sd/chiRNAs
The IC50 values of the siRNAs were from 19.0-336.1 pM, a decrease of 14-49% compared to the IC50 values of the nonmodified siRNAs varied from 9.4-105.6 pM depending on the sequence, indicating that siRNAs used in this study were 37-73% of the siRNA values (Figure 2b-e), suggesting that the 5’ end ribonucleotide in the guide strand played an important role in the RNAi activity. These results show good agreement with the previously reported result that AMP and UMP bind to Ago2 with up to 30-fold higher affinity than either CMP or GMP [24].

**Comparison of the off-target effect activities of the siRNAs, chiRNAs, sdRNAs, and sd/chiRNAs**

To compare the off-target effect activities of siRNAs, chiRNAs, and sdRNAs (Figure 3a), four different sequences, CLTC2416, FL2-153, FL3-36, and VIM270, were used. Off-target effect activities were measured using the psicHECK-SM reporter system. These highly functional siRNAs, siCLTC2416, siFL2-153, siFL3-36, and siVIM270, showed off-target effects in a dose-dependent manner until the siRNA concentration reached 50 nM (Figure 3b-e). The calculated IC50 values of the off-target effects in the nonmodified siRNAs, siCLTC2416, siFL2-153, and siVIM270, varied from 46.0-1261.5 pM, and that of siFL3-36 was not determined. However, the off-target effects of the chiRNAs were considerably reduced as shown in our previous report [16]; their off-target effects were reduced to undetectable levels, and the IC50 value of chiVIM270 was decreased to 0.6% (7968.7 pM) of the IC50 value of siVIM270. Previously, we showed that ability to induce an off-target effect is correlated to the thermodynamic stability of seed-target duplex of siRNA [12]. The melting temperature value of RNA-RNA duplex at seed positions 2-8 (Tm2-8) of siCLTC2416 is 33.2°C, which decreases to 12.7°C when DNA-RNA duplex is formed by chiCLTC2416, indicating the stability of seed-target duplex is strong for siCLTC2416 but weak for chiCLTC2416 consistent with the results shown in figure 3b. Similarly, Tm2,3 values of RNA-RNA duplexes formed between siFL2-153, siFL3-36, and siVIM270 and each target mRNAs were 21.0-26.2°C, and the values of DNA-RNA duplexes formed between chiFL2-153, chiFL3-36, and chiVIM270 and each target mRNAs decreased to 3.1-18.9°C (Figure 3c-e) [12]. Since the Tm2,3 values in the duplexes formed between sdRNAs and target mRNAs are same as those of DNA-RNA duplexes formed between chiRNAs and target mRNAs, it was unexpectedly that the sdRNAs showed strong off-target effects compared to those of chiRNAs. The IC50 values of sdFL2-153 and sdVIM270 were decreased to 41% (3045.0 pM) and 1.7% of the IC50 value of each siRNA, respectively. As shown in figure 4b, the three siRNAs showed the strongest off-target effect activities, while the chiRNAs showed the weakest off-target effect activities; the sdRNAs showed intermediate levels of off-target effect activity. The decreases in off-target effect activities of the sdRNAs compared to those of the siRNAs might be caused by the DNA substitutions in the seed region, suggesting that decreased stabilities in the seed-target duplexes reduced the off-target effects. However, the off-target effects of the sdRNAs increased compared to those of the chiRNAs, suggesting that the 5’ end ribonucleotide of the guide strand and three ribonucleotides at the 3’ end of the passenger strand contributed to the increase in off-target effects. Furthermore, the sd/chiRNAs showed weak or similar levels of off-target effects compared to those found with sdRNAs (Figure 4b).

This result suggests that the 5’ end RNA of the guide strand contributed strongly to the off-target effect, while the three ribonucleotides at the 3’ end of the passenger strand contributed weakly to this effect.
Conclusions

Based on our previous reports, we expected that an sdRNA that included a DNA seed would be an appropriate tool for inducing target-specific RNAi with reduced off-target effects. Our results confirm that the sdRNA showed weak off-target effect compared to siRNAs and strong RNAi activity compared to chRNAs. A 5' end ribonucleotide in the siRNA guide strand contributed strongly to the off-target effects, whereas the three ribonucleotides at the 3' end of the siRNA passenger strand contributed weakly to the off-target effects. Our results indicate that sdRNAs might be a powerful tool for developing ideal RNAi experiments and therapeutic applications.

Supplementary Information Files

S1. Supplementary Information 1. RNA sequences.
S2. Supplementary Information 2. Target sequences in psiCHECK-CM.
S3. Supplementary Information 3. Target sequences in psiCHECK-SM.

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