Profilling of mismatch discrimination in RNAi enabled rational design of allele-specific siRNAs

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ABSTRACT

Silencing specificity is a critical issue in the therapeutic applications of siRNA, particularly in the treatment of single nucleotide polymorphism (SNP) diseases where discrimination against single nucleotide variation is demanded. However, no generally applicable guidelines are available for the design of such allele-specific siRNAs. In this paper, the issue was approached by using a reporter-based assay. With a panel of 20 siRNAs and 240 variously mismatched target reporters, we first demonstrated that the mismatches were discriminated in a position-dependent order, which was however independent of their sequence contexts using position 4th, 12th and 17th as examples. A general model was further built for mismatch discrimination at all positions using 230 additional reporter constructs specifically designed to contain mismatches distributed evenly along the target regions of different siRNAs. This model was successfully employed to design allele-specific siRNAs targeting disease-causing mutations of PIK3CA gene at two SNP sites. Furthermore, conformational distortion of siRNA-target duplex was observed to correlate with the compromise of gene silencing. In summary, these findings could dramatically simplify the design of allele-specific siRNAs and might also provide guide to increase the specificity of therapeutic siRNAs.

INTRODUCTION

RNAi is an evolutionarily conserved process where small interfering RNA (siRNA) specifically represses the expression of target genes (1,2). siRNAs are widely expected to become next generation of biological therapeutics (3,4), and they are initially anticipated to play a major role in treatment of diseases involving single nucleotide polymorphisms (SNPs) where discrimination against single nucleotide variation between wild-type and mutant alleles is demanded (5–8). This dream was then hammered by subsequent reports demonstrating that siRNAs could incur widespread knockdown of unrelated genes, a phenomenon known as ‘off-target effects’ (9–12). Closer scrutiny of ‘off-target effects’ of siRNA has however revealed the Janus-like view of siRNA in term of target specificity. On one hand, siRNA does induce weak down-regulations on sites that are apparently only related to the siRNA by matching to the seed region (13). On the other hand, siRNA could discriminate some very closely related target sites with only two or even one nucleotide mismatches (14,15). Nobody knows how siRNA or the RISC complex could harmonize the two apparently conflicting properties of siRNAs.

Due to utmost importance of siRNA specificity in siRNA drug development and the need for generating allele-specific siRNAs, we have created an experimental system to generate insight of siRNA discrimination of mismatched target sites. Using more than 400 reporter plasmids for 20 siRNAs we have revealed a general rule for mismatch tolerance and discrimination. Using this discovery as a guideline, we have constructed the first model of mismatch-tagged, position-specific discrimination of closely related target sites. The model was then successfully used for designing allele-specific siRNA.

MATERIALS AND METHODS

Oligonucleotides and plasmids

DNA oligonucleotides were from Invitrogen (Beijing, China). RNA oligonucleotides were from Genepharma

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(Shanghai, China) and Proligo Sigma (Paris, France). Plasmid DNAs were extracted using a mini-purification kit (Promega).

RNAi assay

Human embryonic kidney (HEK293) cells were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 2 mM l-glutamine, 100 U/ml penicillin and 100 μg/ml streptomycin (Life Technologies, Gibco). The cells were seeded into 24-well plates at a density of ~1 × 10^5 cells/well one day before transfection. siQuant vector (0.17 μg/well) carrying the target site of tested siRNA was transfected into HEK293 cells at approximately 50% confluence, together with pRL-TK control vector (0.017 μg/well), with or without the siRNA (13 nM). The activity of both luciferases was determined by a fluorometer (Synergy HT, BioTek, USA) before the firefly luciferase activity was normalized to renilla luciferase for each well. Silencing efficacy of each siRNA was calculated by comparison with a sample without siRNA treatment. All experiments were performed in triplicate and repeated at least twice.

Northern blot assay

Twenty-four hours after transfection with siQuant vector and siRNA, total RNA was harvested from HEK293 cells with Trizol reagent (Invitrogen) according to the manufacturer’s instructions. Total RNA (10–20 μg) was separated by electrophoresis in an ethidium bromide-containing agarose–formaldehyde gel. The intensities of the 18S and 28S rRNA bands were checked under ultraviolet light to verify that all samples were loaded equally and that no RNA degradation had occurred. The DNA probe was labeled with biotin-dUTP using the Prime-a-Gene labeling system (Promega). Hybridization and stringent washing were performed according to the following procedure: (i) 1000 steps of initial energy minimization with solvate and ions, holding RNA structure fixed; (ii) 2500 steps of restrained energy minimization (no restraints for the minimization of the wild-type RNA structure); cis-Watson–Crick base pairing hydrogen bonds were used as constrains for the optimization; (iii) 20 ps molecular dynamics simulation by heating the whole system from 0 to 300 K, while keeping the RNA molecule fixed with weak restraints; and (iv) molecular dynamics equilibration with a time step of 2 fs at 300 K until the RMSD curves became stable.

RESULTS

Position-dependent mismatch discrimination pattern of siRNA

Using a previously established gene silencing assay (16), a siRNA target fragment, perfectly matched or single nucleotide mismatched relative to the siRNA guide strand, was fused in-frame with the firefly luciferase gene in a mammalian expression vector. The resulting fusion reporter was then used as an artificial target to evaluate the silencing efficacy of the siRNA on the target. In the present study, target mutations were initially made at three positions, selected from target regions of differing mutation sensitivity (9,15). In total, twenty effective siRNAs were examined for their silencing efficacy on 200 matched or single nucleotide mismatched targets, covering all such mismatches at these positions (Supplementary Data S1). The mismatch formed between the siRNA guide strand and the mutant target is indicated by X:Y, where X stands for the nucleotide in the guide strand and Y stands for the opposite nucleotide in the target.

Surprisingly, when the silencing efficacies were aligned in terms of the position and the identity of the mismatched nucleotide in the siRNA guide strand, consistent discrimination profiles were found in each subgroup (Figure 1, measurement variances in Supplementary Data S2). For example, in the case of four siRNAs having adenine at position 17 of the guide strand (Figure 1, upper left panel), mutation of the target nucleotide from U to C led to an A:C mismatch between the guide strand and the target. Compared with other mutations, mutation from U to C caused the least loss of silencing efficacy for all four siRNAs, while mutation from U to G...
(forming an A:G mismatch) resulted in the most significant loss of silencing activity. Therefore, for siRNAs having adenine at this position, the tolerance order was C > A > G; for siRNAs with cytosine at this position, the order was A > U > C; and for siRNAs with uracil, the order was U > C > G.

When mutations were made on the target nucleotide opposite to position 12 of the guide strand, the tolerance order was A > C = G for adenine, A > U > C for cytosine, G > C = U for uracil, and A > U > G for guanine in the guide strand. Unlike position 17, which is located in a region of low sensitivity, position 12 is in a region of high sensitivity approximating the cleavage site. Accordingly, relatively low tolerance was found at this position.

Position 4 was selected from the seed region of the siRNA guide strand; this position is implicated in off-target gene silencing mediated by translation repression (9,10). When mutations were made here, the tolerance order was A > C = G for A in the guide strand, A > U > C for C in the guide strand, U > G = C for U in the guide strand, and U > A = G for G in the guide strand.

From the layered gene silencing efficacy at these positions, several revealing observations were made. First, a unique tolerance profile was identified at each tested position, demonstrating significant difference in the gene silencing of the most tolerant and the less tolerant target mutations (Supplementary Data S3). For example at position 12, the A:A mismatch had virtually no effect on the silencing efficacy of all five siRNAs having adenine, however, the same mismatch significantly compromised gene silencing of the four siRNAs with adenine at position 17 (15% reduction on average). Second, dramatically different tolerance levels occurred if mismatched nucleotides were swapped between the guide strand and the target. For example at position 17, both A:C and C:A mismatches were well tolerated and mediated strong gene silencing; equivalent tolerance did not occur at position 12. While an A:C mismatch at position 12 was largely tolerated (11% reduction on average), silencing efficacy of the siRNAs was significantly compromised by a C:A mismatch (25% reduction on average). In addition, tolerance of the U:G wobble base pair was not as great as that of the G:U wobble base pair in general. Third, the order of position-specific tolerance was independent of the sequence context of the siRNA or the target.

To rule out potential bias due to the cell type used in the assay, the experiments were repeated with Hela cells,
revealing similar tolerance profiles. Together, these data demonstrated that the tolerance profiles are a common property of RNAi processing.

**Building a general model of single nucleotide mismatch discrimination profile of siRNA**

Taking into consideration of its potential significance in allele-specific siRNA design, other target positions were further examined. For each given identity of the nucleotide on a target position, at least two siRNAs were examined using the same experimental setting. The resulting gene silencing data were aligned accordingly (Figure 2, Supplementary Data S4). As expected, a unique discrimination profile was found at individual positions across the whole target site, demonstrating the generality of the mismatch discrimination in RNA interference.

To the best of our knowledge, this is the most comprehensive data set regarding single point target mutations to date. The most striking finding was that, depending on the nucleotide in the siRNA guide strand, a most tolerant and a least tolerant mismatch, as well as a few most sensitive...
positions occurred. Given A in the guide strand, the most tolerant mismatch was the A:C (purine:pyrimidine) mismatch that resulted in 69% reduction of reporter gene activity on average, and the least tolerant mismatch was A:G (purine:purine, 52% reduction on average). The most sensitive positions were 9, 10, 12, 14 and 15 (Figure 2A). For G in the guide strand, the most tolerant mismatch was G:U (purine:pyrimidine, 72% reduction) and the least tolerant was G:G (purine:purine, 64% reduction); the most sensitive positions were 8–13 (Figure 2B). For U in the guide strand, the most tolerant mismatch was U:G (purine:pyrimidine, 62% reduction) or U:U (pyrimidine:pyrimidine, 62% reduction) and the least tolerant was U:C (pyrimidine:pyrimidine, 49% reduction); the most sensitive positions were 9, 11–13 and 15–16 (Figure 2C). For C in the guide strand, the most tolerant mismatch was C:A (purine:pyrimidine, 80% reduction) and the least tolerant was C:C (pyrimidine:pyrimidine, 62% reduction); the most sensitive positions were 10–13 and 16 (Figure 2D). In general, purine:pyrimidine mismatches were more tolerated than purine:purine or pyrimidine:pyrimidine mismatches. Other interesting observations included: (i) distinct sensitivity profiles were found depending on the nucleotide in the siRNA guide strand. The mismatches formed by G were generally more tolerated than mismatches formed by A in the guide strand (Figure 2A and B). (ii) In contrast to previous suggestions that perfectly complementary around the cleavage site is required in RNAi (17–20), our data showed that these were not always the most sensitive positions. For example, mismatches at position 10 were much more tolerated than at positions 9 and 11 (Figure 2C). (iii) Generally low sensitivity was found in the seed region of the siRNA guide strand.

To investigate the nature of the silencing in our study, northern blot assay was performed. The results showed that the amount of mutated target transcript was significantly reduced by the introduction of the siRNA, indicating that the gene silencing was mediated by target cleavage rather than translational repression (Figure 3A). Furthermore, mRNA quantification by northern blot perfectly matched the gene silencing efficacy measured by luciferase activity (Figure 3B).

**Figure 3.** Silencing of single point mismatched target is mediated by target cleavage. Northern blot was performed with mRNA transcripts harboring perfectly matched or single nucleotide mismatched target sites of siRNA-4. Reporter plasmids mutated at positions 12 and 10 were co-transfected into cells with (+) or without (−) the siRNA, and then the amount of reporter mRNA was evaluated using fluorophore-labeled probe. (A) siRNA treatment markedly decreased the fusion gene mRNA level relative to the control without siRNA. 18s rRNA was included as loading control. (B) The amount of mRNA transcript was quantified and compared with the gene silencing efficacy. Both the northern and luciferase assay were normalized to perfectly matched target.

Rational design of allele-specific siRNA using the mismatch discrimination model as a guide

The remarkable discrimination profile revealed in our studies indicated that, for a given mismatch, some positions are more sensitive (less tolerant) than others. If a mismatch was placed at its most sensitive position, selective gene silencing between the perfectly matched and the single nucleotide mismatched target could be achieved.
Table 1. Recommended positions for allele-specific siRNA design

| Mutation type | Nucleotide on siRNA | Discriminate (least tolerant) | Positions in guide (strand) |
|---------------|---------------------|-----------------------------|---------------------------|
| A→C          | G                   | A                           | 8, 9, 10, 11, 16          |
| A→G          | C                   | A                           | 10, 11, 12, 16            |
| A→T          | A                   | A                           | 9, 10, 14, 15, 16        |
| C→A          | U                   | C                           | 11, 12, 13, 14, 16       |
| C→G          | C                   | C                           | 9, 10, 12, 13, 16        |
| C→T          | A                   | C                           | 9, 10, 11, 12, 14        |
| G→A          | U                   | G                           | 10, 13, 15, 16           |
| G→C          | G                   | G                           | 10, 11, 12, 13           |
| G→T          | A                   | G                           | 9, 10, 11, 12, 14        |
| T→A          | U                   | U                           | 9, 11, 12, 15            |
| T→C          | G                   | U                           | 10, 11, 12, 13           |
| T→G          | C                   | U                           | 10, 11, 12, 13           |

This discrimination is of great value to the silencing of SNP disease alleles, in which a siRNA that is designed to perfectly match the mutant allele will also pair imperfectly with the wild-type allele, and form a single nucleotide mismatch between them. By placing the mismatch in its most sensitive positions (Table 1), the siRNA might mediate effective silencing of the mutant allele, while off-target silencing of the wild-type allele is minimized.

To establish proof of concept, single nucleotide mutated alleles of the PIK3CA (phosphoinositide-3-kinase, catalytic, alpha polypeptide) gene were further studied. PIK3CA encodes the p110α catalytic subunit that regulates essential signaling pathways important for neoplasia (21,22). This gene is frequently mutated in human tumors in the form of a single point mutation, thus providing us with a model system to evaluate the selective gene silencing strategy (23). The two most frequent SNP sites derived from human tumor samples were selected for study: one causes a G to A transition (mutation 1633), and the other an A to G transition (mutation 3140). Based on our earlier data, four allele-specific siRNAs were designed to target mutation 1633 and three were designed to target mutation 3140, placing the mismatched nucleotide at their most sensitive positions in the siRNA guide strand (Figure 4A). To evaluate the selective gene silencing, fusion reporter genes carrying wild-type or mutant target were made as described above. Cell assays successfully identified three allele-specific siRNAs against mutation 1633 (1633-10, 13, 16) and one against mutation 3140 (3140-11) (Figure 4B and C). While pronounced gene silencing occurred with perfectly matched mutant alleles, silencing of mismatched wild-type alleles was insignificant, displaying high selectivity in allele-specific gene silencing. The discrimination was further observed over siRNA concentrations from 0.02 to 80 nM, indicating that it is not a concentration-dependent phenomenon (Figure 4D and E).

**Duplex conformation change is correlated with mismatch discrimination**

In light of a recent study showing that a mismatch between the siRNA guide strand and the target RNA strand causes marked structural change and distorts its helix conformation (24–26), we quantitatively investigated two crucial features of the siRNA-target duplex, helix conformation and thermodynamic stability. Studies were performed with duplex RNAs formed by the guide strand of siRNA-604 and its matched, or single nucleotide mismatched targets, at position 6, 9, 10, 12, 15 or 17 (Supplementary Data S5). At these positions, the perfectly matched duplexes, the most tolerant duplexes (A:C or C:A), as well as the least tolerant duplexes (A:A or C:C) were examined.

To quantify the conformational distortion caused by the mismatches, an experimental CD spectrum assay and computing-based MD simulations were performed. Consistent with previous studies (25,26), a distorted helix conformation was observed in mismatched duplexes when compared with perfectly matched counterparts (Figure 5B). More importantly, matched patterns were revealed between the silencing efficacies and the deviations in negative Cotton values of the CD spectra exhibiting conformational distortion of the helix structure (Figure 5A and B). This finding was further corroborated by MD simulations (Figure 6), suggesting a correlation between the compromised gene silencing and the degree of conformational distortion. Taken together, our data indicated that greater conformational distortion caused more interference with gene silencing activity, thus providing an explanation of why some target mutations are more tolerated than the others.

Besides conformational change, the influence on the thermodynamic stability of the RNA duplexes was also examined in terms of melting temperature. It was interesting to note that changes in melting temperature correlated with the positions but not the identities of the mismatches (Figure 5A and C). For example, at position 15, A:A and A:C mismatches had little effect on melting temperature compared with the wild-type counterpart. The silencing efficacy of the A:A mismatch was, however, decreased by a significant 72%, while the silencing of the A:C mismatch was decreased by only 9%. The apparent inconsistency between the changes of melting temperature and gene silencing efficacies indicated that the perturbed thermodynamic stability of the duplex does not account for the compromised gene silencing.

**DISCUSSION**

In allele-specific gene therapy, specifically designed siRNAs are expected to selectively reduce the expression of disease-causing alleles without affecting the expression of their wild-type counterparts. To achieve this goal, we performed a large-scale RNAi assay to understand the role of target mutation in off-target gene silencing. The results demonstrated a position-specific discrimination profile across the whole target site, and in particular, identified the most tolerant and the most disruptive single point target mutation at each target position. Furthermore, for any given mismatch between siRNA guide strand and target RNA, the study identified the most sensitive positions for a single point target
mutation, therefore establishing a solid basis for allele-specific siRNA design.

A consistent tolerance preference was observed for the same mismatch when it was at the same position in the target site, indicating that processing of similarly mismatched substrate duplexes at the same location within the Ago protein likely results in comparable interference with its functionality. In contrast, when the same mismatch was at different target positions, distinct tolerance preferences were found. This indicated that processing of similarly mismatched duplexes at different locations within the protein resulted in distinct levels of interference.

**Figure 4.** Potent and selective silencing of SNP disease alleles. (A) Four siRNAs were designed to target the disease-causing allele at position 1633 of the PIK3CA gene, and three siRNAs were designed to target the disease allele at position 3140. (B, C) RNAi assays performed with fusion luciferase reporter having wild-type or target mutation at position 1633 and 3140. (D, E) RNAi assays performed with siRNA 1633-10, 16 over a series of siRNA concentrations.
with its functionality. In search of the mechanism involved in the discriminative gene silencing against differently mutated targets, quantitative assays were performed on siRNA-target duplexes, with the focus on the influence on duplex conformation and thermodynamic stability. In agreement with previous studies (25,26), our data showed that mismatch between siRNA guide strand and target RNA caused a marked change in the normal helix structure. Furthermore, our data indicated that the tolerant mismatches caused less structural distortion than the disruptive mismatches, which explains why some mutations were more tolerated than others. Collectively, these lines of evidence led us to speculate that, depending on its position and identity, a mismatch formed between the siRNA guide strand and the target RNA might introduce a conformational change in the

Figure 5. Correlation between gene silencing efficacy and CD spectra or thermodynamic stability. Mismatch-derived alteration of conformational distortion or duplex thermodynamic stability was examined using RNA duplexes formed by the guide strand of siRNA-604 and matched or single nucleotide mismatched targets. Conformational distortion of the RNA duplex was evaluated by measuring CD spectra (B), and the thermodynamic stability was evaluated by melting temperature (C). For comparison, the gene silencing efficacy of the respective target is shown in (A). Mismatched position in siRNA guide strand and base-pairing type are indicated on the horizontal axis.
siRNA-target duplex, which in turn interferes with the functionality of the Ago protein in off-target gene silencing. The influence on Ago function is determined by the level of the conformational change caused by the target mutation.

As a major potential application of therapeutic siRNAs, allele-specific gene therapy has received increasing attention in recent years. Among the 2.6 million SNP polymorphisms characterized in the human genome, the proportions of the substitutions are A/G, 32.77%; C/T, 32.81%; A/C, 8.98%; G/T, 9.06%; A/T, 7.46%; and C/G, 8.92. Around 65% of SNPs occur between the two purines (A or G) or the two pyrimidines (C or T), which were thus named transition substitutions (27). Targeting these abundant SNP alleles raised a serious challenge to siRNA design. For example, to silence a disease-causing mutation from wild-type A to G, siRNA designed against the mutant G allele has a C in the guide strand. When this siRNA interacts with the wild-type A allele, a C:A mismatch is formed between them. As revealed in our study, the C:A mismatch is well tolerated in the processing of RNAi and causes minimal interference with gene silencing efficacy. To confront this challenge, a G→A and an A→G transition mutation were chosen to test in our study. By placing the mismatches in their most sensitive positions, as identified in the study, the specifically designed siRNA showed great discrimination against these two alleles, indicating that the principle proposed here is generally applicable to allele-specific gene silencing.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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