PolymiRTS Database: linking polymorphisms in microRNA target sites with complex traits

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ABSTRACT

Polyorphism in microRNA Target Site (PolymiRTS) database is a collection of naturally occurring DNA variations in putative microRNA target sites. PolymiRTS may affect gene expression and cause variations in complex phenotypes. The database integrates sequence polymorphism, phenotype and expression microarray data, and characterizes PolymiRTS as potential candidates responsible for the quantitative trait locus (QTL) effects. It is a resource for studying PolymiRTS and their implications in phenotypic variations. PolymiRTS database can be accessed at http://compbio.utmem.edu/miRSNP/.

INTRODUCTION

Identification of causal genetic variants underlying complex traits is a major goal in genetic studies. Linkage analysis has long been used to discover chromosomal intervals harboring sequence variants that cause variations in quantitative traits. A typical quantitative trait locus (QTL) interval usually contains many genes ranging from several dozens to several hundreds, hence it is critical to be able to focus on genetic variants in the interval that are most likely to have functional impacts. Among them, nonsynonymous single nucleotide polymorphisms (SNPs) that alter protein sequences and regulatory polymorphisms that affect gene expression are natural high-priority candidates. Although regulatory polymorphisms are much more challenging to be identified and characterized, experimental and analytical tools are being actively developed for this purpose (1–4). Polymorphisms in miRNA target sites (PolymiRTS) represent a specific class of regulatory polymorphisms that may regulate posttranscriptional gene expression. A recent work reports that PolymiRTS can underlie the effects of physiological QTLs (pQTLs) that control classic higher order traits (5). MicroRNAs (miRNAs) are a family of small RNAs that pair to the transcripts of protein-coding genes and cause translational repression or mRNA destabilization (6,7). Hundreds of miRNAs have been identified in humans and mice and many of them have been shown to regulate their target genes that control diverse biological processes such as differentiation, proliferation, apoptosis and morphogenesis (7). PolymiRTS may affect the base-pairing process, hence affect the miRNA-mediated gene repression which in turn cause phenotypic variations. It has been found that miRNA-mediated target mRNA destabilization is widespread in mammals (8–10). Thus, for miRNAs acting by this mechanism, the PolymiRTS may lead to heritable variations in gene expression. Variations in gene expression across a population can be assessed by a newly developed genetical genomics approach (11–13). The genetical genomics approach treats gene expression level as quantitative trait. Linkage mapping is then used to discover the genetic loci regulating gene expression traits (eQTLs). PolymiRTS may induce a cis-acting eQTL that coincides with the gene’s physical location. We proposed a simple conceptual model (Figure 1) that represents information flow from PolymiRTS to complex trait via cis-acting eQTL. Based on this model scheme, we create a database integrating SNP, phenotype and expression microarray data of human and mouse.

DATA SOURCES AND PROCESSING

Identifying and annotating PolymiRTS

The method of identifying and annotating PolymiRTS is shown in Figure 2. SNPs that are located in the 3'-untranslated regions (3'-UTRs) of all known genes by UCSC genome annotation (mouse: mm7 and human: hg18) (14) were extracted from dbSNP build 126 (15). Genomic locations of these SNPs were mapped onto miRNAs. For each SNP, we assessed whether its two alleles lead to different miRNA target sites. To be conservative, we only consider...
the 3′-UTR SNPs that affect the match to the seed region of the miRNA. Mature miRNA sequences were downloaded from the miRBase (16). We used the criteria of TargetScanS (17) in the prediction of miRNA sites. Basically, besides requiring a perfect Watson–Crick match to the seed 2–7 nt of miRNA, we further require that there is either a perfect match to the 8th nt of miRNA, or an anchor adenosine immediately downstream the 2–7 seed in the target. We assigned the PolymiRTS to one of the four classes: ‘D’ (an allele disrupts a conserved miRNA site), ‘N’ (a derived allele disrupts a nonconserved miRNA site), ‘C’ (a derived allele creates a new miRNA site) and ‘O’ (other cases when the ancestral allele can not be determined unambiguously). PolymiRTS of class ‘C’ may cause abnormal gene repression and PolymiRTS of class ‘D’ may cause loss of normal repression control. These two classes of PolymiRTS are most likely to have functional impacts. We used the pre-calculated 17-way Multiz alignments of vertebrate genomes to derive the annotations. For a miRNA site to be conserved, we require that it is present in at least two other vertebrate genomes in addition to the query genome. For mouse SNPs, their ancestral alleles were determined by mouse versus rat (rn3) genome alignment. For human SNPs, their ancestral alleles were determined by human versus chimpanzee (panTro1) genome alignment. Additionally, we categorized PolymiRTS with A/G alleles because they are supposed to be less deleterious with their ability to form G:U wobble base-pairs with miRNAs.

PolymiRTSs within cis-acting expression QTL (eQTL) intervals

The genes with both cis-acting eQTL and PolymiRTS are featured in the database. First, gene expression levels in cerebellum, hippocampus, striatum, eye, whole brain and hematopoietic cell (18) were assessed in recombinant inbred mouse strains (BXD) derived from two parental strains C57BL/6j and DBA/2J. Gene expression levels were treated as quantitative traits and were mapped onto genomic regions (eQTL) using standard marker regression. A gene is said to have a suggestive (significant) cis-acting eQTL, if the LOD (log of odds) peak location is within 10 MB from the gene’s physical location and the LOD > 2.8 (>4.3) (19). Second, gene expression levels in lymphoblastoid cells of 194 human individuals from 14 CEPH families (20) were downloaded from the GEO database (21) and the raw data were processed by using the RMA protocol (22). Genotypes for 1628 autosomal SNP markers were downloaded from The SNP Consortium database (23). We used Merlin (24) to remove genotype errors and perform family-based linkage analysis. A gene is said to have a cis-acting eQTL, if the LOD peak location is within 10 MB from the gene’s physical location and the P-value is < 0.05.

PolymiRTSs in physiological QTL (pQTL) intervals

We first mapped the QTLs (with a LOD > 2.8) for more than 800 published BXD phenotypes (physiological/behavioral traits) (18). For each QTL, we linked it with genes that are physically located in the QTL interval and have at least one PolymiRTS. These genes are candidate causal genes underlying the pQTL.

DATABASE CONTENT AND ACCESS

Table 1 shows the major data fields for a typical PolymiRTS record. The users can access the database by several options. First, a web interface is implemented for browsing the entries. Second, a text search interface is designed for query by SNP ID, miRNA ID, GenBank accession, HUGO gene identifier and gene description. Third, a chromosome location search is offered so that the users can specify a genomic interval of interest and search all the PolymiRTSs within the interval. For mouse, we also provide the inbred strain comparison option. By combining strain comparison with the range...
search, the user can retrieve all the PolymiRTSs that are candidate variants underlying the pQTL identified by them. Finally, we provide flat file downloads for all the data.

**DISCUSSIONS AND FUTURE WORK**

miRNAs regulate posttranscriptional gene expression through mRNA destabilization or translational repression. Destabilization of mRNA may cause variations in the transcript levels while translational repression does not (25–27). The effects of miRNAs that act through translational repression cannot be detected by expression microarrays. Therefore, in the PolymiRTS database, the sequence variations in the target sites of miRNAs that act through translational repression may be linked to pQTLs but not to eQTLs.

A recent study shows that there are two broad categories of mRNA target sites: (i) 5’-dominant sites with sufficient 5’-pairing, and (ii) 3’-compensatory sites with insufficient 5’-pairing which require strong 3’-pairing (28). By using the TargetScanS algorithm alone, we are likely to miss many target sites from the second category. Because some mismatches in the 5’-pairing region of 3’-compensatory target sites are tolerated which is difficult to estimate the influence of SNPs on the function of 3’-compensatory target sites. Hence, at this stage we want to be conservative and focus on the major and well-studied category of 5’-dominant sites.

Currently, we did not take miRNA gene expression pattern into consideration due to lack of large-scale miRNA expression data. However, in many cases, such information would be very valuable. For example, it is known that a set of genes evolutionarily avoid having mRNA target site in their 3’-UTRs (8). They are called mRNA antitargets. Obviously, creation of a new mRNA site in an antitarget is most likely to have severe consequences. Since antitargets tend to be highly and specifically expressed in the tissue where the miRNA is expressed (8), we can use this information to pick PolymiRTS in the putative antitargets and further prioritize strong functional candidates. With the anticipated accumulation of such mRNA profiling data, we would expect a much better annotation of PolymiRTS in the near future.

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