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Software

siDirect 2.0: updated software for designing functional siRNA with reduced seed-dependent off-target effect

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Abstract

Background: RNA interference (RNAi), mediated by 21-nucleotide (nt)-length small interfering RNAs (siRNAs), is a powerful tool not only for studying gene function but also for therapeutic applications. RNAi, requiring perfect complementarity between the siRNA guide strand and the target mRNA, was believed to be extremely specific. However, a recent growing body of evidence has suggested that siRNA could down-regulate unintended genes whose transcripts possess complementarity to the 7-nt siRNA seed region. This off-target gene silencing may often provide incongruous results obtained from knockdown experiments, leading to misinterpretation. Thus, an efficient algorithm for designing functional siRNAs with minimal off-target effect based on the mechanistic features is considered of value.

Results: We present siDirect 2.0, an update of our web-based software siDirect, which provides functional and off-target minimized siRNA design for mammalian RNAi. The previous version of our software designed functional siRNAs by considering the relationship between siRNA sequence and RNAi activity, and provided them along with the enumeration of potential off-target gene candidates by using a fast and sensitive homology search algorithm. In the new version, the siRNA design algorithm is extensively updated to eliminate off-target effects by reflecting our recent finding that the capability of siRNA to induce off-target effect is highly correlated to the thermodynamic stability, or the melting temperature (Tm), of the seed-target duplex, which is formed between the nucleotides positioned at 2-8 from the 5' end of the siRNA guide strand and its target mRNA. Selection of siRNAs with lower seed-target duplex stabilities (benchmark Tm < 21.5°C) followed by the elimination of unrelated transcripts with nearly perfect match should minimize the off-target effects.

Conclusion: siDirect 2.0 provides functional, target-specific siRNA design with the updated algorithm which significantly reduces off-target silencing. When the candidate functional siRNAs could form seed-target duplexes with Tm values below 21.5°C, and their 19-nt regions spanning positions 2-20 of both strands have at least two mismatches to any other non-targeted transcripts, siDirect 2.0 can design at least one qualified siRNA for >94% of human mRNA sequences in RefSeq. siDirect 2.0 is available at http://siDirect2.RNAi.jp/.
Background

RNA interference (RNAi) mediated by double-stranded RNA has become a powerful tool not only for studying gene functions, but also for therapeutic applications [1,2]. In mammalian cells, RNAi is induced by small interfering RNA (siRNA), a duplex of 21-nucleotide (nt) RNAs containing 2-nt 3' overhangs. The siRNAs incorporated into cells are transferred to the RNAi effector complex called RNA-induced silencing complex (RISC) [3,4]. RISC assembles on one of the two strands of siRNA duplex, and is activated upon the removal of the passenger strand [5-9]. The activated RISC is a ribonucleoprotein complex minimally consisting of the core protein Argonaute (Ago) and single-stranded siRNA, which acts as the guide to target complementary sequences within mRNAs [10-13]. The 5’ end of the siRNA guide strand is anchored in the binding pocket of the Mid domain of Archaeoglobus fulgidus Ago-like protein [14,15], and the 3’ end is anchored to the PAZ domain of human [16] and Drosophila [17] Ago in the RISC complex. Thus, in the siRNA guide strand, 19 nucleotides positioned at 2-20 from 5’ end may be responsible for target RNA recognition, leading to the silencing of gene expression by cleaving target mRNA [10-13]. Since RNAi is based on sequence recognition by the siRNA, it can give rise to the silencing of other genes with similar sequences. This phenomenon is referred to as an off-target effect, and the growing evidence from large-scale knockdown experiments indicates that the off-target silencing is induced by the base-pairing between the seed region at positions 2-8 from the 5’ end of the RISC-loaded siRNA strand, and its complementary sequences in the 3’ UTR of the unrelated mRNAs [18-23]. Although RNAi is now widely and routinely used as an experimental tool, the remaining fundamental concern is whether the target gene can be specifically silenced. Especially, accurate knowledge of RNAi specificity is critical for therapeutic technologies.

To avoid off-target effects, one approach may be to select the siRNA whose seed sequence is not complementary to any sequences in the 3’ UTR of all non-targeted genes. However, this approach is problematic because random 7-nt sequence is predicted to appear in every 16,384 bp on average. In fact, we analyzed the human 3’ UTR database and it proved impossible to select such siRNAs. That is, human siRNAs with the most infrequent 7-nt seed sequence still have seed-complementarities with 17 3’ UTR sequences. Recently, we have revealed that the capability of siRNAs to induce seed-dependent off-target effect is highly correlated to the thermodynamic stability of the duplex formed between the seed region of siRNA guide strand and its target mRNA [23]: the melting temperature (Tm) of the seed-target duplex showed strong positive correlation with the induction of seed-dependent off-target effects. The results suggested that the Tm of 21.5 °C may serve as the benchmark, which discriminates the almost off-target-free seed sequences from the off-target-positive ones. Thus, selecting the siRNAs with low Tm of the seed-target duplex should minimize seed-dependent off-target silencing.

We have previously released highly effective, target-specific siRNA design software, siDirect [24], in which siRNA sequences were selected using our guidelines established through extensive experiments to clarify the relationship between siRNA sequences and RNAi activities [7]. In order to exclude potential cross-hybridization candidates, siDirect used the rigorous homology search algorithm to select siRNA sequences that have at least three mismatches to any other non-targeted transcripts [25]. In the updated software, siDirect 2.0, the siRNA design algorithm has been extensively updated to select off-target minimized siRNAs by considering the thermodynamic stability of the seed-target duplex. By using the default parameters, at least one functional siRNA could be designed for >94% of the human mRNA sequences in RefSeq release 30.

Implementation

Overall flow of siRNA selection in siDirect 2.0 is illustrated in Figure 1. All possible 23-mer subsequences, corresponding to the complementary sequence of 21-nt guide strand and 2-nt 3’ overhang of the passenger strand within the target sequence, are generated and filtered in three selection steps described below.

Selection of highly functional siRNAs

In the first step, highly functional siRNA sequences were selected using our algorithm [7] (Figure 1, Step 1). We have revealed that efficient RNAi could be induced by the siRNAs that satisfies the following three sequence conditions simultaneously: A/U at the 5’ terminus of the guide strand; G/C at the 5’ terminus of the passenger strand; at least 4 A/U residues in the 5’ terminal 7 bp of the guide strand. In addition, G/C stretch longer than 9 bp should be absent [7]. The experimental validation showed that 98% of the siRNAs predicted to be functional have reduced the target gene expression [26]. The proportion of functional siRNA sequences selected by this algorithm is 14.7% of all human 23-mer sequences generated from RefSeq 30 (Figure 1A, see Step 1).

Reduction of seed-dependent off-target effects

We have found that the off-target effect is highly correlated with the thermodynamic stability or Tm of the seed-target duplex, which is formed between the nucleotides positioned at 2-8 from the 5’ end of the siRNA guide strand and its target sequence [23]. In the second step, to avoid off-target effect, Tm for the seed-target duplex was calculated using the nearest neighbor model and the thermodynamic parameters for the formation of RNA duplex
as described previously [23] (Figure 1, Step 2). The formula for calculating Tm is: $Tm = \frac{1000 \times \Delta H}{(A + \Delta S + R \ln(C_i/4))} - 273.15 + 16.6 \log [\text{Na}^+]$, where $\Delta H$ (kcal/mol) is the sum of the nearest neighbor enthalpy change, $A$ is the helix initiation constant (-10.8), $\Delta S$ is the sum of the nearest neighbor entropy change [27], $R$ is the gas constant (1.987 cal/deg/mol), and $C_i$ is the total molecular concentration of the strand (100 μM). [Na$^+$] was fixed at 100 mM. As shown in our previous report, calculated Tm of 21.5°C may be a benchmark to discriminate almost off-target-free seed sequences from the off-target-positive ones [23], and thus used as the initial standard in this study. Furthermore, it has been revealed that RNAi silencing is occasionally induced by the passenger strands of functional siRNAs [23], and that the passenger strands also take part in the seed-dependent off-target gene silencing [18,28]. Thus, siRNAs whose seed-target Tm is below 21.5°C for both guide and passenger strands were selected in this study. In consequence, 3.0% of all human 23-mer sequences remained available (Figure 1A, see Step 2). Calculated Tm value for each siRNA is shown in the siDirect 2.0 output page (Figure 2A).

**Elimination of near-perfect matched genes**

Several studies have indicated that the effect of single-base mismatches between the siRNA guide strand and the target mRNA varies, according to the positions of the mismatch and/or the sequence of siRNA [21,29]. However, as shown in our previous report, it is obvious that even when the Tm value of the seed-target duplex is sufficiently low, the target gene silencing can still take place if the non-seed region is completely complementary [23]. Therefore, in the third step, siRNAs that have near-perfect matches to any other non-targeted transcripts were eliminated. In siDirect 2.0, off-target searches are performed for 19-mer sequences at positions 2-20 of both strands of the siRNA duplex (Figure 1B, Step 3), because these 19 nucleotides are thought to be involved in target mRNA recognition. Since widely-used BLAST tends to overlook near-perfect match candidates frequently, we used our fast and sensi-
Results and Discussion

We performed a genome-wide design of siRNAs for human mRNAs in RefSeq release 30 with the following parameters: 1) satisfying our functional siRNA design algorithm [7,24], 2) Tm values at the seed-target duplex of both the guide and the passenger strands below 21.5 °C, and 3) no off-target hits with less than two mismatches.

The degree of off-target effects is shown to be correlated with the thermodynamic stability or the calculated Tm value of the seed-target duplex [23]. The initial boundary Tm value was set to 21.5 °C to discriminate the off-target-free sequences from the off-target-positive ones, according to the validity of the siDirect 2.0 output values as described.
to our previous report [23]. Among the entire siRNA sequence population that have at least two mismatches to any other non-targeted transcripts, the siRNA sequences with seed-target Tm below 21.5°C account for 2.1% of about 56 million 23-mer fragments found in human mRNAs (Figure 3A), and one or more siRNA can be designed for 94.7% of all human mRNAs (Figure 3B). However, the strong correlation between the calculated Tm and the off-target gene silencing activity indicates that the seed-dependent off-target effect is definitively reduced when the siRNA with lower Tm of seed-target duplex are selected. The population of siRNAs among all human 23-mer sequences with the Tm in the seed-target duplex of less than 15°C and 10°C is 0.7% and 0.3%, respectively (Figure 2A), and the fraction of human mRNAs which can be targeted by more than one siRNA within such criteria decreases to 85.1% and 72.7%, respectively. (Figure 3B). It is also desirable to select siRNA that contains as many mismatches as possible to any non-targeted mRNAs. In addition to the Tm value of below 21.5°C, siRNA sequences with at least two mismatches to any other non-targeted transcripts are selectable for 94.7% of human mRNAs (Figure 3B). However, if the siRNAs having near-perfect match hits with less than three mismatches, with their Tm of seed sequences below 21.5°C, are selected, one or more siRNA can be designed for only 77.2% of the human mRNAs (Figure 3B). When siRNAs with seed Tm below 15°C and 10°C were selected, siRNAs can be designed for only 47.0% and 18.5%, respectively (Figure 3B). Furthermore, the percentage of human mRNAs drops severely to 0.15% if the near-perfect match hits with less than four mismatches are filtered. Thus, siDirect 2.0 filters siRNAs with less than two mismatches by default to avoid severe reduction in the number of siRNA candidates. We were unable to design functional, off-target minimized siRNAs for 5.3% of the RefSeq mRNAs using the default parameters. Typical examples of these mRNAs are the histone clusters (NM_003523, etc.) and ribosomal proteins (NM_002952, etc.), which are known to form multigene families. When designing siRNAs targeting such genes, users can manually investigate the detailed list of off-target gene candidates (Figure 2B) and select the siRNA that does not have off-target hits to unrelated transcripts. Although most existing web servers for designing siRNA incorporate BLAST [30] to avoid off-target effects [31-38], several sites including WI siRNA Selection Server [34], siDRM [39], DSIR [40] and Dharmacon siDESIGN Center consider seed-dependent off-target effects. Current version of WI siRNA Selection Server and siDRM enumerates the transcripts with full homology to the seed region, and DSIR and Dharmacon siDESIGN Center calculate seed frequencies for each siRNA candidate. Therefore, we analyzed the relationship between the calculated Tm and the distribution of each seed sequence in human 3′UTRs. Cal-

**Figure 3**
The proportion of selectable siRNAs and mRNAs according to Tm values in the seed-target duplexes. (A) The percentage of selectable siRNA candidates for human mRNAs according to the Tm values in the seed-target duplexes. The total number of siRNA (56,375,087) is set to 100%. (B) The percentage of human mRNAs harboring at least one target sequence of an siRNA whose Tm value of the seed-target duplex is below the indicated value. 100% indicates 25,651 mRNAs. Off-target hits with 0-4 mismatches between nucleotides at positions in the 2-20 of both siRNA strands and human mRNAs were represented as separate lines.
culated Tm of the seed-target duplexes of all possible 7-nt seed sequences \(4^7 = 16,384\) ranged from -12°C to 60°C, and of these, 4,488 (27.4%) 7-mers had the Tm below 21.5°C (Figure 4A). The number of 3’ UTRs bearing at least one target site of any 7-nt sequence was broadly distributed from 17 to 10,882 (Figure 4B), excluding the sequence AAAAAAAA, which is found in almost all 3’ UTRs with poly(A) tails. When the siRNAs were classified into eight groups according to their Tm of the seed-target duplex, as shown in Figure 4C, siRNAs whose seed-target duplexes had higher Tm, ranging from 20°C to 60°C, were less frequent and similarly distributed. On the other hand, the seed sequences with lower Tm were frequently found in human 3’ UTRs (Figure 4C).

**Conclusion**

We have extensively updated siDirect 2.0 based on our experimental knowledge, and provided a promising website for reducing siRNA off-target silencing. The website selects: 1) functional siRNAs that satisfy our guideline [7], 2) siRNAs with reduced seed-dependent off-target effects by considering the thermodynamic stability of the seed-target duplex, 3) siRNAs that do not hit any non-targeted genes with near-perfect matches. When the candidate functional siRNAs could form seed-target duplexes with Tm values below 21.5°C, and their 19-nt region spanning positions 2-20 of both strands have at least two mismatches to any other non-targeted transcripts, siDirect 2.0 can design at least one qualified siRNA for >94% of human mRNA sequences in RefSeq. This website should provide a wide scope of applications in RNAi studies.

**Figure 4**

**Calculated Tm values and appearances of 7-nt seed sequences.** (A) Calculated Tm values of the duplex formed by all possible 7-nt sequences. The dotted line indicates that the number of 7-nt sequences with duplex Tm below 21.5°C is 4,488 (27.4%). (B) Appearance of 7-nt seed sequences in human 3’ UTRs. The numbers of 3’ UTR sequences containing at least one given 7-nt sequence are shown. (C) Relationship between the appearance of each 7-nt sequence in the 3’ UTRs containing at least one 7-nt sequence and its calculated Tm. Histograms in the right panel show the appearance of each 7-nt sequence in human 3’ UTR, divided into 10°C Tm intervals. The seed sequence whose duplex has lower Tm (colored blue) is more frequently observed in the 3’ UTRs as compared to those with higher Tm (colored orange).
Availability and requirements
Project name: siDirect
Project home page: http://siDirect2.RNAi.jp/
Operating system(s): Platform independent
Programming language: Perl

Any restrictions to use by non-academics: Contact license@RNAi.jp

Authors’ contributions
YN developed the webserver and performed the computational analyses. JY and SM developed the core framework. KU-T supervised the entire study. YN and KU-T drafted the manuscript. All authors read and approved the final manuscript.

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