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An efficient RNA interference screening strategy for gene functional analysis

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Abstract

Background: RNA interference (RNAi) is commonly applied in genome-scale gene functional screens. However, a one-on-one RNAi analysis that targets each gene is cost-ineffective and laborious. Previous studies have indicated that siRNAs can also affect RNAs that are near-perfectly complementary, and this phenomenon has been termed an off-target effect. This phenomenon implies that it is possible to silence several genes simultaneously with a carefully designed siRNA.

Results: We propose a strategy that is combined with a heuristic algorithm to design suitable siRNAs that can target multiple genes and a group testing method that would reduce the number of required RNAi experiments in a large-scale RNAi analysis. To verify the efficacy of our strategy, we used the *Orchid* expressed sequence tag data as a case study to screen the putative transcription factors that are involved in plant disease responses. According to our computation, 94 qualified siRNAs were sufficient to examine all of the predicated 229 transcription factors. In addition, among the 94 computer-designed siRNAs, an siRNA that targets both TF15 (a previously identified transcription factor that is involved in the plant disease-response pathway) and TF21 was introduced into orchids. The experimental results showed that TF15 is involved in plant defense responses. Interestingly, our second-round analysis, which used an siRNA specific to TF21, indicated that TF21 is a previously unidentified transcription factor that is related to plant defense responses.

Conclusions: Our computational results showed that it is possible to screen all genes with fewer experiments than would be required for the traditional one-on-one RNAi screening. We also verified that our strategy is capable of identifying genes that are involved in a specific phenotype.

Keywords: RNA interference, RNAi screening, SiRNA design, Gene functional analysis, Group testing

Background

The study of plant genomes has increased dramatically over the last decade [1,2]. Many genome sequencing projects for important crops, such as maize [3,4], wheat [5] and rice [6,7], have recently been completed. The available plant genomic sequence data will continue to increase explosively because of the recently developed next-generation sequencing techniques [8-10]. Therefore, a major challenge to all biologists is how to efficiently translate the functions of the genetic code sequences. It is especially important to rapidly screen for genes that are related to certain important phenotypes, such as salt-tolerance, disease-resistance and flower color, because these genes will help to improve crops. For example, by transferring *mannitol-1-phosphate de-hydrogenase gene* from *Escherichia coli*, researchers have created many genetically modified plants with increased salt tolerance, which is beneficial to farmers and consumers [11,12].

One of the most efficient tools in high-throughput gene functional screening is the application of RNA interference (RNAi) [13]. RNAi can be induced by introducing a small double-stranded (ds) RNA into a cell [14] and has been widely employed in genome-wide scale studies of



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model organisms, including animals and plants [15-20]. The primary goals of these applications are usually related to the screening of a single gene or multiple genes that convey(s) a specific phenotype. In these applications, each dsRNA is designed to target one specific gene and to facilitate the observation of the resulting phenotype change. This one-on-one approach works very well for model organisms, because whole genome RNAi libraries are available for these model organisms [21-23]. However, even when one-on-one RNAi experiments are feasible in model organisms, it is still laborious and costly because more than ten thousand coding and non-coding genes are typically found in an organism (e.g., approximately 20,000-25,000 genes were identified in the human genome [24] and approximately 25,500 genes were predicted in the Arabidopsis genome [25]). Thus, matches to the gene number in RNAi experiments should at least be performed. Although using a vector carrying multiple shRNAs/multi-miRNA hairpins can be applied to silence multiple genes simultaneously ("combinatorial RNAi") [26,27] and can reduce the required experiments for highthroughput analysis, these approaches still need to design and synthesize as many siRNAs as the number of target gene sets.

RNAi libraries are not available for non-model organisms, and the cost and scale of high-throughput RNAi screening experiments are still difficult for most laboratories. In reality, RNAi screening can focus on screening specific groups of genes through strategies such as literature surveys or transcriptome analyses [28,29]. However, these approaches usually identify hundreds to thousands of candidate genes; the subsequent large-scale RNAi analysis would be still a challenge for most laboratories by using a one-on-one approach. In fact, siRNAs do not need to perfectly complement their targets. Genes that contain a sequence that is partially similar to the designed siRNA could also be down-regulated (off-target effect) [30]. The off-target effects have been widely discussed and studied in both animals and plants [13,31,32], and they are widely considered to be a general problem for analyses; however, off-target effects also facilitate the silencing of multiple genes simultaneously with a carefully designed siRNA. Thus, utilizing the off-target effect to carefully design siRNA could provide opportunities for us to design a minimum of siRNA(s) to target a maximum of genes, which would reduce the labors and costs for highthroughput gene functional screening and would decrease the obstacles for most laboratories in facing highthroughput screening. This approach would be especially useful for applications on non-model organisms for which RNAi libraries are not available.

Here, we utilize the off-target effects to develop a hierarchical RNAi screening strategy to reduce the cost and labor of high-throughput RNAi analyses. Our strategy utilizes a heuristic algorithm to design siRNAs that have multiple target genes and uses group testing methods [33] to minimize the RNAi analysis for identifying genes that are involved in a specific phenotype. Our strategy can be applied to whole-genome RNAi analyses to reduce the number of initial experiments that are needed for phenotype screening, and it is especially suitable for a group of candidate genes that were first identified by an experimental approach. Moreover, to our knowledge, no design of an siRNA for multiple-gene targets has been described previously. We have verified the applicability of this method by analyzing transcription factors that are known to be involved in plant defense responses in orchids, and we also identified a new transcription factor that is involved in this process.

Methods

Orchid ESTs library

To verify the applicability of our approach, we used an established Orchid (Phalaenopsis equestris) expressed sequence tags (ESTs) library in our initial test. There are 233 previously predicted transcription factor (TF) sequences [34] and 8017 mRNA sequences that are available for download from the NCBI ESTs database. We used the BLAST tool [35] to remove the redundant sequences. First, we used each transcription factor sequence to perform a BLAST search against the transcription factor data to delete the redundant sequences. If the BLAST result revealed that two sequences had a >95% match ratio and a >85% coverage ratio, we removed the shorter sequence as a redundant sequence. The match ratio is defined as the ratio of the number of identical bases in the aligned sequences to the total length of the aligned sequences, and the coverage ratio is defined as the total length of the aligned sequences relative to the length of the longer sequence among the two transcription factor sequences. After the pre-processing, there were 229 non-redundant transcription factor (TF) sequences remaining. The sequences that we used are available at OrchidBase (http://140.116.25.218/EST/) [36].

Because the mRNA data could overlap with TF data, we continued to remove the mRNA sequences that overlapped with the TF data. We performed a BLAST search against the mRNA data for each TF sequence. If the BLAST result revealed that one mRNA sequence had an >85% coverage ratio or 90% overlapping ratio with the TF sequence, then we removed the mRNA sequence as an overlapping sequence. The overlapping ratio was defined as the total length of the aligned sequences relative to the length of the shorter sequence among these two sequences.

Qualified siRNA model for our strategy

Our strategy was designed to screen all of the genes in the genome or to screen candidate genes that had been

narrowed down using previous data or experimental analyses. Therefore, without loss of generality, we defined two disjoint sets of genes. We first defined the set of candidate-genes, $C = \{c_1, c_2, \ldots, c_n\}$, which contained the candidate genes of the RNAi analysis. Second, we defined the set of excluded-genes, $E = \{e_1, e_2, \dots, e_m\}$, which contained all of the genes except the candidategenes. Because silencing these excluded-genes would have likely complicated further analysis, our RNAi design attempted to target the candidate-genes and avoid silencing the excluded-genes. Furthermore, to utilize the off-target effect to design the proper siRNAs for the analysis, we employed two user-defined parameters, d_T and d_{N_2} to determine the specificity of the designed siRNAs. First, the user can employ d_T to define the maximum number of sequence mismatches that will be tolerated between the designed siRNA and its target. Second, the user can adjust d_N to define the minimum number of sequence mismatches that will be allowed between the designed siRNA and its non-target to prevent the designed siRNA from targeting unanticipated genes. Before we can provide a formal definition for a qualified siRNA and its target gene(s), we must first introduce the definition of a qualified sequence as follows:

Given a set of *n* candidate-genes, $C = \{c_1, c_2, ..., c_n\}$, a set of *m* excluded-genes, $E = \{e_1, e_2, ..., e_m\}$, and three integers, d_T , d_N and L ($L > d_N > d_T$), a sequence *r* of length *L* is determined to be a qualified sequence if and only if there exists a subset of candidate-genes *T*, $T \subseteq C$, such that for each gene $t_i \in T$, $HD(r, u_i) \leq d_T$ for some length *L* substring u_i of t_i and for each gene $g_i \subseteq \{E \cup C - T\}$, $HD(r, u_i) \geq d_N$ for any length *L* substring u'_i



Figure 1 Schematic diagram of each step of our heuristic method. (a) The process of enumerating the subsequences from the candidategenes. The green sequence represents a candidate-gene, and the blue sequences represent the subsequences that were derived from this candidate-gene by the sliding scan. (b) Diagram of the relationship between the subsequences. The blue lines $C_1, C_2, ..., C_8$ represent the candidate-genes, and $S_1, S_2, ..., S_{11}$ are the subsequences that were enumerated from these candidate-genes. A solid line between two subsequences indicates that the two subsequences are **neighbors**, and the dotted line indicates that the two subsequences are **far_neighbors**. Each subsequence is marked as a candidate for a qualified sequence. (c) Diagram of the relationship between the marked subsequences after the **far_neighbor** examination. The subsequences S_5, S_9 and S_{10} are unmarked because they all contained a **far_neighbor**. However, S_2 is still marked because one of its **neighbors** was located in C_4 , and its **far_neighbor** S_5 was also located in C_4 . In this situation, we are not concerned about whether the siRNA that was designed based on S_2 will recognize S_5 because C_4 is already the target gene of this siRNA. Therefore, S_2 is still marked as a candidate for the qualified sequence. (d) Diagram of the relationship between the marked subsequences after the powerful subsequence examination. S_1, S_4 and S_6 are unmarked because they are not powerful subsequences and are all dominated by S_2 . (e) Diagram of the relationship between the marked subsequences after the excluded-gene hit examination. E_i is one of the excluded-gene is less than d_{N_i} this scenario also indicates that the Hamming distance between a marked subsequence and a substring that is located in an excluded-gene is less than d_{N_i} this scenario also indicates that this marked subsequence contains an excluded-gene hit. Because any subsequences that contain an excluded-gene hit will

of g_i . Here HD(,) represents the Hamming distance between two strings.

We defined a qualified siRNA r' as the reverse complement sequence of a qualified sequence. Therefore, by definition, finding a qualified siRNA is equal to finding a qualified sequence. Moreover, because there are no more than d_T mismatches between each gene $t_i \in T$ and the qualified siRNA r', T would be the target gene set of the qualified siRNA r'. Conversely, $\{E \cup C-T\}$ is the non-target gene set of r' because each gene $g_i \subseteq \{E \cup C-T\}$ has at least d_N mismatches with the qualified siRNA r'. Each qualified siRNA can silence some of the candidate-genes and avoids silencing the excluded-genes; therefore, they can be used to perform an RNAi experiment that will identify genes that are involved in a specific phenotype.

Enumerating subsequences and computing subsequence similarity

To obtain sequences for the qualified siRNAs, we first enumerated all of the subsequences of length L from the candidate-genes with a sliding scan (Figure 1a). The subsequences that contained an undetermined nucleotide "N" were discarded due to the bad sequencing quality of the sequence. Each subsequence from where the sequence was derived and its original position was recorded. Therefore, every subsequence could be differentiated by its original sequences and its position. We temporarily marked these subsequences as candidates for qualified sequences. Next, we computed the Hamming distances of the pairwise subsequences to measure their sequence similarity. If the Hamming distance between two subsequences was equal to or smaller than d_T these two subsequences were considered *neighbors*. However, if the Hamming distance between the two subsequences was larger than d_T and smaller than d_N , these two subsequences were considered *far_neighbors*.

According to the sequence similarity, we obtained the neighborhood relationship of all of the similar subsequences (Figure 1b). Let N(r) denotes the set of all **neighbors** of the subsequence r. According to the off-target effect, because r is very similar to its **neighbors**, when the reverse complement sequence of r is used as an siRNA in an RNAi experiment, it is possible that this siRNA will also target the **neighbors** of r. Therefore, the neighborhood relationship of these subsequences allowed us to determine the potential target genes of the siRNAs that were designed based on these subsequences.

Far_neighbor and powerful subsequence examination

To prevent the targeting of unanticipated candidategenes, each qualified sequence was required to contain a Hamming distance of at least d_N with any substrings of each gene with the exception of its own target genes. However, because d_T and d_N are defined by users, a subsequence may have *far_neighbors*. If we designed an siRNA that was derived from a subsequence that contained *far_neighbors*, this designed siRNA would not be guaranteed to target these *far_neighbors*. Therefore, in this situation, our algorithm avoids selecting subsequences that contain *far_neighbors* to design the siRNA, and it would unmark the subsequence with *far_neighbors* unless the *far_neighbors* of a subsequence were derived from the genes from which its *neighbors* were derived (Figure 1c).

To reduce the computing time that was required to perform the next steps, we attempted to remove the less useful subsequences. To achieve this, we first looked for a "powerful subsequence". Any subsequence P that satisfied equation (1), where \cup is the union operator, was defined as a powerful subsequence, and the powerful subsequence dominated its *neighbors*. Because the powerful subsequence is expected to target all of its *neighbors*' target genes, the removal of these subsequences (their target genes can be covered by the powerful subsequence) will save computational resources without causing any side effects. (Additional file 1, Figure S1 shows an example of a powerful subsequence and the subsequences that would be unmarked.)

$$\cup_{\nu \in N(P)} N(\nu) \subseteq \{N(P) \cup P\}$$
(1)

To identify the powerful subsequences, we examined each subsequence to determine whether it satisfied equation (1). After identifying the powerful subsequences, we continued to unmark the subsequences that were dominated and were not powerful subsequences (Figure 1d). These unmarked subsequences were ignored in our later analysis.

Excluded-gene hit examination

To prevent the targeting of excluded-genes, we continued to examine whether each marked subsequence displayed a Hamming distance at least a d_N to every excluded-gene. For any subsequence *t*, if there existed a subsequence *u* of an excluded-gene such that $HD(t, u) < d_N$, HD(t, u) represented the Hamming distance between t and u, then t is determined to contain an excluded-gene hit and this subsequence t would be abandoned. To perform this work, we enumerated all of the substrings of length L from the list of excluded-genes by utilizing a sliding scan. Next, we computed the Hamming distance between these substrings and the marked subsequences. We unmarked the subsequences that contained an excluded-gene hit (Figure 1e). The remaining marked subsequences were therefore determined to be qualified sequences, and the reverse complement sequences of these gualified sequences were qualified siRNAs.

Redundancy reduction

All of the qualified siRNAs are expected to silence the candidate-gene(s) without silencing the excluded-gene; however, several siRNAs could target the same genes. Because these qualified siRNAs are redundant to each other, only the effective siRNA will be selected. To properly evaluate the effective siRNA, we use existing siRNA designing rules to evaluate the effective siRNA. In plants, design rules for only artificial microRNAs, amiRNAs, (microRNAs are another class of small regulatory RNA [37]) have been proposed [38]. The proposed amiRNA targeting rules have been shown to share several features with the proposed siRNA designing rules in animal systems [39,40]. We used the siRNA designing rules proposed by Reynolds et al. [40] for selecting siRNA in our analysis. Three siRNA designing rules proposed by Reynolds [40], Hsieh [41] and Takasaki [42] are currently included in our program, allowing users to select their preferred design rules.

If more than one qualified siRNA passed the evaluation, then we continue to calculate their Hamming distance sum. The Hamming distance sum, HD-sum for short, is the sum of all of the Hamming distances between a qualified sequence and all of its neighbors. A smaller HD-sum indicates that a qualified sequence is more similar to its *neighbors*, and the qualified siRNA that is derived from it may have a higher gene silencing efficiency. Therefore, when two qualified sequences had the same target genes, we retained the qualified seguence that displayed the smaller HD-sum. If two gualified sequences both displayed the same HD-sum, we discarded the subsequences that displayed the maximum Hamming distance to one of its neighbors. After we eliminated the redundancies, we obtained the nonredundant qualified sequences.

Construction of a pB7GWIWG2-derived clone

The primer pairs PhaTF15-hpRNA F/PhaTF15-hpRNA R, PhaTF21-hpRNA F/PhaTF21-hpRNA R, PhaTF60 hpRNA F/PhaTF60 hpRNA R and PhaRNA hpRNA F/PhaRNA hpRNA R were used to anneal to doublestranded (dsDNA) (Table 1). Two microliters of each primer (15 μ g/ μ l) was mixed with its complementary primer at 72°C for 10 min and the solution was transferred to 25°C for an additional 10 min. The dsDNA was then cloned into the pCR8[®]/GW/TOPO[®] Gateway entry vector (Invitrogen) by following the manufacturer's recommendations. The pCR8[®]/GW/TOPO dsDNA construct was sequenced to confirm the sequence of the cloned fragments. The LR Clonase II enzyme (Invitrogen) was used to transfer the cloned fragments into pB7GWIWG2 [43] to generate pB7GWIWG2-PhaTF15hpRNA.

Table 1 Primers used in this study

Primer Nucleotide sequence		
TF106 F	5'-CCAAAACGGCCGAGGACCCC-3'	
TF106 R	5'-CGCCACACCTGACGGTCC-3'	
TF157 F	5'- TCATGAACTCGTCGCCTTCCGA-3'	
TF157 R	5'-GCTCGCCCTTCATACGTGGCA-3'	
TF187 F	5'-TGCAAGTTCCAGCAATGCTCCT-3'	
TF187 R	5'-AGAACAACAGGGATGGTGTGCATC-3'	
PhaTF15-hpRNA F	5'-CCTGCGATTTCGACTATAAGT-3'	
PhaTF15-hpRNA R	5'-ACTTATAGTCGAAATCGCAGG-3'	
PhaTF21-hpRNA F	5'-TTACGGAGACCATCTTAAAAC-3'	
PhaTF21-hpRNA R	5'-GTTTTAAGATGGTCTCCGTAA-3'	
PhaTF60 hpRNA F	5'-CAACTTATGCCACTATGCAAT-3'	
PhaTF60 hpRNA R	5'-ATTGCATAGTGGCATAAGTTG-3'	
PhaRNA hpRNA F	5'-TGGTTCCGGTGGTGGGGGCTA-3'	
PhaRNA hpRNA R	5'-TAGCCCCTCCTCCGGAACCA-3'	
CymMV CPF	5'-GAAATAATCATGGGAGAGCC-3'	
CymMV CPR	5'-AGTTTGGCGTTATT CAGTAGG-3'	
PR1 F	5'-AGGACCCTGGCGTCTAAAG-3'	
PR1 R	5'-TATTACAAATCAAACCGCTAAAG-3'	
NPR1 F	5'-CGCATTTGTGTCGTAGTTTAT-3'	
NPR1 R	5'-GGCCTTGCTCCTTTAGTTA-3'	
TF15 F	5'-TGAATTAGCACAAGATCCGAT-3'	
TF15 R	5'-AGCAATGGAAACAGACCACCAAT-3'	
TF21 F	5'-AGGGACGGAGATGCCAAGAAGGC-3'	
TF21 R	5' -CAACCATCCAGCTAATCTATC-3'	
TF60 F	5'-ATGATTTCAAAATTTTATCAGTT-3'	
TF60 R	5'-CAATGGAATACTATTCTGACCA-3'	
Pha RNA F	5'-AAGCCGTCAAGTACAAGGGC-3'	
Pha RNA R	5'-GCTAATCAGTTGCAAGAATTA-3'	
Ubiqutin10 F	5'-CCGGATCAGCAAAGGTTGA-3'	
Ubiqutin10 R	5'-TCAGGCGGAGGACAAGATG-3'	

Transient gene silencing assay

pCambia-CymMV-Gateway, pB7GWIWG2 and their derivatives were transformed into *Agrobacterium tume-faciens* LBA4404 by electroporation. The *A. tumefaciens* strains were cultured in 2 ml of YEB of medium containing 100 mg/l kanamycin and 100 μ M acetosyringone overnight at 28°C. One milliliter of the bacterial culture was then transferred into 10 ml of YEB medium containing 100 mg/l kanamycin and 100 μ M acetosyringone and was further incubated at 28°C until the OD reached approximately 1.0–1.2. The *A. tumefaciens* cultures were then centrifuged at 3000 ×*g* for 10 min; the cells were resuspended in 1 ml of infiltration medium (10 mM MES, 10 mM MgCl₂, 100 mM acetosyringone) and incubated at room temperature for 3 h. Each *Phalaenopsis* plant was then infiltrated with 100 μ l of the

A. tumefaciens suspension. The plant RNA was extracted and analyzed 4 days after the agroinfiltration.

RNA isolation and semiquantitative RT-PCR analysis

To conduct the semiguantitative RT-PCR experiment, the total RNA was extracted from the orchid plant as previously described [44]. One microgram of RNA was treated with RNase-free DNase (Ambion, 1 unit) for 0.5 h to eliminate any genomic DNA contamination. Next, 0.5 µg of DNA-free RNA for each sample was used for the synthesis of the first strand of cDNA using Moloney murine leukemia virus (MMLV) reverse transcriptase following the manufacturer's instructions (Invitrogen). The PCR amplification was performed using gene-specific oligonucleotide primers according to a previously described method [45]. The PCR products were separated by electrophoresis and visualized with ethidium bromide staining. Ubiquitin 10 was used as an internal control. The primer pairs used in the semiguantitative RT-PCR analysis are listed in Table 1.

Results

A case study using the orchid ESTs library

To verify the applicability of our strategy, we used an established *Orchid* (*Phalaenopsis equestris*) expressed sequence tags (ESTs) library in our initial test. The ESTs library contained 229 predicted transcription factor sequences and 7721 mRNA sequences (see the subsection "*Orchid* EST library" in the Methods section).

To use our strategy to screen for transcription factors that are involved in plant defense responses, we defined the predicated transcription factors as candidate-genes and the remaining mRNAs as excluded-genes. A previously described transcription factor, TF15, was used as a positive control [46]. Before we applied our strategy to screen for transcription factors, we first needed to design qualified siRNAs to silence the candidate-genes (see the subsection "Qualified siRNAs for our strategy" in the Methods section). Obtaining qualified siRNAs that target the maximum number of genes was desirable to minimize the number of experiments that were performed for the RNAi screening. However, this problem is NP-hard [we used the distinguishing substring selection problem (DSSP) [47,48] to prove that finding a qualified siRNA that has the maximum number of target genes is NP-complete, and the proof is presented in Additional file 2]. Moreover, to analyze every candidategene by RNAi screening, we needed to obtain sufficient numbers of qualified siRNAs to silence all of the candidate-genes. Therefore, we needed at least k qualified siRNAs such that $\bigcup_{1 \le i \le k} Ti = C$, where T_i is the target gene set of siRNA r'_i , and C is the set of candidategenes. To obtain a sufficient number of qualified siRNAs and their corresponding target genes within a reasonable time, we provided a heuristic algorithm. This heuristic algorithm was incorporated into the workflow that was used to design the siRNAs that were needed in the firstround analysis of our strategy. The flowchart is depicted in Figure 2.

Because previous experiments indicated that 21 nucleotides were sufficient to trigger RNAi in plants, we set the length (*L*) of the qualified siRNA to 21 [49]. Previous reports also indicated that siRNAs can target genes with four mismatches [13,50]. Therefore, we first tolerated the presence of four mismatches (d_T) between an siRNA and its target gene. To avoid the target of unanticipated genes by the siRNA, each siRNA was designed to contain at least five mismatches (d_N), with every gene excluding its own target genes. Therefore, in this case study, we selected the parameters $d_T = 4$, $d_N = 5$ and L = 21 for our heuristic algorithm to design the qualified siRNAs.

In the first stage of our heuristic algorithm, we enumerated 191,087 subsequences of length 21 from these 229 TF sequences and then computed the pairwise Hamming distances between these subsequences (see the subsection "Enumerating subsequences and computing subsequence similarity" in the Methods section). According to the similarity relationship of these subsequences, we obtained the information regarding the potential target genes of the siRNAs that were designed based on these subsequences. Next, to ensure that the designed siRNAs contained at least a d_N distance to their non-target genes and to remove the less powerful subsequences, we performed a *far_neighbor* and powerful subsequence examination of these subsequences, and we unmarked the disqualified subsequences (see the subsection "Far_neighbor and dominating subsequence examination" in the Methods section). After this examination, 188,889 subsequences were still marked. To prevent the designed siRNAs from targeting the excluded-genes, we continued to perform the excluded-gene hit examination of the remaining marked subsequences in the third stage (see the subsection "Excluded-gene hit examination" in the Methods section). In total, 106,188 subsequences passed this examination, and these subsequences were the qualified sequences.

In stage 4, the subsequences were selected from the qualified sequences, and they were used as templates to design the qualified siRNAs. However, some of the qualified siRNAs that were derived from these qualified sequences may have targeted the same genes, which would have resulted in redundancy; therefore, we incorporated some pre-proposed siRNA design rules [40-42] in the redundancy reduction stage to select the best sequences for this use (see the subsection "**Redundancy reduction**" in the Methods section). After the reduction



was performed, 2,147 non-redundant qualified siRNAs remained. To screen every candidate-gene at least once, we needed at least k qualified siRNAs, $r_1 \ldots r_k$, such that $\bigcup_{1 \le i \le k} Ti = S$, where T_i is the target gene set of siRNA r_i . Moreover, to minimize the cost, we also wanted to minimize the number of first-round RNAi experiments. The problem of satisfying $\bigcup_{1 \le i \le k} Ti = S$ with a minimum of k qualified siRNAs is exactly the set cover problem

that has been proven to be an NP-hard problem [51]. Therefore, we adopted the greedy algorithm for the set cover problem to select the siRNAs [52,53]. Furthermore, to increase the examination frequency of each candidate-gene, we slightly modified this greedy algorithm (see Additional file 3 for more detail). A total of 94 siRNAs were selected from the 2,147 non-redundant qualified siRNAs by our modified greedy algorithm (the

list of the selected siRNAs is reported in Additional file 4: Table S1). According to our computational result, these 94 selected siRNAs are sufficient to examine all 229 candidate-genes in the first-round of the RNAi experiments (Table 2). Any researchers who need our program can email to us or visit the website http://algorithm.cs. nthu.edu.tw/RNAi_screening.php.

Hierarchical RNAi analysis strategy

Once we had selected the 94 siRNAs (See Additional file 4: Table S1 for more detail), we were able to perform our strategy, hierarchical RNAi analysis, which contained two rounds of RNAi analysis (Figure 3 shows the schematic diagram of the hierarchical examination process). In the first round of RNAi analysis, we observed phenotypes and correlated those phenotypes to at least one of the genes that were affected in the experiment. Thus, we can rapidly exclude the candidate-genes that did not affect the target phenotype from the suspect list, and we can also determine which candidate-genes could be related to the target phenotype. To identify the associated gene more precisely, a second round of analysis was needed. However, in an RNAi experiment, the selected siRNA and its reverse complement sequence (the selected siRNA sense strand) are employed to form a dsRNA that triggers RNAi. Previous studies have demonstrated that both of the dsRNA strands can be loaded into the RNA-induced silencing complex (RISC) and that both strands can facilitate RNAi [54-56]. Therefore, in addition to the target genes of the selected siRNA, we also needed to examine the possible target(s) of the complementary strand of the selected siRNA. The potential target genes contained a subsequence that had at most d_T mismatches (in this case, $d_T = 4$) with the selected siRNA sense strand. To precisely identify the associated gene from these affected genes, we designed dsRNAs that would specifically target all of the possible target genes in the second-round analysis (Figure 3). According to our statistical results from each sense strand of the qualified siRNA, the average number of potential targets of the sense strand was 0.6511; thus, only a limited number of genes could have been affected by the siRNA sense strands in each RNAi experiment in the first-round analysis.

Experimental verification

To verify if siRNA can silence multiple genes with four mismatches, we first randomly selected an siRNA, siRNA-G41. The predicted targets of siRNA-G41 are TF106, TF157, and TF187. The number of mismatches between siRNA-G41 and its targets, TF106, TF157, and TF187, are 4, 0 and 2, respectively. The experimental results showed that siRNA-G41 successfully silenced *PhaTF106, PhaTF157* and *PhaTF187* (Figure 4), demonstrating that it is feasible to simultaneously silence multiple genes with a carefully designed siRNA and that siRNA can silence the target gene even with 4 mismatches to its target genes.

Among the 94 siRNAs, we observed one siRNA, siRNA-G55, which targeted two transcription factors, TF15 and TF21. TF15 has been previously reported to be involved in the critical salicylic acid (SA)-related plant defense response [46]; the depletion of TF15 in SA-treated plants resulted in the decreased expression of orchid pathogenesis-related gene 1 (*PhaPR1*), which is a marker of the plant defense response, and the important central regulator *PhaNPR1*. To confirm that our strategy can help us to determine that TF15 is involved in plant defense responses, we selected siRNA-G55 to perform a first-round RNAi experiment that utilized *Agrobacterium*-

Table 2 The detailed results of our testing on the Phalaenopsis E	ESTs data with different parameters
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5			
Parameters	$d_T = 3, d_N = 5, L = 21$	$d_T = 4, d_N = 5, L = 21$	$d_T = 4, d_N = 6, L = 21$
Number of enumerated subsequences	191,087	191,087	191,087
Number of subsequences (after <i>Far_neighbor</i> and powerful subsequence examination)	173,106	188,889	117,203
Number of qualified siRNAs	100,890	106,188	6,443
Number of non-redundant qualified siRNAs	487	2,147	352
Average number of target genes of a qualified siRNA	1.0583	1.1163	1.0852
Number of qualified siRNAs (number of target gene > 1)	5,565	10,863	491
Average number of target genes of qualified siRNAs (number of target gene > 1)	2.0577	2.1370	2.1181
Number of selected siRNAs in first round analysis	142	94	162
Number of candidate-genes examined in first round analysis	229	229	227
Average examination frequency of each candidate-gene	1.153	1.170	1.110

From these result we can see that the number of qualified siRNA has important influence on the number of used siRNAs in first round analysis.



mediated dsRNA delivery. The dsRNA that was derived from siRNA-G55 silenced both TF15 and TF21 but not the randomly selected transcription factor TF60.

The first-round experimental result showed that siRNA-G55 successfully silenced both *PhaTF15* and *PhaTF21*, and the RNA levels of *PhaPR1* and *PhaNPR1* were affected (Figure 5). This scenario verified that we could observe a change in the target phenotype when a related gene is affected in the first-round analysis.



According to our strategy, we then select two siRNAs from qualified siRNAs that are specific to *PhaTF15* or *PhaTF21* for the second-round analysis. Because the complementary strand of siRNA-G55 also targeted a



Phalaenopsis gene (*PhaRNA*, accession number CB034920 [34]), we also designed an siRNA that was specific to this RNA sequence.

The second-round experimental data indicated that both *PhaPR1* and *PhaNPR1* were silenced in the *PhaTF15-* and *PhaTF21-*silenced plants, but not in the *PhaRNA-*silenced plant (Figure 6). These data suggested that *PhaTF15* and *PhaTF21* are involved in the SArelated plant defense response pathway but that *PhaRNA* is not. This result showed that our hierarchical strategy is capable of identifying genes that are involved in a specific phenotype.

Discussion

Our strategy provides a method for discovering new phenotypes that is an alternative to the traditional oneon-one RNAi experiments. The analysis of the mutant collection of model organisms revealed that single-gene knockouts did not usually exhibit an observable phenotype [57-59]. Our strategy facilitates the more efficient screening of single genes that are involved in a phenotype and the simultaneous targeting of several genes; this simultaneous targeting provides an opportunity to



observe additional phenotypes that can be observed only when multiple genes are silenced.

Although an siRNA can target its complementary genes, the number of mismatches that are allowed between the siRNA and its target gene(s) has not been strictly defined. According to our experimental results, at least 4 mismatches are allowed between the siRNA and its target gene(s) (Figure 4). Besides, previous reports have indicated that the targeting of the siRNA to the RNA could be affected by three, four or even five mismatches [13,50]. Therefore, using a threshold number of mismatches to distinguish the target and the nontarget genes of an siRNA could generate a prediction error. To prevent the unexpected target genes of a certain siRNA, in our qualified siRNA model, two userdefined parameters, d_T and d_N , were employed to predict the target and non-target genes of the qualified siRNAs, respectively (the genes within a d_T distance of the siRNA are its targets, whereas the genes that have at least a d_N distance to the siRNA are its non-targets). Theoretically, a smaller d_T / greater d_N should benefit the accuracy of the prediction of target/non-target genes because the probability of being a target gene tends to increase as the number of mismatches decreases. Nevertheless, according to our experimental results for the Orchid EST data (Table 2), a smaller d_T / greater d_N decreased the number of non-redundant qualified siRNAs and limited the reduction of the RNAi experiments. When the pairings of d_T and d_N were designed differently, they affected the number of RNAi experiments that would be required to screen the entire genome; however, as long as the d_T is set within a reasonable range ($d_T < 5$), each candidate-gene will be examined by at least one qualified siRNA.

In addition, our strategy allows the user to design two disjoint sets of genes, which are the candidate-gene set and the excluded-gene set. After that researchers narrow down the genes that could be associated with a specific phenotype into a set of candidate-genes by utilizing existing knowledge or an experimental analysis; this design can help them to focus their resources on the study of these candidate-genes. However, if a user cannot exclude any genes in advance or wants to perform a whole-genome analysis, our heuristic algorithm and hierarchical strategy can still assist in the reduction of the number of RNAi experiments that are required to perform a whole-genome analysis. We found that additional RNAi experiments can be eliminated when the excluded-gene set is empty. For example, for the orchid ESTs data, when the candidategenes included all of the mRNA sequences and the TF sequences and no genes were excluded, our heuristic algorithm identified 3,092,859 qualified siRNAs with the parameters L = 21, $d_T = 4$ and $d_N = 5$. Only 1,674 RNAi experiments would be required to analyze 7,796

candidate-genes in the first-round RNAi analysis because there were no excluded-gene hit restrictions on the qualified siRNAs; therefore, more qualified siRNAs that can target more genes could be obtained.

Previous studies have indicated that both strands of a dsRNA can be loaded into the RNA-induced silencing complex (RISC), and they can both facilitate RNAi [54-56]. Therefore, whenever a mutant phenotype is observed, the target genes of the qualified siRNA and the potential target genes of the complementary strand of the qualified siRNAs must be examined in the second-round RNAi analysis. This step is necessary because our qualified siRNA was initially designed to be complementary to the candidate-genes. According to our analysis, the average number of possible targets of the complementary strand of a qualified siRNA was 0.6511, which indicates that only a few additional genes required screening in the second-round analysis.

Schwab et al. [60] used artificial microRNAs (amiRNA) to investigate the specificity of miRNA and showed that amiRNA can target mRNAs with a few mismatches. Artificial microRNAs can knockdown single and multiple genes and thus are an effective tool for knocking down several related but not identical genes. Schwab et al. [60] developed a tool, Web MicroRNA Designer (WMD), to facilitate the design of amiRNAs for silencing of single and multiple genes. Our strategy and that by Schwab et al. [60] are similar in that they both design small RNA to target single or multiple genes. However, WMD allows users to design effective amiRNA to knockdown their interested genes, whereas our strategy aims to reduce the initial experiments needed in a genome-wide screening.

Our initial screening using hpRNA-G55 allowed us to knockdown PhaTF15 and PhaTF21 simultaneously (Figure 5), and a second round of analysis using siRNAs specific to *PhaTF15* and *PhaTF21* individually indicated that TF21, similar to the previously identified transcription factor TF15, is involved in plant disease resistance (Figure 6). Interestingly, the alignment of TF15 and TF21 revealed that these two TFs share a conserved region of approximately 180 nucleotides (see Additional file 1: Figure S2 for more detail). We also found that this conserved region contains a zinc finger AN1 domain, and the target region of siRNA-G55 is located in the region of this domain. To determine whether siRNA-G55 could silence all of the genes that contain the zinc finger AN1 domain, we queried the SWISS-PROT database for all of the genes of Arabidopsis thaliana that contain the zinc finger AN1 domain. Seventeen genes were reported to contain this domain, and only two of these seventeen genes may be affected by the siRNA-G55 (when set at d_T = 4). It will be interesting to further determine their roles in the plant defense response.

Conclusions

In this study, we developed a strategy to minimize the efforts required to perform high-throughput RNAi screening to identify genes involved in a specific phenotype. Our strategy took advantage of the biological phenomenon of the off-target effect of RNAi and utilized the set cover algorithm and the group testing method. We also provided a heuristic algorithm to design the siRNAs that can target multiple genes for our strategy. In the case study using the Orchid ESTs library, our computational result showed that 94 qualified siRNAs are sufficient to screen all 229 candidate-genes in the first-round analysis, where 229 siRNAs are required in traditional one-on-one RNAi screening. Furthermore, we verified our strategy with a previously reported transcription factor (TF15) involved in plant defense responses. In the first-round analysis, we used siRNA-G55, which is one of the 94 selected siRNAs, to silence TF15 and TF21 simultaneously, and the experimental data showed that the RNA levels of PhaPR1 and PhaNPR1 were subsequently affected. Moreover, in the second-round analysis, we successfully confirmed that TF15 is involved in plant defense responses only with three additional RNAi experiments. Additionally, we also indicated that TF21 is a previously unidentified transcription factor that is involved in plant defense responses.

Additional files

Additional file 1: Figures S1 and S2. A simple example of a powerful subsequence *P*, and the dotplot of TF15 and TF21 sequences.

Additional file 2: Proof to prove that the problem of finding a qualified sequence *r* maximizing the size of *T* is NP-complete.

Additional file 3: Methods the detail of our modified greedy algorithm for selecting qualified siRNAs to perform the first-round RNAi analysis.

Additional file 4: Tables S1-3. The lists of the siRNAs that were selected for first-round RNAi analysis with different parameters.

Abbreviations

RNAi: RNA interference; dsRNA: double-stranded RNA; RISC: RNA-induced silencing complex; TF: Transcription factor; ESTs: Expressed sequence tags; DSSP: Distinguishing substring selection problem; MMLV: Moloney murine leukemia virus; amiRNA: Artificial microRNA.

Competing interests

The authors declare that they have no competing interests

Authors' contributions

CHC designed and implemented the algorithms for the analysis, performed the computational analysis, and drafted the paper. HHY and HIW conceived of the study, participated in the design and drafted the paper. HHY, HCL and CEC performed the wet experiments. HHC provided the transcription factor sequence data. HHY and CYT provided constructive opinions and refined the manuscript. All authors read and approved the final manuscript.

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